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# A Rescorla-Wagner Drift-Diffusion Model of Conditioning and Timing

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#### Abstract

Computational models of classical conditioning have made significant contributions to the theoretic understanding of associative learning, yet they still struggle when the temporal aspects of conditioning are taken 10 into account. Interval timing models have contributed a rich variety of 11 time representations and provided accurate predictions for the timing of 12 responses, but they usually have little to say about associative learning. 13 In this article we present a unified model of conditioning and timing that 14 is based on the influential Rescorla-Wagner conditioning model and the 15 more recently developed Timing Drift-Diffusion model. We test the model 16 by simulating 10 experimental phenomena and show that it can provide 17 an adequate account for 8, and a partial account for the other 2. We argue 18 that the model can account for more phenomena in the chosen set than 19 these other similar in scope models: CSC-TD, MS-TD, Learning to Time 20 and Modular Theory. A comparison and analysis of the mechanisms in 21

these models is provided, with a focus on the types of time representation

and associative learning rule used.

### <sup>24</sup> Author Summary

How does the time of events affect the way we learn about associations between 25 these events? Computational models have made great contributions to our 26 understanding of associative learning, but they usually do not perform very 27 well when time is taken into account. Models of timing have reached high levels 28 of accuracy in describing timed behaviour, but they usually do not have much to 29 say about associations. A unified approach would involve combining associative 30 learning and timing models into a single framework. This article takes just this 31 approach. It combines the influential Rescorla-Wagner associative model with a 32 timing model based on the Drift-Diffusion process, and shows how the resultant 33 model can account for a number of learning and timing phenomena. The article 34 also compares the new model to others that are similar in scope. 35

## <sup>36</sup> 1 Introduction

Classical conditioning theories aim to understand how associations between 37 stimuli are learned. Ever since Pavlov (1927) the process of association forma-38 tion has been understood to depend crucially on the temporal relations between 39 stimuli (Savastano and Miller, 1998; Balsam et al., 2006; Kirkpatrick, 2013). 40 Yet, classical conditioning theories have so far struggled to work when time is 41 taken into account as an attribute of the stimulus representation. The study of 42 time as a mental representation is the object of a separate area of study known 43 as interval timing. Interval timing theories have produced a rich variety of time 44 representations (Gibbon et al., 1984; Killeen and Fetterman, 1988; Machado, 45

22 23 <sup>46</sup> 1997; Staddon and Higa, 1999; Matell and Meck, 2004), and therefore are a
<sup>47</sup> natural place to look for ways to integrate time into classical conditioning. In
<sup>48</sup> this paper we first analyse previous efforts in this direction before introducing
<sup>49</sup> a new hybrid classical conditioning and timing model.

The process of association formation is understood to be of fundamental sur-50 vival value for both human and non-human animals. Prediction, which forms 51 the core of classical conditioning, allows the organism to adapt to significant 52 events in its surroundings. A prototypical experiment in classical conditioning, 53 a type of associative learning, involves a neutral stimulus and an unconditioned 54 stimulus (US) which is capable of eliciting an unconditioned response (UR). 55 After repeated pairings of both stimuli in a specified order and temporal dis-56 tance, the neutral stimulus comes to elicit a response similar to the UR. This 57 response is called the conditioned response (CR) and the neutral stimulus is 58 said to have become a conditioned stimulus (CS). Classical conditioning theo-59 ries typically conceptualize this process as the formation of a link (association) 60 between the internal representations of CS and US. Their basic building blocks 61 are (Pearce and Bouton, 2001; Brandon et al., 2002): (a) the representations 62 of stimuli, and (b) a learning rule to update the association weights between 63 these representations. Although most theories do not attempt to find neuro-64 physiological correlates, these constructs are nonetheless commonly assumed to 65 be instantiated by (a) neural activity in the form of spike rates, and (b) synaptic 66 plasticity (Moore, 2002; Klopf, 1988; Gallistel and Matzel, 2013). These have 67 found some support in the neuroscientific literature, particularly studies of the 68 role of dopamine in reward prediction (Schultz et al., 1997; Dayan and Niv, 69 2008; Niv, 2009; Eshel, 2016). However it is important to note that there is still 70 no widely accepted complete neural mechanism for classical conditioning and 71 that most theories stay at the computational level of explanation. 72

Stimulus representations are generally thought of as neural activation that 73 is elicited by the stimulus, which may linger for a short time as a 'trace' af-74 ter stimulus offset. Representations are commonly one of two types: molar or 75 componential. Molar (or elemental) trace theories treat the stimulus as a single 76 conceptualized unit whose activity is usually assumed to peak quite early fol-77 lowing stimulus onset, and then gradually decrease (Hull, 1943; Wagner, 1981; 78 Sutton and Barto, 1981; Schmajuk and Moore, 1988; McLaren and Mackintosh, 79 2000; Harris and Livesey, 2010). In contrast, componential trace theories break 80 down the CS representation into smaller units, each capable of being associated 81 with the US, with some units more active early during the CS and others late, 82 but all leaving a trace after activation (Desmond and Moore, 1988; Grossberg 83 and Schmajuk, 1989; Vogel et al., 2003; Ludvig et al., 2008). 84

Learning rules may be classified according to different criteria. An important 85 period in the recent history of the field gave rise to one of these criteria. Prior to 86 1970's conditioning used to be rooted in the stimulus-response tradition, which 87 attributed crucial importance to the temporal pairing, or contiguity, of stimuli 88 for the development of associations. The linear operator learning rule (Hull, 89 1943) is one of the products of that period. In the late 1960's and early 1970's 90 important experimental discoveries using compound stimuli, that is, a stimulus 91 formed by combining other individual stimuli, showed the contiguity view to 92 be incomplete (Rescorla, 1988; Gallistel and Gibbon, 2001). These compound 93 experiments indicated that the formation of associations also depended on the 94 reinforcement history of the individual elements forming the compound stimu-95 lus. This led to the development of new learning rules (Rescorla and Wagner, 96 1972; Mackintosh, 1975; Pearce and Hall, 1980) capable of combining individual 97 reinforcement histories in compounds, which the linear operator rule cannot. 98 The first, and arguably still the most influential, of these learning rules is the 99

Rescorla-Wagner (RW, Rescorla and Wagner, 1972). It has become famous for
being the first model able to provide an account for the blocking effect (Kamin,
1968), where a novel CS does not become associated with the US if it is reinforced only in compound with a previously conditioned CS.

The CR is usually not a single event. Organisms time their responses so 104 that they emerge gradually during the duration of the CS and reach maximum 105 frequency or intensity around the time of reinforcement. Interval timing theories 106 have attempted to provide an account for this *timing* of the CR. One of the 107 fundamental properties of timing behaviour is that it is approximately timescale 108 invariant, i.e. the whole response distribution scales with the interval being 109 timed (Gibbon, 1977; Allman et al., 2014). One of the consequences of timescale 110 invariance is that the coefficient of variation, that is the standard deviation 111 divided by the mean, of the dependent measure of timing is approximately 112 constant. A number of timing models have put forth explanations for timescale 113 invariance and other timing properties (how time is encoded, how it is stored in 114 memory and how it gets translated into behaviour) by recourse to an internal 115 pacemaker. The most influential pacemaker-based timing theory to date is 116 Scalar Expectancy Theory (SET, Gibbon et al., 1984; Gibbon and Church, 117 1984). The pacemaker is supposed to mark the passage of time by emitting 118 pulses. These pulses can be gated to an accumulator via a switch which closes 119 at the start of a relevant interval and opens when the interval is finished. The 120 accumulator count is kept in working memory. At the end of the interval the 121 current count is transferred to a long-term reference memory. Behaviour is 122 guided by the action of a comparator which actively compares the count in 123 working memory to the one retrieved from reference memory. 124

In spite of the considerable overlap, interval timing and classical conditioning are not easily integrated. Most conditioning theories are trial-based, that is they

consider the trial as the unit of time. A trial is generally taken to be the state 127 where a CS is present (or CSs in compound) and which may or may not contain 128 a US (or USs). The most influential model in this category is the Rescorla-129 Wagner (RW, Rescorda and Wagner, 1972). In order to account for different 130 stimulus durations, trial-based theories like RW must resort to some sort of 131 time discretization, usually by subdividing the trial into 'mini-trials'. Each 132 mini-trial is treated as a trial in its own right, which are then used to update 133 associative links. This gives rise to the problem of deciding on a particular 134 discretization. Also, given that humans experience time passing as a continuous 135 flow, it is unlikely that animals discretize their conditioning experience in such 136 a way. A more realistic approach to timing is taken by real-time theories. These 137 theories attempt to formalize the concept of a continuous flow of time. 138

The Temporal Difference model (TD, Sutton and Barto, 1990,9) was one 139 of the earliest and still most influential real-time classical conditioning model. 140 It may be thought of as a real-time version of RW. When used with stimulus 141 representations such as the Complete Serial Compound (CSC, Moore et al., 142 1998), Microstimuli (MS, Ludvig et al., 2008,0) and the Simultaneous and Se-143 rial Configural-cue Compound (SSCC, Mondragón et al., 2014) it is capable of 144 reproducing some timing phenomena like the gradual increase in anticipatory 145 responding that occurs before a signalled reinforcer, and the lower response rates 146 observed during longer CSs. However, only MS-TD has a time representation 147 capable of approximating the most fundamental property of timing, timescale 148 invariance. Another issue with the stimulus representations for TD is that their 149 approach to timing resembles the strategy used by trial-based models, i.e. they 150 all split the stimulus into a number of smaller units or states, the number of 151 which being directly proportional to the duration of the stimulus. Given that 152 conditioning is observed in a timescale that ranges from milliseconds to hours 153

(Kehoe and Macrae, 2002, p. 189) this can lead to a very high number of units 154 being required. The stimulus as a whole no doubt is a complex entity, and the 155 brain may be employing a large number of neurons to represent it, but to ded-156 icate so many resources only for timing might not be the most energy-efficient 157 strategy. Also, TD and its stimulus representations do not usually account for a 158 change in timing that is not tied to reinforcement. Animals time the occurrence 159 of different events, such as onset and offset of stimuli (see for example Meck and 160 Church, 1984), but TD usually only allows for the timing of rewards. 161

On the other hand, timing models have made even fewer attempts at inte-162 grating aspects of classical conditioning. A notable exception is the Learning 163 to Time (LeT, Machado, 1997; Machado et al., 2009) model. It represents the 164 passage of time by transitioning between internal states according to a stochas-165 tic pacemaker, an idea borrowed from an earlier timing model called the Be-166 havioural Theory of Time (Killeen and Fetterman, 1988). Learning takes place 167 by associating reinforcement presentation with the current internal state accord-168 ing to the linear operator, a standard classical conditioning rule. LeT offers an 169 account of the basic dynamics of association formation, but it cannot explain 170 cue-competition phenomena like blocking. In a blocking procedure, a CS is first 171 paired with a US until a CR is acquired. The same CS is then presented together 172 with a novel CS and both are paired with the US for a few trials. If the novel 173 CS is now presented alone it elicits little or no responding, and so it is said to be 174 blocked by the first CS. LeT's learning rule, the linear operator, has largely been 175 supplanted by RW in classical conditioning modelling because it cannot explain 176 cue-competition phenomena. Like TD, LeT also employs a representation that 177 requires as many units as time-steps, making it a resource-intense model. 178

Modular Theory (MoT, Guilhardi et al., 2007; Kirkpatrick, 2002) is a timing
model which because of its explicit goal of integrating timing and learning may

be called a hybrid theory. MoT has introduced novelties that allow it to account 181 for some aspects of the dynamics of classical conditioning that LeT cannot. Its 182 architecture is different than the connectionist one (states or units connected 183 by modifiable links) assumed by RW, TD and LeT. Instead, it uses a more cog-184 nitive architecture, with separate information processing stages that deal with 185 perception, memory and decision. It postulates two separate memories: a pat-186 tern memory which stores CS durations, and a strength memory which stores 187 the associative strength between each pattern memory and the US. This sepa-188 ration allows MoT to deal with more complex situations involving the dynamics 189 of learning during acquisition and extinction. However, MoT also relies on the 190 linear operator to update its strength memory, which, like LeT, prevents it from 191 accounting for cue-competition phenomena. 192

Although the models mentioned above, namely TD, LeT and MoT, have 193 accomplished a great deal in terms of bringing together timing and conditioning, 194 they each have their different strengths and weaknesses as we have touched 195 above. In this paper we introduce a model that tries to address some of these 196 weaknesses while preserving the strengths. More specifically, the model has the 197 following strengths. It represents time in real-time. Like MoT and unlike LeT 198 and TD, its time representation does not require an arbitrary large number of 199 units or states. Similarly to TD but unlike LeT and MoT, it uses a learning rule 200 that preserves the main features of RW which allow it to account for compound 201 phenomena. It can time the onset and offset of all stimuli, not only of rewards, 202 and store a memory for each. It includes two update rules: one for timing that 203 is updated by time-markers, and another for associations that is updated by 204 the US. Hence, simple stimulus exposure causes the model to learn and store 205 its duration. This capability is not present in models that depend only on an 206 associative learning rule to also learn about time, such as TD and LeT. 207

This new model is essentially a way to connect one of the most influen-208 tial classical conditioning theories, the Rescorla-Wagner model (Rescorla and 209 Wagner, 1972), with a recently developed timing theory called Timing Drift-210 Diffusion Model (TDDM, Rivest and Bengio, 2011; Simen et al., 2011). The 211 TDDM is based on the drift-diffusion model, widely used in decision making 212 theory, and it provides an adaptive time representation that has commonalities 213 with pacemaker-based models like SET and LeT (Simen et al., 2013). These 214 models postulate the existence of a pacemaker that emits pulses at a regular 215 rate, which are then counted to mark the passage of time. To preserve timescale 216 invariance they either postulate a specific type of noise in the memory saved for 217 intervals and a ratio-based decision process (SET) or adapt the rate of pulses 218 (LeT). The TDDM takes the latter route but sets a fixed threshold on pulse 219 counting. To emphasize the unification of these two theories we call our pro-220 posal the Rescorla-Wagner Drift-Diffusion Model (RWDDM). 221

We evaluate RWDDM based on how well it can simulate the behaviour of 222 animals in a number of experimental procedures. Many classical conditioning 223 phenomena have been identified which collectively represent a significant chal-224 lenge for any single model to explain. A recent list (Alonso and Schmajuk, 2012) 225 has compiled 12 categories, which include acquisition, extinction, conditioned in-226 hibition, stimulus competition, preexposure effects, temporal properties, among 227 others. Of particular interest to a theory of timing and conditioning are phe-228 nomena that involve elements of both timing and conditioning. As we detail 229 later, we have searched the literature for documented effects that can challenge 230 the main mechanisms embodied in RWDDM. 231

We proceed by first introducing the new model. We compare its formalism with four models that have similar scope, namely CSC-TD, MS-TD, MoT and LeT. In the results section we present the phenomena we will simulate, followed <sup>235</sup> by the results of our simulations, and compare them to the current explanations<sup>236</sup> given by LeT, MoT and TD.

## 237 **2** Model

We follow most classical conditioning theories in conceptualizing the condition-238 ing process as the formation of an association between the internal representa-239 tions of CS and US. Arguably, one of the most influential rules describing the 240 evolution of this association through training is the Rescorla-Wagner (Rescorla 241 and Wagner, 1972) rule. As mentioned previously, other models exist which 242 have a similar scope to RW, both trial based (Mackintosh, 1975; Pearce and 243 Hall, 1980) and real-time (Buhusi and Schmajuk, 1999; McLaren and Mack-244 intosh, 2000,0). However, our goal was to take advantage of TDDM's time 245 representation, so we sought a theoretical associative framework that could in-246 corporate such a representation. Since trial-based conditioning theories lack 247 any time representation, they are a natural place to start. Out of those theories 248 the RW is perhaps the simplest whilst also retaining the greatest possible ex-249 planatory power. Its basic formalism consists of the following rule for updating 250 associative strength: 251

$$\Delta V_i(n) = \alpha \beta \left( \lambda - \sum_{j=1}^l V_j(n) x_j(n) \right) x_i(n) \tag{1}$$

where  $V_i(n)$  denotes associative strength for  $CS_i$  at trial  $n, \lambda$  the asymptote of learning which is set by the US representation,  $x_i(n)$  which marks the presence  $(x_i = 1)$  or absence  $(x_i = 0)$  of the i-th CS representation at trial n, 0 < $\alpha < 1$  a learning rate set by the CS and  $0 < \beta < 1$  a learning rate set by the US. The summation term in the equation (1) sums over all CSs present in the trial. The top panel of figure 1 shows a diagram of a basic neural net

for classical conditioning which serves as the architectural framework for both 258 RW and RWDDM. The RW rule is used to update the links  $V_1, ..., V_l$  that 259 connect the CS input nodes  $CS_1,...,CS_l$ . The summation term in the RW rule 260 is represented in the diagram as a summation unit or junction  $\Sigma$ , that sums 261 the inputs it receives from the CSs j = 1, ..., l present in the trial. This sum 262 allows RW to combine (additively) the reinforcement history of each individual 263 CS present in a compound trial. In the neural network literature, equation (1)264 is also referred to as the Widrow-Hoff rule (Widrow and Hoff, 1960) and the 265 Least-Means-Square (LMS; Sutton, 1992). The relationship to the LMS rule 266 is easier to see if we let  $y(n) = \sum_{j=1}^{l} V_j(n) x_j(n)$  be the output of a learning 267 unit that aims to predict a target  $\lambda$  given inputs  $x_i$  by adapting the weights 268  $V_i$ . In classical conditioning,  $\lambda$  represents the maximum learning driven by a 269 given outcome (the US),  $x_i$  is the CS and  $V_i$  the associative strength. If we 270 let  $\delta(n) = \lambda - y(n)$  be the error between output and US, equation (1) can be 271 obtained with the method of gradient descent by minimizing the squared error 272  $\delta^2(n)$  with respect to the weight  $V_i$ . 273

In spite of the relative success in explaining a wide range of conditioning 274 phenomena (for a list of successes, and failures, see Miller et al., 1995), the 275 Rescorla-Wagner rule lacks a mechanism to account for the microstructure of 276 real-time responding during conditioning procedures. In terms of the order of 277 CS-US presentation conditioning procedures may be either forward (CS followed 278 by US) or backward (US followed by CS). Two common types of forward condi-279 tioning are delay and trace. In delay conditioning the US always occurs a fixed 280 time after CS onset. In trace conditioning the US occurs at a fixed duration after 281 CS offset. After sufficient training with delay or trace conditioning, responding 282 begins some time after CS onset, increases rapidly in frequency until it reaches 283 a maximum level where it stays until US onset (Gormezano et al., 1983). The 284



Figure 1: Connectionist diagram of RWDDM. Each CS unit is connected to a summing junction (labelled  $\Sigma$ ) via a modifiable link V. The output of the summing junction is the CR. The US is represented as a teaching signal with a fixed weight H. Each CS unit has its own timer  $\Psi$  and representation x. The bottom panel shows a zoomed-in view of the timer  $\Psi_l$  and CS representation  $x_l$ associated with CS<sub>l</sub>. The timer slope  $A_l$  is tuned to a 5-second CS duration.

RW rule alone does not account for CR level as a function of time. This role 285 is usually fulfilled by the choice of CS representation. We base our choice on a 286 timing model called Timing Drift-Diffusion Model (TDDM, Simen et al., 2011; 287 Rivest and Bengio, 2011; Luzardo et al., 2013; Balci and Simen, 2016). We chose 288 the TDDM because it possesses a number of interesting features. It is part of a 289 family of pacemaker based models like SET and LeT (Simen et al., 2013) which 290 are arguably two of the most successful timing theories to date. The TDDM 291 is a modified version of the drift-diffusion models that have been extremely 292

successful at modelling reaction time in decision making tasks (Ratcliff, 1978; 293 Voss et al., 2013). Evidence of climbing neural activity related to timing that 294 resembles the TDDM has been extensively reported (Komura et al., 2001; Leon 295 and Shadlen, 2003; Brody et al., 2003; Wittmann, 2013; Jazayeri and Shadlen, 296 2015). The TDDM consists of a drift-diffusion process with an adaptive drift or 297 rate. The drift-diffusion process is defined by a continuous random walk called 298 Wiener diffusion process. The two main components of Wiener diffusion are 200 the drift and the normally distributed noise. The Wiener diffusion process may 300 be visualized by imagining a two-dimensional grid with time in the horizontal 301 axis and displacement on the vertical axis. If we imagine a purely linear and 302 non-random walk that starts at the origin and moves up at a constant rate then 303 the resulting walk would be a straight line and the drift would be equal to the 304 slope of the line. With normally distributed noise, the walk becomes a random 305 walk and it looks like a jagged curve, since at each time step there is now only 306 a probability that the displacement will be up or down. For the purposes of 307 timing, the slope is always positive and the random walk can be interpreted as 308 a noisy accumulator (or timer)  $\Psi(t)$ , which starts at the beginning of a salient 309 stimulus and stops (and resets) at the end. In a conditioning experiment the CS 310 is usually the most salient stimulus in the uneventful context of the conditioning 311 chamber, so it is well placed to serve as a time marker. When timing starts, 312 accumulator increments are performed at each time-step according to 313

$$\Delta \Psi_i(t) = A_i(n) \cdot \Delta t + m \cdot \sqrt{A_i(n) \cdot \Delta t} \cdot \mathcal{N}(0, 1), \qquad (2)$$

where  $A_i(n)$  is the rate (slope) of accumulation for  $CS_i$  in trial n, m is a noise factor,  $\Delta t$  is the time-step size and  $\mathcal{N}(0, 1)$  denotes a sampling from the standard normal distribution. An interval is timed by the rise in the accumulator to a certain fixed threshold, say  $\Psi_i(t) = \theta$ . The TDDM adjusts to new intervals by keeping the threshold fixed but adapting the rate of accumulation  $A_i(n)$ . The bottom left panel of figure 1 shows a typical trajectory (or realization) of a CS's TDDM timer after one 5-second trial.

In its original formulation (Rivest and Bengio, 2011; Simen et al., 2011) the 321 accumulation process was not allowed to continue beyond the threshold value  $\theta$ , 322 constraint that gave rise to two distinct rules for rate adaptation, one for when 323 the US arrived earlier than expected and another for when it arrived later. The 324 constraint fixing a maximum level of accumulation was driven by the neurophys-325 iological assumption that a linear neural accumulator is not likely to continue 326 to perform effectively beyond a certain level. The neural implementation so far 327 proposed for TDDM's linear accumulator (Simen et al., 2011) is based on a feed-328 back control mechanism that is tuned to balance excitation and inhibition in a 329 neuron population. Tuning of this kind requires great computational precision, 330 which may not be easily kept for very long in a biological system. Neurophysiol-331 ogy notwithstanding, we will drop that requirement here for simplicity and use 332 instead only one update rule. We demonstrate how this single update rule can 333 be derived by the method of gradient descent. The model learns a new interval 334 by adapting its slope  $A_i$  so that the accumulator  $\Psi_i$  reaches the threshold value 335  $\theta$  at the target time  $t^*$ , which may be the time of reinforcement for example. 336 The target slope will therefore be  $\theta/t^*$ . The error  $\delta(n)$  between the target slope 337 and the current slope is  $\delta(n) = \theta/t^* - A_i(n)$ . By minimizing the squared error 338  $\delta^2(n)$  using gradient descent we can derive the slope update rule. The squared 339 error as a function of  $A_i$  forms a curve. Moving in the direction opposite the 340 slope of this curve and taking a step of size  $\alpha_t/2$  we form the equation: 341

$$A_i(n+1) = A_i(n) - \frac{\alpha_t}{2} \frac{d\delta^2(n)}{dA_i(n)}.$$
(3)

342 Solving the derivative yields

$$A_{i}(n+1) = A_{i}(n) - \frac{\alpha_{t}}{2} 2\delta(n)(-1)$$
  
=  $A_{i}(n) + \alpha_{t} \left(\theta/t^{*} - A_{i}(n)\right).$  (4)

Since the organism only has access to the psychological time given by its internal timing mechanism, and not the physical time t, we assume that an internal estimate for t is formed by dividing the current pacemaker count by the current slope,  $t = \Psi_i(t)/A_i(n)$ . Substituting this estimate into equation (4) we get:

$$A_{i}(n+1) = A_{i}(n) + \alpha_{t} \left(\frac{\theta A_{i}(n)}{\Psi_{i}(t^{*})} - A_{i}(n)\right)$$
$$= A_{i}(n) + \alpha_{t} A_{i}(n) \left(\frac{\theta}{\Psi_{i}(t^{*})} - 1\right)$$
$$= A_{i}(n) + \alpha_{t} A_{i}(n) \frac{(\theta - \Psi_{i}(t^{*}))}{\Psi_{i}(t^{*})}.$$
(5)

Hence, the update rule for slope  $A_i$  to be applied at target time  $t^*$  (the end of the trial or of the interval being timed) is

$$\Delta A_i(n) = \alpha_t A_i(n) \frac{(\theta - \Psi_i(t^*))}{\Psi_i(t^*)}.$$
(6)

Equation (6) is the slope update rule we use. Note that n above is indexing 349 the number of occurrences of a specific interval that the timer is timing. These 350 intervals may be the duration between CS onset and US onset (the usual 'trial' in 351 delay conditioning for example), but they may be any other salient time interval 352 such as CS or intertrial duration. Figure 2 shows timer slope adaptation during 353 three timing scenarios: timing a novel stimulus (row 1), timing a long-short 354 change in stimulus duration (row 3), and timing a short-long change in stimulus 355 duration (row 5). 356



Figure 2: RWDDM timer and CS representation during three 12-trial timing scenarios. Top two rows: timing a novel 6 second stimulus. Timer starts with a low baseline slope (A = 0.001) on trial 1 and gradually adapts over training to reach approximately the required slope. Middle two rows: stimulus duration change from 6 to 3 seconds. Bottom two rows: stimulus duration change from 6 to 12 seconds. Parameters:  $\alpha_t = 0.215$ ,  $\theta = 1$ ,  $\sigma = 0.25$ , m = 0.15.

In the top row of figure 2 and throughout the paper we assume that the initial 357 value of slope A for a novel stimulus is so low as to overestimate the stimulus 358 duration. This overestimation will only last for a few trials, the number of 359 which can be made arbitrarily small by choosing a high adaptation rate  $\alpha_t$ . 360 Alternatively, it would be possible to use a very high initial value for A so as to 361 underestimate the stimulus duration. However this alternative does not seem 362 neurophysiologically plausible as the brain would need to keep a pool of neurons 363 firing very rapidly as its 'standby' timer. 364

In TDDM, timescale invariance arises from the nature of the noise in the 365 accumulator. After repeated training, say in delay conditioning with a CS of 366 fixed duration, equation (6) will converge to a value of  $A_i$  which will make the 367 accumulator reach the threshold value  $\theta$  at the time of stimulus offset, but only 368 on average. In some trials the accumulator will reach the threshold sooner, in 369 which case the organism will underestimate the stimulus duration. In other trials 370 the accumulator will reach the threshold later, causing overestimation. The 371 variability of this time estimate relative to the mean is given by the coefficient 372 of variation (CV). It has been well established experimentally that the CV of 373 time estimates in humans and other animals is approximately constant over a 374 wide timescale (Gibbon, 1977; Gallistel and Gibbon, 2000; Allman et al., 2014). 375 The CV of TDDM's time estimate is (see equation 3 in Luzardo et al., 2017) 376

$$CV = \frac{m}{\sqrt{\theta}},\tag{7}$$

which depends only on the choice of threshold  $\theta$  and noise factor m. As these are constant, the CV of TDDM's time estimate is also constant. Note that because the timer adapts its slope gradually, if the duration of a CS is changed, CV measurements will only match the one given by equation (7) after the slope has finished adapting. The number of trials to adaptation will vary depending on the adaptation rate  $\alpha_t$ .

We substitute the presence representation used in the original RW model by a Gaussian radial basis function. Its input is provided by the TDDM accumulator:

$$x_i(\Psi_i) = \exp\left(-\frac{(\Psi_i(t) - \theta)^2}{2\sigma^2}\right).$$
(8)

This representation may be interpreted as the receptive field of time-sensitive neurons that read the signal coming from the accumulator neurons. Their re-

ceptive fields are tuned to the accumulator threshold value  $\theta$ . The bottom right 387 panel in figure 1 shows the representation for  $CS_l$  generated from the input 388 provided by the timer on the left. Note how  $x_l$  reaches its maximum value at 389 the same time that  $\Psi_l$  crosses the threshold at 1. Figure 2 shows  $x(\Psi)$  adapting 390 in the three different timing scenarios explained previously. As can be seen, 391  $x_i$  is a dynamic representation of  $CS_i$  that adapts to the temporal information 392 conveyed by the stimulus. Other representation shapes could be used, like a 393 sigmoid for example, but a Gaussian is mathematically simple and has been 394 used before by at least one other timing model (MS-TD, Ludvig et al., 2008). 395

We follow Gibbon (1977) and Gibbon and Balsam (1981) in assuming that 396 time sets the asymptote of learning,  $\lambda$ , in equation (1). They were led to this 397 hypothesis by investigating CR timing in fixed interval conditioning schedules, 398 a type of delay conditioning. After enough training in this procedure, subjects 399 begin responding some time after CS onset, with a slow rate at first which then 400 increases rapidly until it reaches asymptotic level some time before reinforcement 401 delivery. Gibbon (1977) proposed that subjects make an estimate of time to 402 reinforcement which is used to generate an expectancy of reinforcement. The 403 expectancy for a particular  $CS_i$  with duration  $t^*$ ,  $h_i$ , was hypothesised to be 404  $h_i = H/t^*$ , where H was a motivational parameter which was assumed to depend 405 on the reinforcing properties of the US. The reinforcing value of the US is 406 thus spread evenly over the CS length. It was assumed that this expectancy 407 would be updated as time elapsed during the CS, such that  $h_i(t) = H/(t^* - t)$ 408 t). Hence, expectancy would increase hyperbolically until the estimated time 409 to reinforcement  $t = t^*$ . Responding would reach asymptotic level when the 410 expectancy crossed a threshold value  $h_i(t) = b$ . 411

Here we will not use Gibbon's concept of expectancy update. A similar role is fulfilled by the TDDM accumulator in our formalization. But we hold on to

his argument that the reinforcing value of the US is spread over the CS length. 414 Within the Rescorla-Wagner modelling framework, Gibbon's expectancy value 415 may be interpreted as setting the asymptotic level of learning in equation (1), 416 namely  $\lambda = H/t^*$ . Under this interpretation,  $\lambda$  may be said to implement 417 hyperbolic delay discounting of rewards. Similarly to the argument used above 418 in the derivation of the slope update rule, we use the psychological time estimate 419 from TDDM in place of the physical time  $t^*$ , such that  $t^* = \Psi_i(t^*)/A_i(n)$ . 420 The value we use is then  $\lambda = \frac{HA_i(n)}{\Psi_i(t^*)}$ . Another possibility would be simply 421  $\lambda = HA_i(n)$ . Both alternatives yield the same asymptotic value, but  $HA_i(n)$ 422 converges gradually (with the rate set by  $\alpha_t$ ) whilst  $\frac{HA_i(n)}{\Psi_i(t^*)}$  immediately. Our 423 version of equation (1) for updating associative strength then becomes: 424

$$\Delta V_i(n) = \alpha_V \left( \frac{HA_i(n)}{\Psi_i(t^*)} - \sum_{j=1}^l V_j(n) x_j(\Psi_j) \right) x_i(\Psi_i).$$
(9)

In the trial-based RW model, equation (1) is applied at the end of a 'trial', which is usually taken to be the event starting at CS onset and ending at US delivery. We follow the same practice here and apply equation (9) at the end of a trial, i.e. at US delivery. Note that because  $x_i(\Psi_i)$  is a dynamic CS representation, its activation (or strength) level at the end of the trial will vary from trial to trial, as can be seen in figure 2. Equation (9) is applied using the activation level of  $x_i(\Psi_i)$  current at the end of the trial.

We assume that real-time responses to a  $CS_i$  are emitted according to the product of its associative strength  $V_i(n)$  and representation  $x_i(\Psi_i)$ , that is, it is the output of the summing junction in figure 1:

$$CR_i(t) = V_i(n)x_i(\Psi_i).$$
(10)

435

Equations (2), (6), (8), (9), (10) fully define the basic model. Its six free

<sup>436</sup> parameters are:  $m, \alpha_t, \theta, \sigma, \alpha_V, H$ .

#### 437 2.1 Relationship with Other Models

Among the theories capable of providing an account of both timing and conditioning, arguably four stand out for their scope or influence. They are CSC-TD,
MS-TD, LeT and MoT.

TD has been developed primarily as a learning model, without the explicit intention of addressing timing. It may be visualized as a real-time rendition of the RW rule. Its basic learning algorithm, is given by:

$$V_t(\mathbf{x}_t) = \sum_i w_t(i) x_t(i), \tag{11}$$

$$\delta_t = \lambda_t - (V_t(\mathbf{x}_{t-1}) - \gamma V_t(\mathbf{x}_t)), \tag{12}$$

$$\mathbf{w}_{t+1} = \mathbf{w}_t + \alpha \delta_t \mathbf{e}_t \tag{13}$$

where  $V_t$  is the US prediction at time t, formed by a linear combination of 444 the weights w(i) and the CS representation values x(i). This update algorithm 445 is performed at each time step, and not only at the end of a trial like RW 446 and RWDDM. Another important difference is that equation (12) computes a 447 difference between the current US value and the temporal difference between 448 predictions. Hence,  $\delta_t > 0$  if the US is higher than this temporal difference in 449 prediction, and  $\delta_t < 0$  if the US is lower. The constant  $0 < \gamma < 1$  is termed a 450 discount factor. Equation (13) updates the weights for the next time step. The 451 vector  $\mathbf{e}_t$  stores *eligibility traces*, which are functions describing the activation 452 and decay of representations  $\mathbf{x}_t$ . The three most common eligibility traces used 453 are: accumulating traces, bounded accumulating and replacing traces. These 454 three types accumulate activation in the presence of the CS and discharge slowly 455 in its absence, the first accumulates with no upper bound, the second only until 456

the upper bound and the third is always at the upper bound whilst the CS is
present (Sutton and Barto, 1998, pp. 162-192).

The richness of TD's timing account relies on the choice of CS representation 459 x. The Complete Serial Compound representation (CSC, Moore et al., 1998) 460 postulates one CS element x(i) per time unit of CS duration. Each element is 461 only switched on at its activation time unit, and then decays afterwards following 462 its choice of eligibility trace e(i) (usually an exponential decay function). This 463 componential representation, which increases in size linearly with CS duration, 464 should be contrasted with RWDDM's molar representation (equation (8)) which 465 requires only one element. CSC may be called a time-static representation, 466 whilst RWDDM is a time-adaptive representation, with a rule to change its 467 structure based on a change in time (equations (6) and (8)). CSC-TD also lacks 468 any mechanism to explain timescale invariance of the response curve, which is 469 present in RWDDM. A modification of CSC has recently been developed, the 470 Simultaneous and Serial Configural-Cue Compound (SSCC, Mondragón et al., 471 2014). SSCC-TD formalizes the idea that when multiple stimuli are presented 472 together in time, a configural cue-a novel stimulus that is unique to the current 473 set of present stimuli-is formed. SSCC follows on the CSC representation, but, 474 unlike any other TD model, it allows for the representation of compounds and 475 configurations of stimuli. Because SSCC-TD is a real-time model, it also allows 476 for the simulation of CR timing during compounds and configurations. However, 477 its approach to timing is still the same as CSC, i.e. it breaks down the stimuli 478 into a series of elemental units which are activated in series. Therefore, with 479 respect to timing only we will consider SSCC to belong to the family of CSC 480 representations. 481

The Microstimuli representation (Ludvig et al., 2008,0) introduced a more realistic description of time. Unlike CSC, it uses a fixed number of elements  $_{484}$  x(i) per stimulus. The ith microstimulus is given by:

$$x_t(i) = \frac{1}{\sqrt{2\pi}} \exp\left(-\frac{(y_t - i/m)^2}{2\sigma^2}\right) \cdot y_t \tag{14}$$

where m is the total number of microstimuli, y is an exponentially decaying time 485 trace set at 1 at CS onset. It will be noted that a microstimulus is a Gaussian 486 curve modulated by the decaying trace  $y_t$ . The set of microstimuli generated by 487 the CS will then give rise to partially overlapping Gaussians, with decreasing 488 heights and increasing widths across time. The fact that only a fixed number 489 of microstimuli are required per CS is an improvement to the potentially large 490 numbers of elements in CSC. The MS representation tries to capture the idea 491 that as time elapses, the stimulus leaves a more diffuse and faint impression. 492 However, even though it is more realistic than CSC, it still lacks a mechanism 493 to produce exact timescale invariance. 494

Learning to Time is primarily a theory of interval timing which can also 495 account for some aspects of conditioning. Here we will deal with its most recent 496 version in Machado et al. (2009), which differs somewhat from the earlier version 497 in Machado (1997). Its CS representation resembles CSC in postulating a long 498 series of elements (or states) that span the whole stimulus duration. Unlike 499 CSC, it transitions from state to state at a rate that varies from trial to trial, 500 and that is normally distributed. Hence, time during a trial is represented as 501 a noiseless linear increase from states n = 1, 2, 3, ... (one per time-step) at a 502 fixed rate. This linear time representation resembles the linear accumulator in 503 RWDDM, except that the latter has noise built into the linear accumulator, 504 whilst LeT assumes noise only at the intertrial level. Each state n is associated 505 with the US via an associative link. At the end of a trial, the strength w of 506 these links are updated as follows: 507

• For the active state at reinforcement,  $n^*$ , the update rule is

$$\Delta w(n^*) = \beta (1 - w(n^*)), \tag{15}$$

509 where  $\beta$  is a constant.

• For inactive states,  $n < n^*$ , the update rule is

$$\Delta w(n) = -\frac{\alpha}{n^*} w(n), \tag{16}$$

sin where  $\alpha$  is a constant.

512

• For states that did not become active during the trial,  $n > n^*$ , the rule is

$$\Delta w(n) = 0. \tag{17}$$

Note that unlike RWDDM's associative update rule, equations (15) to (17) do 513 not include a summation term. This places a severe limitation on the ability 514 of LeT to deal with compound conditioned stimuli. LeT's strength lies on its 515 being able to explain timescale invariance of the response curve. Machado et al. 516 (2009) showed that it is possible to derive timescale invariance using only the 517 assumption of intertrial normality of state transition rate. Finally, LeT assumes 518 that responses are emitted at a constant rate if the current active state has 519 associative strength w(n) greater than a threshold  $\theta$ . The fact that responding 520 depends on the associative strength of the current state, and that this strength 521 only changes with US associations, prevents LeT from accounting for changes 522 in timing that are not related to US occurrence. For example, there is evidence 523 that animals learn the timing of a preexposed CS (Bonardi et al., 2016) and are 524 sensitive to changes in timing during extinction (Guilhardi and Church, 2006), 525 two situations that do not involve the occurrence of a US. 526

Modular Theory is another primarily timing theory that can also deal with some aspects of conditioning. It treats the onset of a stimulus as signalling a time expectation to reinforcement. Its time representation T is, like LeT, an accumulator that increases linearly with time t, T = ct, where c is a constant. When reinforcement is delivered the current reading from the accumulator is stored in what is called *pattern memory*. Pattern memory is updated at each trial n according to

$$m(n) = m(n-1) + \alpha(T^* - m(n-1))$$
(18)

where  $\alpha$  is a learning rate and  $T^*$  is reinforcement time. Equation (18) may be 534 contrasted to (6) from RWDDM. The main difference is that pattern memory 535 in MoT stores a moving exponential average of intervals, whilst the slope in 536 RWDDM stores a moving exponential harmonic average of intervals. However, 537 both models are similar in that they can potentially time the occurrence of any 538 event, not only rewards. MoT's pattern memory and RWDDM's slope can be 539 made, for example, to adapt to mark the end of stimuli that are not necessarily 540 paired with a reward. 541

A stochastic threshold b is used to mark response initiation. The threshold 542 distribution is set so as to yield timescale invariance of the response curve. 543 Its mean, B, is a fixed proportion of the value in pattern memory, B = km(n), 544 where k is the proportionality constant, and its standard deviation is  $\gamma B$ , where 545  $\gamma$  is the coefficient of variation of B. Hence, the coefficient of variation of 546 the threshold, i.e. of response initiation, is constant for all intervals, which 547 is the timescale invariance of the response curve. RWDDM derives timescale 548 invariance of response curve from noise in the accumulator (equation (2), not 549 from the threshold. 550

551

This account of time from MoT is an instantiation of Scalar Expectancy

Theory, arguably one of the most successful timing models to date. Being a purely timing theory, SET does not address associative learning directly, so it does not have a rule for changes in association between stimuli. MoT bridges this gap by adding a rule to update what is termed *strength memory*, w(n). Strength memory holds the associative strength between stimulus and reinforcement. The rule consists of a linear operator:

$$\Delta w(n) = \begin{cases} \beta_e(0 - w(n-1)) & \text{if US is absent,} \\ \beta_r(1 - w(n-1)) & \text{if US is present,} \end{cases}$$
(19)

with  $\beta$  a constant that can determine different rates of update for acquisition ( $\beta_r$ ) and extinction ( $\beta_e$ ). Equation (19) may be compared with (9). Note that, unlike RWDDM, equation (19) does not contain the summation term from RW based rules.

MoT also includes a rule for response rate that is more realistic than RWDDM's given by (10). It is partly derived from an empirical analysis of real-time responding in animals. We refer the interested reader to Guilhardi et al. (2007) for a fuller description. We will only mention here that MoT generates a two-state response pattern, low and high. The transition between states is determined by the crossing of threshold B, and the high state is proportional to strength memory w(n).

Other theories exist which are similar in scope to CSC-TD, MS-TD, LeT and MoT. Two notable examples are the Componential version of the Sometimes Opponent Process model (C-SOP, Brandon et al., 2003) and the Adaptive Resonance Theory - Spectral Timing Model (ART-STM Grossberg and Schmajuk, 1989). C-SOP builds a CS representation based on two sets of elements, or components, one that includes elements activated as a function of time and another whose elements are randomly activated. Associative strength for each element is

updated using the standard trial-based RW rule. Simulations in Brandon et al. 576 (2003) have demonstrated that C-SOP can produce some degree of timescale 577 invariance. ART-STM is a neural net with an input layer and one hidden layer, 578 which allows it to explain nonlinear conditioning phenomena (such as negative 579 pattern) that a single-layer RW neural net cannot. It employs a CS represen-580 tation that is very similar to the microstimuli used in MS-TD, so it also shows 581 a degree of timescale invariance. Other theories could be mentioned (for two 582 influential examples see Buhusi and Schmajuk, 1999; McLaren and Mackintosh, 583 2000,0) but we will limit the analysis to CSC-TD, MS-TD, LeT and MoT for 584 two reasons: a) these four models collectively embody most of the conditioning 585 and timing mechanisms used in modelling these areas, and b) our goal here is 586 not to provide a comprehensive review, but rather focus on the mechanisms that 587 are shared by our proposed model and the others. 588

Table 1 summarizes the main mechanisms/features of the models described 589 above. In terms of the type of time representation, it may be observed that 590 the models fall roughly into two categories: (a) those that employ a chain of 591 units or states activated sequentially (CSC-TD, MS-TD, LeT), and (b) those 592 that employ an accumulator (MoT and RWDDM). Those in category (b) may 593 be considered more economical both computationally and biologically, as they 594 don't require a number of units that increase with time. In terms of what 595 the representations can time, two categories may be discerned: (a) those that 596 time only rewards (CSC-TD, MS-TD and LeT), and (b) those that can time 597 any stimuli (MoT and RWDDM). Models in category (b) have more flexibil-598 ity to create a temporal map involving all stimuli present, including those not 599 signalling reward. In terms of timescale invariance, the models are basically di-600 vided between those that can account for it (MS-TD, LeT, MoT and RWDDM) 601 and the one that cannot (CSC-TD). Finally, in terms of the type of associative 602

model	type of time representation	what it can time	timescale in- variant	associative learning rule
CSC-TD	units/states, one per time step	only rewards	no	TD/RW, cue competition
MS-TD	units/states, fewer than one per time step	only rewards	approximately	TD/RW, cue competition
LeT	units/states, one per time step	only rewards	yes	linear operator, no cue competi- tion
MoT	linear accumu- lator	any stimuli, not only re- wards	yes	linear operator, no cue competi- tion
RWDDM	noisy linear ac- cumulator	any stimuli, not only re- wards	yes	RW, cue competition

Table 1: Summary of the main features of the models.

learning rule used, models are divided between those that use a RW-type rule
(CSC-TD, MS-TD, RWDDM) and those that use the linear operator (LeT and
MoT). The ones that use RW are wider in scope, being able to account for
cue-competition phenomena, which form the core of classical conditioning.

The main innovation of RWDDM over its predecessors is the combination of 607 a noisy linear accumulator for timing with the RW rule for associative learning. 608 As table 1 shows, linear accumulator theories are the only ones in our sample 609 of the models that can fully account for timescale invariance. But because 610 they rely on the linear operator rule, they cannot account for cue-competition 611 and other compound stimuli phenomena in conditioning. Therefore RWDDM 612 extends the application of the linear accumulator to compound stimuli, covering 613 a wider range of conditioning phenomena. 614

In summary, the model we propose is, to the best of our knowledge, the only one that unites the flexibility, computational economy and timescale invariance of the linear accumulator as a time representation, to the RW associative

RWDDM feature	phenomenon for which it can ac- count
independent update rules for time and associative strength	faster reacquisition, time change in extinction, latent inhibition and timing
RW rule for associative strength	blocking with different durations, time specificity of conditioned inhi- bition
intertrial variability in time estimation asymptote of associative strength set by time	compound peak procedure ISI effect, mixed FI
a memory that learns the rate of reinforcement	VI and FI, temporal averaging

Table 2: Model features and the experimental findings they can explain.

learning rule, which accounts for many more conditioning phenomena than the
linear operator. In the next section we evaluate the models against a number
of phenomena in conditioning and timing.

### 621 **3** Results

The long history of experimental work in classical conditioning has allowed the discovery of a rich variety of phenomena–a recent review (Alonso and Schmajuk, 2012) has catalogued approximately 87. This forces theorists to be selective when deciding which phenomena to simulate when presenting a new model. We searched the literature for phenomena that could test each feature of the model. Table 2 lists the main RWDDM features, together with the corresponding phenomena found in the literature that can test each.

Table 3 contains the design for each simulation performed with the model. The model parameters used in all simulations were kept almost constant but in some cases a few adjustments were found necessary to obtain a better agreement between model and data. We report their values in each simulation below. The time-step was the same for all simulations:  $\Delta t = 10$  msec. Simulations were performed using MATLAB version R2016b. The code to generate the figures in each result section is available as supplementary material.

#### <sup>636</sup> 3.1 Faster reacquisition

A conditioned response emerges gradually over the course of several trials where 637 the CS signals the arrival of a US. If a measure of CR strength (such as rate or 638 magnitude) is plotted against the number of trials, the shape and rate of this 639 acquisition curve will depend largely on the CR and organism, but it usually 640 follows a negatively accelerated curve (Pavlov, 1927; Kehoe and Macrae, 2002). 641 Pavlov (1927) believed timing of the CR would emerge only later in acquisition, 642 through a process he described as *inhibition of delay* whereby the initial part 643 of the CS would become inhibitory. Recent and more detailed analyses suggest 644 that an estimate for the time to reinforcement is acquired very early in training, 645 possibly even after one or two trials, although the expression of such estimation 646 may not be observable until later in training (Holland, 2000; Ohyama and Mauk, 647 2001; Balsam et al., 2002; Drew et al., 2005). 648

If the CS no longer signals reinforcement, CR strength gradually decreases over the course of these extinction trials, until it finally disappears. If the CS is made to signal the US again, the CR returns, a process that is called reacquisition. It is a consistent finding that reacquisition is faster than acquisition (Ricker and Bouton, 1996; Guilhardi et al., 2007; Kehoe and Macrae, 2002, p. 185).

Learning is loosely defined as an enduring change in behaviour as a result of experience. Acquisition of a CR is the most basic demonstration that classical conditioning is a form of learning. As such, all classical conditioning models provide an account of it.

#### 659 3.1.1 Simulations

Figure 3 (top left panel) shows a plot of RWDDM's associative strength as given by equation (9), in a simulation of acquisition and extinction. Acquisition consisted of 80 presentations of a 5-sec CS followed by reinforcement, after which there were 100 extinction trials where H was set to zero. The simulations match with experimental data from acquisition and extinction (bottom left panel of figure 3). The simulated acquisition curve asymptotes around the theoretical value given by setting  $\Delta V(n) = 0$  in equation (9) and solving for V, yielding

$$V_{\infty} = \frac{HA_{\infty}}{x(\Psi_{t^*})\Psi(t^*)},\tag{20}$$

which in this particular case is  $V_{\infty} \approx 1$ , since H = 5,  $A_{\infty} \approx 1/5$ ,  $\Psi_{t^*} = \Psi(t^*) \approx$ 667 1,  $x(\Psi_{t^*}) \approx 1$ , where  $t^*$  is the time of reinforcement. Because  $\Psi(t^*)$  is a random 668 variable,  $x(\Psi_{t^*})$  and  $V_{\infty}$  are also random variables and their values are reported 669 as approximations to their expected values (but not the actual expected values). 670 Figure 3 (top middle panel) shows the adaptation of timer slope A given by 671 equation (6). This equation precludes the initial value of A from being zero, 672 so we set it to the very low value of  $A(1) = 10^{-6}$ . We also set the threshold 673  $\theta = 1$ , which by equation (6) means that  $A_i(n)$  encodes the exponential moving 674 average of the rate of reinforcement signalled by  $CS_i$ . Or, equivalently,  $1/A_i(n)$ 675 encodes the moving harmonic average of the intervals since last reinforcement 676 during  $CS_i$ . In this simulation, since there is only one US which is delivered 677 always at the same time at CS offset (5000 msec), A converges to  $A_{\infty} = 1/5000$ . 678 Note that the value of A does not decline after extinction begins at trial 80. It 679 continues to be updated since the stimulus is still present, even if its presence 680 no longer signals reinforcement. 681

The top right panel of figure 3 shows the acquisition and reacquisition curves using RWDDM. Reacquisition produced by the model is evidently faster than



Figure 3: Acquisition and reacquisition. Top left: simulated associative strength V in acquisition and extinction. Top middle: adaptation of RWDDM slope A. CR extinction began at trial 80 but has no effect on the RWDDM slope. Top right panel: simulated V curves in acquisition and reacquisition. Bottom left panel: response strength data from an experiment in acquisition and extinction, redrawn from figure 1 in Ricker and Bouton (1996). Bottom right panel: data from an experiment in acquisition, redrawn from the top panel of figure 3 in Ricker and Bouton (1996). Model parameters: m = 0.15,  $\theta = 1$ ,  $\sigma = 0.3$ ,  $\alpha_t = 0.1$ ,  $\alpha_V = 0.1$ , H = 4 in acquisition and H = 0 in extinction.

the simulated acquisition, but not as fast as the reacquisition seen in the data on the bottom left of figure 3.

#### 686 3.1.2 Discussion

In RWDDM acquisition and extinction of associative strength follow from the same mechanism as RW. The only difference is the noisy stimulus representation  $x(\Psi_{t^*})$ , which induces noise into the acquisition curve. Changes in associative strength and timing are treated independently. In particular, the memory for time encoded by the slope A is not affected by extinction. This leads to a faster reacquisition following extinction. This is because RWDDM's time-adaptive <sup>693</sup> CS representation  $x(\Psi_{t^*})$  reaches its maximum activation value right from the <sup>694</sup> beginning of reacquisition, since the timer slope A is already tuned to the current <sup>695</sup> CS duration (see equation (8)).

Modular theory (Guilhardi et al., 2007) is another model that treats timing and associative strength separately. It postulates two memories, one for the pattern of reinforcement and another for the strength of the association between CS and US. The pattern memory stores an exponential moving average of the intervals to reinforcement which, like RWDDM, does not change with extinction. However, its strength memory w(n) is updated according to the linear operator rule,

$$w(n+1) = w(n) + \beta(\lambda - w(n))$$
(21)

which, unlike RWDDM, does not include a term for a time-adaptive CS representation. Thus, the way MoT accounts for rapid reacquisition is by using different learning rates  $\beta$  for acquisition and reacquisition. The same strategy may be employed with the TD and LeT models.

In summary, RWDDM explains reacquisition as the persistence of a memory
 for time, whilst TD, LeT and MoT explain it as a permanent change in the
 learning rate for associative strength.

#### 710 3.2 Time change in extinction

When a previously conditioned stimulus is no longer followed by reinforcement, the conditioned response gradually decreases. An important theoretical question for hybrid timing/conditioning models concerns what happens to the timing of responses in extinction. Using the peak procedure Ohyama et al. (1999) found that although the maximum (peak) response rate decreased in extinction, peak time and sensitivity (measured by the coefficient of variation) remained virtu-

ally unchanged. Drew et al. (2004) investigated the behaviour on extinction by 717 changing CS duration between acquisition and extinction. Groups where the 718 CS changed to a shorter or longer duration were compared to another where 719 the duration did not change. They found that CS duration had little effect on 720 the rate of extinction, with all groups taking about the same number of tri-721 als to achieve CR extinction. However, when the CS used in extinction was 722 considerably longer (4 times) than the one acquired, extinction was facilitated. 723 Guilhardi and Church (2006) performed a similar experiment (experiment 2) 724 and observed that when stimulus duration is changed from acquisition to ex-725 tinction, the pattern of responding during extinction gradually shifts to the new 726 duration over extinction trials. Following the same procedure, Drew et al. (2017) 727 also used partial reinforcement to slow down the rate of acquisition, and thus 728 observe if response patterns really do shift gradually to the new duration. They 729 confirmed that when CS duration was increased from acquisition to extinction, 730 the within-trial response peak shifted gradually to the right over the course of 731 extinction. When the CS was shortened, the results were not conclusive. Also, 732 when CS duration was changed from training to extinction, the speed of extinc-733 tion increased, but this appeared to be explained at least in part by the shifting 734 of response patterns. 735

In summary: a) peak timing and CV are not altered in extinction when using 736 a peak procedure, b) changing the CS duration from training to extinction causes 737 the within-trial response peak to shift to the new duration, and c) changing the 738 CS duration in extinction can speed up extinction, but this may be due to the 739 shifting of the response peak and not to changes in associative strength. These 740 results pose a challenge to the models analysed here. Out of CSC-TD, MS-TD, 741 LeT and MoT, only MoT has a mechanism that would allow it to account for 742 time change in extinction. 743

#### 744 3.2.1 Simulations

RWDDM provides an account for these findings as follows. In the case of the peak procedure, the occurrence of the longer peak trials may be considered too infrequent to cause a shift to the longer time. In this case, equation (6) is not applied in peak trials so RWDDM predicts that both slope *A* and CV will remain unaltered in extinction. In the case of a permanent change in CS duration from acquisition to extinction, the slope update rule is applied and the response peak will shift gradually to the new duration.

We have simulated RWDDM in two extinction conditions, one where the CS presented in extinction was longer than the one acquired (20 sec to 40 sec, short-long) and another where the extinction CS was shorter than the acquired CS (20 sec to 10 sec, long-short). Figure 4 summarizes the main results.

The panels on the left column show response strength during a trial in condi-756 tions short-long (top) and long-short (bottom). In the early stages of extinction 757 (early) the response curves peak around the time of US arrival in acquisition 758 (20 sec). This is more evident in the condition short-long (top left) because in 759 the other condition (bottom left) the trial ends 10 seconds before the peak at 20 760 seconds occurs. Had the stimulus remained on for a full 20 seconds, the response 761 curve in the early stages of long-short would have continued to increase until the 762 20 second mark. In middle and late extinction the response peak slowly shifts 763 to the new duration in both conditions, and their heights decrease. Compare 764 the simulated curves in the left column of figure 4 to the actual experimental 765 data in the right column. The panels on the middle row of figure 4 show the 766 adaptation of time estimate 1/A in conditions short-long (top) and long-short 767 (bottom). They demonstrate that RWDDM adapts exactly to time change in 768 extinction. 769

770

To investigate if the rate of acquisition changes with CS duration, we have



Figure 4: Time change in extinction. Left column: simulated response strength averaged over trials in extinction short-long (top) and long-short (bottom). Middle column: time estimate adaptation of the model during extinction short-long (top) and long-short (bottom). Right column: experimental data from an experiment where the CS duration changed from 12-sec in acquisition to either 24-sec (top) or 6-sec (bottom) in extinction. Data plots redrawn from figure 10 in Drew et al. (2017). Model parameters: m = 0.25,  $\theta = 1$ ,  $\sigma = 0.35$ ,  $\alpha_t = 0.08$ ,  $\alpha_V = 0.09$ , H = 30.

plotted the extinction curves for each CS duration in the left panel of figure 771 5. Decreasing CS duration from acquisition to extinction slightly facilitates 772 extinction, but increasing CS duration markedly delays extinction. However, 773 these are only the V values, a theoretical construct that accounts for the as-774 sociative strength of the stimulus as a whole. Actual behaviour measurements 775 of extinction are based on how much response frequency changes from trial to 776 trial. But response frequency also changes within the trial. As pointed out by 777 Drew et al. (2017), the value obtained for the rate of extinction may be affected 778 by which portion of the CS was measured. To analyse this, Drew et al. (2017) 779 measured response frequency only during the first 6-sec (half the duration of 780 the CS in acquisition) of each CS duration in extinction. We have followed the 781
r82 same procedure and the results can be seen on the middle panel of figure 5.
r83 They show a marked delay on extinction when the CS duration was shortened,
r84 but not when it was lengthened. Compare these curves with the actual data
r85 analysed by Drew et al. (2017) and displayed in the rightmost panel of figure 5.
r86 The simulations conflict in part with the same analysis in Drew et al. (2017),
r87 which showed no delay on extinction, only facilitation in the case of extending
r88 CS duration.



Figure 5: Extinction curves. Left panel: model V values for each CS duration in extinction. Middle panel: simulated CR values calculated only for the first 10 seconds of the CS. Each data point is calculated by summing the output of equation (10) over the first 10 sec of each trial, then averaging these trial values two by two, and dividing by 100 to rescale. Right panel: actual CR data for the first 6 sec of the CS in extinction, redrawn from figure 8 (C) in Drew et al. (2017)

#### 789 **3.2.2** Discussion

RWDDM predicts that a change in CS duration from acquisition to extinction 790 will always cause a rescaling of the response curves in extinction. This is largely 791 in agreement with the data. However, RWDDM seems to predict a degree of 792 delay on extinction, whilst the data seems to point to a facilitation of extinction 793 when the CS changes duration. When only the first half of the CS response 794 curves are analysed, the data suggests that extending CS duration in extinction 795 can speed up extinction, whilst RWDDM predicts that shortening CS duration 796 will delay extinction. 797

RWDDM's prediction for a delay in extinction following a change in CS duration is due to the shifting of the response curve. At the beginning of extinction, a trial ends either before the CS representation has reached its peak (CS shortening) or after its peak (CS lengthening). This makes equation (9) update with a small value for  $x(\Psi)$ , resulting in a smaller update than with the higher  $x(\Psi)$  value of the unchanged CS.

As mentioned above, time change in extinction is a difficult phenomenon for 804 the current models to explain. CSC-TD does not have a mechanism to change 805 the peak of responding when a US is not present. Neither does MS-TD or LeT. 806 These models assume that extinction can only weaken existing links between CS 807 and US representations. Because in these models timing usually depends on the 808 sequential activation of these links, changing the CS duration in extinction would 809 not alter the timing but only the magnitude of responding. RWDDM explains 810 time change in extinction because its rule for time adaptation is independent of a 811 change in associative strength. Thus, when the duration changes in extinction, 812 RWDDM's accumulator slope tracks this change, whilst associative strength 813 decays as a function of US absence. Regarding the extinction facilitation caused 814 by a change in CS duration, none of the models analysed here currently have a 815 mechanism to explain this either. 816

It would be possible to allow the average rate of state transition in LeT to vary as a function of CS duration, which would cause timing to adapt to the new time in extinction. However, in its latest formulation (Machado et al., 2009) LeT relies on a fixed average rate of state transition to explain timescale invariance. Thus, if the rate is made to change as a function of CS duration, this would break timescale invariance.

As for MS-TD, one interesting modification that would likely allow it to explain time change in extinction is to make the microstimuli themselves timeadaptive. Like RWDDM's time-adaptive CS representation, the microstimuli
could be made to 'stretch' or 'compress' when stimulus duration shortens or
lengthens.

Modular Theory is likely to account for time change in extinction, since its pattern memory for time could be made to update even in extinction. That would shift the response pattern to the new time whilst strength memory, which depends only on US presentation, would decay.

# <sup>832</sup> 3.3 Latent inhibition and timing

When a subject is exposed to repeated and non-reinforced presentations of a 833 stimulus it has never encountered before, this procedure is called preexposure. 834 If reinforcement is subsequently paired with the preexposed CS, the initial rate 835 of CR acquisition is usually lower compared to acquisition to a nonpreexposed 836 stimulus, a phenomenon called latent inhibition (Lubow and Moore, 1959). The 837 asymptotic level of conditioning, however, is not normally affected by preexpo-838 sure (Lubow, 1989). Latent inhibition is an important representative of a class 839 of phenomena involving latent effects. Collectively, these phenomena demon-840 strate that something is learned about the stimulus even when it does not sig-841 nal reinforcement. Therefore, latent inhibition cannot be accounted by the 842 Rescorla-Wagner model, since the theory only applies when there are changes 843 in associative strength. 844

A question relevant for real-time conditioning models is what happens to timing when a preexposed stimulus is conditioned. To answer this question, Bonardi et al. (2016) used CSs of variable and fixed durations (the variable duration CS had the same mean as the duration of the fixed CS) to vary the temporal conditions between preexposure and conditioning phases. Latent inhibition was observed even when the temporal information from the two phases was different. Crucially, timing, as measured by the response gradient within
a trial, appeared to improve in the preexposed CS even when the temporal
information was different between the two phases.

As alluded to above, latent inhibition cannot be accounted by the associative 854 learning update rule used in RWDDM, the Rescorla-Wagner. However, we show 855 here that RWDDM is compatible with the Pearce-Hall rule (Pearce and Hall, 856 1980; Pearce et al., 1982), one of the most widely used models for explaining 857 latent inhibition and other latent learning effects. We demonstrate that this 858 modification maintains the basic framework of the RWDDM, and that it can 859 account for latent inhibition and improved timing with preexposure. None of 860 the other models analysed here can account for latent inhibition without mod-861 ifications. Improved timing with preexposure could be accounted by Modular 862 Theory, but not by the the current version of the other models. 863

### 864 3.3.1 Simulations

The Pearce-Hall model is basically a rule for adapting the learning rate  $\alpha_V$  based on the error  $\delta$  between the predicted US outcome and the actual US outcome. It was originally formulated by Pearce and Hall (1980) and updated by Pearce et al. (1982). We have maintained equation (9) for associative strength, but changed  $\alpha_V$  on every trial *n* according to

$$\alpha_V(n+1) = \alpha_V(n) + \gamma(|\delta| - \alpha_V(n)), \tag{22}$$

$$\delta = \left(\frac{HA(n)}{\Psi(t^*)} - V(n)x(\Psi)\right) \tag{23}$$

where  $0 < \gamma < 1$  is a parameter that sets the rate of learning rate adaptation. Equation (22) is basically the Pearce-Hall rule, except that instead of using 1 as the asymptote of learning we use  $\frac{HA(n)}{\Psi(t^*)}$ .

We simulated latent inhibition with a 5-sec CS. Preexposure consisted of

<sup>874</sup> 80 trials of the CS without reinforcement (H = 0). The preexposed CS was then reinforced for 250 trials. Figure 6 (top left panel) compares the acquisition curves for the preexposed CS and a control CS in the reinforced trials. The preexposed CS acquisition curve increases at a lower rate than the control CS, the latent inhibition effect (see data from a corresponding experiment at the bottom left panel of figure 6).



Figure 6: Latent Inhibition. Top row: simulated associative strength in latent inhibition (left), simulated CR averaged over the first 30 trials of conditioning phase (middle), and simulated CR averaged over the last 30 trials of conditioning phase (right). Bottom row: acquisition curves from an actual experiment in latent inhibition (left), and response rate data during the CS (right). Data plots redrawn from figures 1 and 2 respectively in Bonardi et al. (2016). Model parameters:  $\alpha_t = 0.1$ ,  $\alpha_V = 0.08$ ,  $\mu = 1$ ,  $\sigma = [0.6 - 0.35]$ , m = 0.2, H = 4,  $\alpha_{PH} = 0.4$ ,  $\gamma = 0.03$ .

Improved timing with preexposure follows directly from the fact that RWDDM adapts its accumulator slope A to the CS duration during preexposure. However, our choice of a Gaussian for stimulus representation does not allow for this change to become visible. Bonardi et al. (2016) demonstrated improved timing by showing that the slope of the response curve from the preexposed CS was

higher in the first few trials of acquisition than the one from the control CS 885 (see bottom right panel of figure 6). In general, animal response curves tend 886 to be quite flat during the beginning of acquisition. There is evidence that the 887 response curves appear to change from negatively accelerated to a sigmoidal 888 shape over the course of training (see figure 1 in Meck and Church, 1984, for 889 an example). This means that in the early stages of acquisition, within-trial 890 response frequency increases very early in the trial and then stays at a constant 891 level until the end. As training progresses, the increase in frequency moves 892 slowly to the right, giving rise to the sigmoidal shape that peaks just before 893 the end of the trial. In these cases a higher slope of the response curve would 894 indicate improved timing. But in our model the curves are sigmoidal from start 895 of acquisition, so they will always peak at the end of the trial, even if the timer 896 slope has not adapted to the interval yet, as is the case with a novel stimulus. 897 Therefore, during the acquisition phase of latent inhibition, RWDDM predicts 898 that only the peaks of the response curves will gradually increase over the tri-899 als. Because of the learning decrement caused by preexposure, the peak of the 900 control CS will increase faster than the preexposed CS, as the top middle panel 901 of figure 6 demonstrates. The response curve of the control CS will have a 902 higher slope than the preexposed CS, even though the preexposed CS's timer 903 rate has been adapted to its duration. Hence, the improved timing found in the 904 data is explained by adaptation of RWDDM's timer slope, but RWDDM's CS 905 representation cannot make this visible. 906

We have tried adding an adaptable  $\sigma$  in equation (8) so as to decrease the width of the gaussian curve gradually over trials. We chose a simple linear operator rule to adapt the Gaussian width:

$$\sigma(n+1) = \sigma(n) + \alpha_{\sigma}(0.35 - \sigma(n)), \qquad (24)$$

and set  $\sigma(1) = 0.6$  and  $\alpha_{\sigma} = 0.025$ .

Figure 6 (top middle panel) shows response strength of control and preexposed CSs averaged over the first 30 trials of the conditioning phase. The preexposed CS already shows a clear sigmoidal shape, whilst the control is slightly wider and linear. But the effect is too small to be able to account for the one seen in the data from Bonardi et al. (2016). Towards the end of the conditioning phase the two curves converge (figure 6, top right panel).

#### 917 3.3.2 Discussion

The simulations show that the model can account for latent inhibition ade-918 quately if the Pearce-Hall rule is used (in which case the model would be more 919 appropriately named PHDDM). The PH rule adapts the learning rate  $\alpha_V$  based 920 on the level of associative learning between stimulus and reward. When the 921 subject encounters a novel stimulus, it is assumed that  $\alpha_V$  has some non-zero 922 starting value  $\alpha_V^{\text{novel}}$ , which allows learning in equation (9) to take place. If this 923 novel stimulus does not signal reward, as is the case in the preexposure phase 924 of latent inhibition,  $\sigma = 0$  and equation (22) will simply decay the value of the 925 learning rate across trials until it reaches zero. If at this point the stimulus 926 begins to be followed by reward,  $\sigma > 0$  and equation (22) will begin to raise 927 the value of the learning rate, which in turn will allow equation (9) to begin 928 increasing the value of V. Since the increase in the value of the learning rate 929 is gradual, determined by the rate  $\gamma$ , there will be a number of trials in the 930 beginning of the conditioning phase where  $\alpha_V < \alpha_V^{\text{novel}}$ , which leads to the ini-931 tial impairment in the learning curve when compared to the learning curve of a 932 non-preexposed CS, as seen in the top left panel of figure 6. 933

The separate rule for time adaptation allows the model to account for improved timing after preexposure, but the model cannot make this effect visible even if we allow for Gaussian width adaptation. In view of this it seems more likely that a two-state CS representation may be a better solution. As mentioned above, figure 1 in Meck and Church (1984) suggests that during the
initial stages of training a CS representation may be modelled by the following
leaky integrator

$$x_i(t+1) = x_i(t) + \frac{1}{\tau}(I_i - x_i(t))$$
(25)

where  $I_i$  is the indicator function marking the presence of  $CS_i$ , and  $\tau$  a time 941 constant. In the latter stages of training, when timing is expressed, the organism 942 switches to the Gaussian representation given by equation (8). When the switch 943 between representations is made and how abruptly remains to be investigated. 944 Latent inhibition cannot be accounted by any of the other models analysed 945 here without modifications. Also, models that rely on the US for time adap-946 tation, like CSC-TD, MS-TD and LeT, cannot account for improved timing by 947 preexposure. Modular Theory is the only one that can time any stimulus like 948 RWDDM, so it could account for the improved timing. But it would also need a 949 modification like (22) to adapt its learning rate to account for latent inhibition. 950

# **3.4** Blocking with different durations

Arguably, the most important compound conditioning phenomenon is blocking. 952 It is part of a class of cue competition and compound phenomena discovered in 953 the late 1960s which challenged the view that conditioning was driven by the 954 pairing, or contiguity, of CS-US. These results suggested that conditioning with 955 compound stimuli was influenced by the reinforcement histories of the elements 956 forming the compound (Rescorla, 1988; Gallistel and Gibbon, 2001). This led 957 to the development of a new generation of models that could account for those 958 findings (Rescorla and Wagner, 1972; Mackintosh, 1975; Pearce and Hall, 1980). 959 The rule we use, the Rescorla-Wagner, provides an explanation for blocking that 960

 $_{961}$  is based on the summation term in equation (1).

In a blocking procedure a CS is first paired with a US in phase 1 of training. 962 During phase 2 a novel CS is presented in compound with phase 1 CS and paired 963 with the US for just a few trials. Subsequently, when tested alone the novel CS 964 elicits less responding than if it had been trained in compound with another 965 novel stimulus (Kamin, 1968). The previously reinforced CS is said to block 966 the novel CS. The temporal information encoded by each CS has an effect on 967 the amount of blocking observed. Schreurs and Westbrook (1982) varied the ISI 968 in the pre-training and compound phases, and observed less blocking when the 969 durations were different in both phases than when they were the same. Barnet 970 et al. (1993) performed a similar experiment but with forward and simultaneous 971 conditioning varying between phases, and also found that blocking was stronger 972 when blocked and blocking CSs had the same temporal history. Jennings and 973 Kirkpatrick (2006) used compounds where the elements had different durations. 974 They observed that a long blocking CS could block a co-terminating short Cs, 975 but a short blocking CS failed to block a co-terminating long CS (see rows 1 and 976 3 in figure 7). Amundson and Miller (2008) performed four blocking experiments 977 using trace conditioning. In two of them the blocking CS trace duration changed 978 between phases, and blocking was not observed. In the other two experiments 979 the trace duration was held fixed between phases, and the blocking and blocked 980 CSs were presented serially and not in a compound (see rows 2 and 4 of figure 981 7). Blocking was observed when the blocking CS followed the blocked CS, but 982 not in the reverse condition. 983

The studies reviewed above appear to show that changing the ISI of the blocking CS between phases may attenuate blocking. Another finding is the apparent asymmetry of blocking when the ISI of the blocking CS is kept constant between phases. Rows 1 and 2 of figure 7 suggest that a long blocking ISI can



Figure 7: Experimental designs from two blocking experiments. CS X was blocked (B) in rows 1 and 2, and not blocked (NB) in rows 3 and 4. Blue bar indicates US presence.

<sup>988</sup> block a short blocked ISI. Rows 3 and 4 suggest that a short blocking ISI does
<sup>989</sup> not block a long blocked ISI.

As mentioned above, RWDDM can account for blocking because it uses the 990 RW rule. The summation term in equation (1) formalizes the widely held view 991 that a given US can only confer a limited amount of associative strength which 992 CSs must compete for. Different theories exist that take other approaches to 993 blocking (see for example Mackintosh, 1975; Harris, 2006; Stout and Miller, 994 2007) but among the ones analysed here (for their ability to handle timing also) 995 only CSC-TD and MS-TD are equipped to deal with it. We show next that 996 RWDDM can account for the blocking of a short CS by a long CS, and that 997 by making the reasonable assumption of second-order conditioning it can also 998 account for the lack of blocking of a long CS by a short CS. CSC-TD and MS-TD 999 are also capable of providing an account of both blocking conditions. 1000

## 1001 3.4.1 Simulations

Because RWDDM is based on the RW rule, it produces virtually the same results 1002 as the latter when the CSs have the same duration. Our interest here is to test 1003 whether it can reproduce the finding that a long CS can block a shorter CS but 1004 a shorter CS does not block a longer one. We performed a simulation following 1005 the design in rows 1 and 3 of figure 7. In the first phase a CSA (blocking 1006 CS) of duration either 10 or 15 seconds was followed by reinforcement until its 1007 associative strength V reached asymptote. In phase 2 CSA was joined with a 1008 CSX (blocked CS), of either 15 or 10 seconds, in a coterminating compound and 1009 followed by US. The top left panel of figure 8 shows the acquisition of associative 1010 strength for CSX and its control during phase 2 for the condition CSA-15sec 1011 and CSX-10sec. A considerable amount of blocking is observed, matching with 1012 the data (bottom left panel). 1013

The top right panel of figure 8 shows the results for condition CSA-10sec and CSX-15sec. In this condition the model diverges considerably from the data (bottom right panel) and predicts that CSX should actually become inhibitory.

### 1017 **3.4.2 Discussion**

The blocking and inhibition seen in figure 8 is a result of a discrepancy in the asymptote of learning between the CSs. After phase 1, CSA has associative strength  $V_A \approx HA_A$ . During phase 2, CSX's associative strength changes according to:

$$\Delta V_X \approx \alpha (HA_X - (V_A + V_X))$$
$$= \alpha (HA_X - (HA_A + V_X))$$
$$= \alpha (H(A_X - A_A) - V_X)$$



Figure 8: Blocking with different durations. Left column: simulation (top) with a 15 sec blocking CS and 10 sec blocked CS, and animal data (bottom) from an experiment with the same design. Right column: simulation (top) with a 10 sec blocking CS and 15 sec blocked CS, and animal data (bottom) from an experiment with the same design. Data panels redrawn from the top right panel in figure 5 in Jennings and Kirkpatrick (2006). Model parameters:  $\alpha_t = 0.2$ ,  $\alpha_V = 0.1$ ,  $\mu = 1$ ,  $\sigma = 0.35$ , m = 0.2, H = 10.

1022 and since  $(A_X - A_A) < 0, V_X$  becomes negative.

However, it could be argued that the short CSA becomes a secondary reinforcer which is signalled by the onset of the long CSX. In this case, the onset of CSX would serve as the time marker for the onset of CSA, and not for the onset of US. Hence, during the first 5 seconds of CSX responding would be under the control of this 5-sec stimulus representation which would not overlap, thus not compete, with CSA's later representation. It would follow from this account that no blocking would be observed, and that responding during test phase with
CSX would peak at the 5-sec mark. This is a testable prediction that, if shown
to be the case, could validate RWDDM's account.

Also note that the time-dependent associative strength asymptote assumed by RWDDM implies that learning during a compound where the elements are of different durations is not stable. In particular, if CSA and CSX are the two elements of the compound phase of blocking, their associative strengths are updated by RWDDM as

$$\Delta V_A = \alpha_V (HA_A - (V_A + V_B))$$
$$\Delta V_B = \alpha_V (HA_B - (V_A + V_B)),$$

<sup>1037</sup> which in the steady state form an inconsistent system of linear equations,

$$V_A + V_B = HA_A$$
$$V_A + V_B = HA_B.$$

Since the compound phase of blocking only lasts for a few trials, RWDDM 1038 could produce the blocking seen on the left panel of figure 8. But if training 1039 with the compound was carried out for longer, the V values would grow without 1040 bound. However, there is evidence that in compounds formed by elements with 1041 asynchronous onsets, like in the compound phase of the blocking experiments 1042 here, the shorter stimulus comes to control CR timing and there is no summation 1043 of associative strengths (Fairhurst et al., 2003). Hence, it appears that with 1044 compounded asynchronous CSs, the shorter CS, more proximal relative to the 1045 US, comes to dominate and a summation rule like RW would not be applicable 1046 beyond the first few trials of training. 1047

<sup>1048</sup> A model that is well placed to explain these results is CSC-TD. A long

blocking CS will completely overlap a short blocked CS, blocking all units in the 1049 blocked CS. But in the case of a short blocking CS, there will be free units in the 1050 beginning of the blocked CS which will acquire associative strength, attenuating 1051 blocking. Given its similarity, MS-TD would likely produce comparable results. 1052 MoT and Let would not be able to account for any type of blocking given their 1053 current choice of rule for associative strength. Unlike RWDDM and the TD 1054 models, they both rely on the linear operator rule, which antedates the transition 1055 to the rules that sum associative strengths in the compounds as mentioned 1056 previously. MoT and LeT would need, at the very least, to replace the linear 1057 operator by the RW or other equivalent rule to be able to account for blocking 1058 and other compound phenomena. 1059

## <sup>1060</sup> 3.5 Time specificity of conditioned inhibition

Learning occurs not only when a CS signals the occurrence of a US, but also when a CS signals the omission of a US. It is commonly assumed that the excitation caused by the former is counteracted by an inhibition produced by the latter. This is again formalized by the summation term in the RW rule. Conditioned inhibition is thus one of the phenomena that, together with blocking and other compound phenomena, challenged the contiguity interpretation of classical conditioning.

A conditioned inhibition procedure involves reinforced trials with a CS, say A+, intermixed with non-reinforced trials with a compound AB-. Conditioned responding develops during A+ trials but not during AB-. Hence, conditioned inhibition is a key conditioning phenomenon since it is also a form of discrimination learning.

<sup>1073</sup> Conditioned inhibition poses higher technical challenges for a model of learn-<sup>1074</sup> ing and timing as responses cannot be directly observed. To assess conditioned

inhibition two types of measures are used (Denniston and Miller, 2007): sum-1075 mation and retardation tests. There are different procedures that can generate 1076 inhibition, so we refer here specifically to the inhibition produced by alternating 1077 A+ with AB- trials. CSA is called a training excitor, and CSB an inhibitor. 1078 In summation tests, this inhibitor is then presented together with a different 1079 excitor, and the inhibitor is said to pass the test if there is a decrement in re-1080 sponding compared to the excitor alone. In retardation tests, the inhibitor by 1081 itself is now paired with the US, and it is said to pass the test if acquisition 1082 is slower than with a neutral stimulus. Denniston and Miller (2007) reviewed 1083 a series of studies that varied the durations of the training excitor and that 1084 between the inhibitor and the training excitor. The studies showed that condi-1085 tioned inhibition is observed when the temporal relations between training and 1086 testing are preserved, and not otherwise. 1087

However, the studies reviewed by Denniston and Miller (2007) used as mea-1088 sure of conditioned inhibition the time to resume drinking (licking suppression) 1089 when presented with the inhibitor. Williams et al. (2008) investigated inhibi-1090 tion caused by reinforcement omission in excitatory conditioning, a more direct 1091 measure than licking suppression. In their experiments the inhibitor stimulus 1092 signalled the omission of one of two USs (at 10 or 30 seconds) that had been as-1093 sociated with the excitor stimulus. Using summation tests they found that the 1094 inhibitor would suppress responding only at the specific time of predicted US 1095 omission. Retardation tests confirmed that the time of US omission is encoded 1096 by the inhibitor. 1097

We show here that RWDDM can account for inhibition and its time specificity. CSC-TD and MS-TD are also equipped to deal with these results. MoT and LeT do not currently have the necessary mechanisms to explain inhibition.

### 1101 3.5.1 Simulations

We demonstrate time specificity of inhibition with simulations of Williams et al. 1102 (2008) experiment. Excitors E1 and E2 signalled reinforcement after 10 and 30 1103 seconds respectively, and inhibitors I1 and I2 signalled US omission after 10 1104 and 30 seconds respectively. During phase 1, E1 and E2 were always reinforced, 1105 whilst the compounds E1I1 and E2I2 were never reinforced (see table 3). In 1106 phase 2 a transfer excitor E3 was trained on a mixed FI schedule, where in 1107 half the trials E3 lasted 10 seconds and in the other half 30 seconds. Phase 3 1108 consisted of nonreinforced peak trials that lasted 90 seconds, a third with E3 1109 compounded with I1, a third with E3I2, and a third with E3 alone. Figure 9 1110 summarizes the results. Responding during E3 alone shows the two peaks char-1111 acteristic of mixed FIs. As figure 9 shows, the compound excitor and inhibitor 1112 inhibits responding only at the time encoded by the inhibitor. 1113

### 1114 3.5.2 Discussion

The account provided of inhibition by RWDDM relies on the traditional summation term inherited from the RW rule. Time specificity comes from the inhibitor CS timer being treated just like any other CS timer, except that instead of timing the arrival of the US it times the arrival of US omission.

RWDDM predicts that the representation of an inhibitor CS has the same shape as of an excitor CS. This implies that inhibition is the exact opposite of excitation. This is a testable prediction which the empirical results above provide some validation.

The TD models provide a similar account of these data. Both CSC and MS TD have CS representations that allow for time specificity of US omission. Because the TD relies on the RW summation term, they can account for inhibition. LeT and MoT can also represent such time specificity, but because they



Figure 9: Conditioned inhibition. Left column: simulation (top) and data (bottom) from conditioned inhibition with a long inhibitor. Right column: simulation (top) and data (bottom) from conditioned inhibition with a short inhibitor. Data plots redrawn from figure 4 Williams et al. (2008). Model parameters:  $\alpha_t = 0.09, \alpha_V = 0.06, \mu = 1, \sigma = 0.35, m = 0.16, H = 30.$ 

rely on the older linear operator rule, they do not have a mechanism to accountfor inhibition.

# <sup>1129</sup> 3.6 Disinhibition of delay and compound peak procedure

The two related phenomena described here are important in that they appear to challenge the summation effect. A common observation is that a compound of two previously conditioned CSs usually produces more responding than its individual components (Rescorla, 1997; Kehoe and Macrae, 2002, p. 204). However, failure to obtain summation is also common (Rescorla and Coldwell, 1995; Pearce et al., 2002), and the precise conditions when it is observed or not is still a current topic of debate (see Harris and Livesey, 2010, for a discussion). Here we consider two cases in which summation was not observed and that RWDDM can offer a possible explanation.

Aydin and Pearce (1995) used an autoshaping procedure to condition pigeons to stimuli of 30 second duration. They observed little or no summation in compound trials, but a response curve with a consistent shift to the left. This earlier start of responding was observed even when one of the components was a neutral preexposed CS. The shift of the response curve to the left was termed disinhibition of delay.

Meck and Church (1984) performed an analogue experiment using the peak 1145 procedure. They trained rats to associate a light and a sound (both of 50 second 1146 duration) individually to a reinforcement, and then used a peak procedure to 1147 investigate what happens to timing in their compound. Like Aydin and Pearce 1148 (1995) they also found no summation and a shift to the left in the compound. 1149 Furthermore, rats also stopped responding earlier in the compound peak trials. 1150 Taken together, these results appear to show that in some cases summation 1151 is not observed, and responding in the compound starts earlier than in the com-1152 ponent CSs. One possible explanation for this effect is that the subject fails 1153 to recognize the two individual components of the compound, what is known 1154 as generalisation decrement. If this is the case then it would be a performance 1155 effect, and not a learning phenomenon. We cannot rule this out, but we show 1156 that RWDDM's trial variability in time estimation provides a plausible mecha-1157 nism to explain this effect. The only other models in our analysis set that can 1158 account for this are MoT and LeT. 1159

### 1160 3.6.1 Simulations

RWDDM is capable of accounting for the earlier responding in compounds by 1161 noise in the timer. When a compound formed by CSA and CSB is presented, 1162 its two timers  $\Psi_A(t)$  and  $\Psi_B(t)$  will run in parallel. However, their rates  $A_A$ 1163 and  $A_B$  will have slightly different values due to noise. This implies that on 1164 every compound trial, one timer will be running slightly faster than the other. 1165 In contrast, on trials where only one CS is present, the timer will run faster in 1166 some trials and slower in others. Therefore, if on compound trials responding 1167 is guided by the faster timer, the average response curve for compounds will be 1168 shifted to the left when compared to the averaged response curve for a single 1169 CS. 1170

Figure 10 shows simulations of disinhibition of delay and compound peak 1171 procedure. The figures were constructed by averaging the responses produced 1172 by equation (10) over 50 trials. The simulations reproduce in part the an-1173 ticipation in responding during the compound that is observed in the data in 1174 both experiments (see top right and bottom left panels of figure 10). Meck and 1175 Church (1984) reported a median peak time of  $40\pm4$  seconds for the response 1176 curves in compound trials, and  $50\pm3.5$  seconds in the individual trials. We ran 1177 15 simulations as the one shown at the bottom row of figure 10, and analysed 1178 the peak times produced by each. We found an average peak time of  $42\pm3$  sec-1179 onds in the compound trials, and  $47\pm4$  in the individual trials. Both results are 1180 within the error bounds in Meck and Church (1984). Aydin and Pearce (1995) 1181 did not analyse peak times or shift in the response curves, so we cannot make 1182 a quantitative comparison with our simulations. 1183



Figure 10: Disinhibition of delay and compound peak procedure. Top row: simulation (left) and data (right) of disinhibition of delay. Bottom row: simulation (left and middle) and data (right) of a compound peak procedure. The middle panel is a normalized (proportion of maximum response strength) version of the left panel. Data plot redrawn from figure 13 in Meck and Church (1984). Model parameters: m = 0.25,  $\theta = 1$ ,  $\sigma = 0.18$ ,  $\alpha_t = 0.75$ ,  $\alpha_V = 0.1$ , H = 5.

## 1184 3.6.2 Discussion

RWDDM can offer a good account for the lack of summation and earlier re-1185 sponding in compound trials in the two cases analysed here. It does so by 1186 having trial to trial variability in time estimation. However, the model shows a 1187 slightly higher maximum response frequency in compounds than in their com-1188 ponents (top and bottom left of figure 10) something not observed in the data. 1189 This is not the product of summation, but of the slightly different asymptotes of 1190 learning in the faster and slower timers in the reinforced trial immediately pre-1191 ceding the peak trial. Our assumption was that in compound trials the timer 1192 running faster, with a higher slope A, would be the one guiding responding. 1193 When timing adaptation has reached asymptotic levels, the updates on slope 1194 A are due to noise in the value of the timer at reinforcement time,  $\Psi(t^*)$ . The 1195

two slopes,  $A_A$  and  $A_B$ , will have very similar values. In the reinforced trial 1196 preceding the compound peak trial, whichever timer produces a value of  $\Psi(t^*)$ 1197 lower than the threshold will have its slope A adjusted up by the slope 1198 update rule, likely causing it to overtake the other slope. This slightly higher 1199 slope will then be chosen in the peak trial that follows. But the corresponding 1200 V associated with that timer will have been updated on the previous reinforced 1201 trial based on the lower  $\Psi(t^*) < \theta$  value. Because that is the denominator in 1202  $HA/\Psi(t^*)$ , the V value of the chosen timer will be consistently slightly higher 1203 on the compound peak trials. 1204

Other theories that might account for the data in this phenomenon are LeT and MoT. Both theories postulate intertrial variability in timer rate, the same mechanism used by RWDDM to explain this data. TD in any of its current versions lacks a mechanism to explain these data.

# 1209 3.7 ISI effect

The interval between CS onset and US onset is called Inter Stimulus Interval 1210 (ISI). In general, measures of CR strength such as response frequency and ampli-1211 tude decrease with longer ISIs (Smith, 1968; Gormezano et al., 1983; Kehoe and 1212 Macrae, 2002). Response timing is commonly analysed by using fixed interval 1213 (FI) schedules of reinforcement, which rely on a fixed ISI. It is a well established 1214 result that the peak in the response curve decreases with longer FIs (Catania 1215 and Reynolds, 1968; Gibbon et al., 1997). However, the entire response curve 1216 approximately scales with FI. This is obtained by plotting different FI response 1217 curves as the proportion of maximum response strength versus the proportion 1218 to FI, a normalization procedure. The resultant normalized curves roughly su-1219 perimpose (Rakitin et al., 1998; Matell and Meck, 2000,0; Allman et al., 2014). 1220 This is sometimes called scalar timing, and it is one of the manifestations of the 1221

<sup>1222</sup> more general property of timescale invariance.

CSC-TD does not have a mechanism to explain either timescale invariance or the ISI effect. Its more recent development, MS-TD, can approximately reproduce both timescale invariance and the ISI effect. LeT is also a timescale invariant model, but does not appear to show the decrease in response peak as a function of FI. MoT, at least in its earlier version (Kirkpatrick, 2002), can reproduce both the ISI effect and timescale invariance.

#### 1229 3.7.1 Simulations

To demonstrate how RWDDM can reproduce the ISI effect we have simulated a 1230 delay conditioning procedure using three fixed interval stimuli. Figure 11 shows 1231 RWDDM simulations with FIs 5, 10 and 20 seconds. The top left panel shows 1232 within-trial response rate (given by equation (10)) averaged over 50 trials for 1233 each FI. The response curves show the same pattern as the data (bottom panel) 1234 from the ISI effect: a sigmoidal shape with a maximum that decreases as a 1235 function of FI duration. Note that because the curves are averages of 50 trials, 1236 the noise is averaged out. 1237

The top middle panel of figure 11 shows the associative strength acquisition curves for each FI. Their asymptotic levels are given by equation (20).  $V_{\infty}$  is approximately a linear function of  $A_{\infty}$ , the TDDM slope. The different asymptotic levels of associative strength are responsible for the different response peaks in the left panel of figure 11.

RWDDM also reproduces the superposition observed when FI response curves are normalized by maximum response rate and time to reinforcement (top right panel of figure 11).

## 1246 **3.7.2 Discussion**

<sup>1247</sup> Gibbon and Balsam (1981) attributed the ISI effect to the expectancy to re-



Figure 11: ISI effect. Top row: simulated average response rate during CSs (left), associative strength over trials (middle), and superimposition of response curves (right). Bottom row: average response rate data from an FI experiment, redrawn from bottom right panel of figure 4 in Kirkpatrick and Church (2000). Model parameters: m = 0.15,  $\theta = 1$ ,  $\sigma = 0.3$ ,  $\alpha_t = 0.2$ ,  $\alpha_V = 0.1$ , H = 5.

inforcement. A specific reinforcer carries, according to their view, an amount 1248 of expectancy H. This expectancy is spread back in time over the stimulus 1249 that signals US occurrence. Hence, for a CS of fixed duration T and US with 1250 expectancy amount H, the total expectancy during the CS is  $h_T = H/T$ . Our 1251 RWDDM account follows the same principles. The time to reinforcement T is 1252 computed by the ratio between the accumulation height at time of reinforcement 1253  $\Psi(t^*)$  and the timer slope at the current trial A(n). This leads to the asymptote 1254 of learning in equation 9 being set to  $HA_i(n)/\Psi_i(t^*)$ . Superimposition of the 1255 response curves follows directly in RWDDM from the nature of noise in the 1256 linear accumulator. This noise guarantees that the time estimate produced by 1257 the model is timescale invariant (Simen et al., 2013). 1258

The ISI effect can also be explained by the TD model with the Presence representation (Sutton and Barto, 1990) and with the more recently developed

Microstimuli representation (Ludvig et al., 2012). The Presence representation 1261 consists of a single element x which has the value 1 when the CS is present, 1262 and 0 otherwise. Its associative strength V is updated by the TD rule at every 1263 time step within a trial. In longer trials (longer FIs) the strength V will decay 1264 more, since it is updated more times in the absence of the US. This will lead to 1265 a lower asymptotic value for V. However, Presence TD cannot account for the 1266 superimposition of intratrial response curves. The CSC-TD fares even worse, 1267 unable to account for either ISI effect or superimposition (see Ludvig et al., 1268 2012, for a comparison between MS, CSC and Presence TD). The Microstimuli 1269 representation treats the stimulus as if it were composed of many units activated 1270 in sequence. Their activations follow a Gaussian shape which partially overlap. 1271 Later units have lower peaks and are wider than earlier ones. Because the 1272 number of Microstimuli are fixed, in longer FIs there is less temporal resolution 1273 which causes the US prediction to be lower than in shorter FIs, so it can explain 1274 the ISI effect. MS-TD's account of superimposition is only partial, although 1275 clearly better than CSC and Presence-TD. 1276

LeT in its current version lacks a mechanism to produce decreasing response 1277 peaks with increasing FIs. But it can account very well for superimposition, 1278 since its time representation is timescale invariant. The earlier version of Mod-1279 ular Theory, called Packet Theory, has been shown to produce the ISI effect (see 1280 top row of figure 3 in Kirkpatrick, 2002). This prediction comes from longer in-1281 terval durations decreasing the probability of response packet generation in the 1282 model. MoT is also timescale invariant, so it generates superimposition quite 1283 easily. 1284

To summarise, the ISI effect is explained either by time setting the asymptote of learning (RWDDM) or by a time representation that gets more diffuse with time, lowering the US prediction (MS-TD). Superimposition is explained either <sup>1288</sup> by the type of noise in the linear accumulator (RWDDM, LeT) or by stimulus <sup>1289</sup> units which have an approximately timescale invariant activation profile (MS-<sup>1290</sup> TD).

# 1291 3.8 Mixed FI

Procedures where a stimulus signals reinforcement at more than one location in 1292 time are called mixed FI or two-valued interval schedules. A mixed FI involves 1293 only one CS which could be of short or long duration, and the subject has no way 1294 of knowing which duration it is currently experiencing until the US is delivered. 1295 Catania and Reynolds (1968) conditioned pigeons in a mixed FI and reported 1296 a pattern of responding during the long CS that resembles a combination of 1297 two distinct FIs (with two peaks) when the separation between the intervals 1298 was in the ratio 8:1 but not at smaller proportions. Cheng et al. (1993) found 1299 a similar result (experiment 2) when the intervals were in 5:1 proportion and 1300 Leak and Gibbon (1995) showed that with intervals in the 8:1 proportion the 1301 scalar property (measured by the CV) holds approximately even for three-valued 1302 interval schedules. Whitaker et al. (2003) ran three experiments with Mixed FIs 1303 in rats and found two peaks with the same CV when the proportion between 1304 the durations was greater than 4:1, but not for smaller proportions. They also 1305 found that the peak height at the short duration was higher than at the long 1306 duration in most cases. Whitaker et al. (2008) used intervals in the very small 1307 proportion 2:1 and still found two peaks that became more distinct when the 1308 short interval was presented more often than the long. 1309

These results are interesting because they challenge in particular models of timing. They have served to provide evidence in favour of SET, and against BeT and the first version of LeT (Leak and Gibbon, 1995). Subsequently, they provided motivation for the development of the current version of LeT Machado et al. (2009). LeT can now account for the multiple response peaks in Mixed FIs, and their superimposition, but it cannot produce peaks with decreasing heights. Modular Theory has the necessary mechanisms to account for all the features of the data above. The TD models, MS and CSC, could both account for multiple peaks, but their account of superimposition would vary, with MS being superior than CSC. We show next that RWDDM can account for all features of the data in Mixed FIs.

### 1321 3.8.1 Simulations

In this simulation one CS was used which was followed by reinforcement either 1322 after 15 or 75 seconds randomly chosen, a proportion of 5:1. Our assumption 1323 was that in Mixed FI experiments subjects form two independent stimulus rep-1324 resentations, one for the short interval  $x_S$ , and another for the long interval  $x_L$ , 1325 each with its respective associative strength  $(V_S, V_L)$  and timer  $(\Psi_S, \Psi_L)$ . At 1326 CS onset, both timers begin timing, generating the two representations  $x_S$  and 1327  $x_L$ , and at each point in time behaviour is guided by the representation with the 1328 highest activation value. When a reinforcement occurs, the CS representation 1329 with the highest activation value is the one to which credit is assigned. 1330

The left panel of figure 12 shows the simulated responses averaged over 50 trials of the long 75-second duration. Two peaks, centred roughly at 15 and 75 seconds, of decreasing heights and increasing widths are clearly seen, matching roughly with the data (right panel).

### 1335 **3.8.2 Discussion**

RWDDM's mechanism for dealing with mixed FIs is in essence the same as for single FIs. The only difference is that instead of only one timer (and CS representation) in Mixed FIs RWDDM uses as many timers (and CS representations) as rewards. We have not however addressed explicitly how one CS can give rise



Figure 12: Mixed FI. Left: simulated response strength during long trials. Right: response strength data from a mixed FI experiment, redrawn from figure 3 in Leak and Gibbon (1995). Model parameters:  $\alpha_t = 0.2$ ,  $\alpha_V = 0.1$ ,  $\mu = 1$ ,  $\sigma = 0.425$ , m = 0.2, H = 30.

to two distinct representations. One possible explanation is that the slope adaptation rule (equation (6)) is only applied when the difference between the two intervals is below a certain amount. If the difference is above this amount, then the model would create a new representation. In fact, the data reviewed here suggests that animals may not be able to distinguish two intervals if they are in proportion below 2:1.

To the best of our knowledge, the only other model from our analysis set 1346 that has tried to address the behaviour in mixed FIs is LeT. Machado et al. 1347 (2009) have succeeded in obtaining the two peaks with the same CV using LeT. 1348 Their account relies on a single accumulator in the form of a series of states 1349 activated at a fixed rate. This rate is fixed within a trial, but varies from 1350 trial to trial. After repeated training with a mixed FI, the states around the 1351 reinforced times receive on average more associative strength than the ones away 1352 from them. This activation pattern generates the response peaks seen in the 1353 data. However, as the authors note, 'in mixed-FI schedules, the response rate 1354 [produced by LeT] at the first peak is equal to or lower than the response rate 1355

at the second peak, but never higher,' which is the opposite of what the data 1356 shows. The authors suggest that a decaying arousal function might need to be 1357 added to the model so as to allow response rate to decay with interval duration. 1358 Modular Theory is capable of accounting for the behaviour in Mixed FIs 1359 since its pattern memory for time is based on SET, which has been shown to 1360 account for these data (Leak and Gibbon, 1995). MoT's account is similar to 1361 RWDDM's in that both rely on a separate accumulator (and memory) for each 1362 time of reinforcement. CSC-TD would likely produce two peaks, since it relies 1363 on a perfect discretization of time into as many units as time-steps. But the 1364 curves would not superimpose when scaled as there is no mechanism to account 1365 for timescale invariance. MS-TD would also account for the two peaks but 1366 superimposition would likely not be fully obtained as its simulations of the ISI 1367 effect have only partially reproduced it (see section 3.7 and Ludvig et al., 2012). 1368

# 1369 3.9 VI and FI

Schedules of reinforcement specify the conditions of reinforcement delivery. There 1370 are a number of different types of schedules, some are based on the time elapsed 1371 between reinforcements, some on the number of responses emitted between re-1372 inforcements, but there can be other possibilities. Of particular interest for a 1373 timing and conditioning model are the two most commonly used time-based 1374 schedules: variable and fixed interval. Variable Interval schedules of reinforce-1375 ment (VI) consist in the delivery of a US following a CS that varies in duration 1376 from trial to trial. The CS durations are usually derived from an arithmetic or 1377 geometric sequence. In contrast, Fixed Interval schedules of reinforcement (FI) 1378 use a CS of fixed duration in all trials. Skinner and Ferster (2015) reported that 1379 VIs tend to produce behaviour with a constant rate throughout the trial, whilst 1380 FIs produce scalloped curves with a pause following each reinforcement and a 1381

1382 rapid increase in rate until the next reinforcement.

Catania and Reynolds (1968) performed a detailed analysis of behaviour under VIs and found that response rate declined with the average reinforcement rate. Within a trial response frequency increased with time, following approximately a negatively accelerated curve. When normalized by maximum response rate and time to reinforcement, these curves showed a considerable degree of superimposition.

<sup>1339</sup> Matell et al. (2014) trained rats on a VI in which intervals were sampled from <sup>1390</sup> an uniform distribution  $\mathcal{U}(15, 45)$ , and then tested using a peak procedure. They <sup>1391</sup> compared the VI response peak curve to the peak curve from a control group <sup>1392</sup> trained on an FI 30 (the mean of the VI distribution). Although the two curves <sup>1393</sup> were not significantly different statistically, the VI response peak curve peaked <sup>1394</sup> slightly earlier and was slightly higher than the control group.

Jennings et al. (2013) compared timing performance between VI and FI in three experiments, but found VI timing only in a VI where the average interval was 30 seconds. The other experiments from the same paper produced results more in agreement with the earlier work by Skinner and Ferster (2015) showing a constant rate of responding during VI trials.

Taken together, these studies appear to show that timing may sometimes be present during VI schedules. In this case, animals appear to be learning the average of the interval distribution. Here we demonstrate with simulations that RWDDM can account for such findings. The only other model in our analysis set that can account for this result is Modular Theory.

### 1405 **3.9.1** Simulations

In this simulation a random VI was produced by sampling intervals from a discrete uniform distribution  $\mathcal{U}(15, 45)$ . Non-reinforced peak trials of duration 135 seconds were interspersed during the VI, with a probability of 0.25. Our assumption here is that subjects will keep adapting the timer rate A over trials. In this case, equation (6) calculates the exponential moving harmonic average of the CS durations. Since it is a moving average, the predicted peak time will depend on the actual intervals used and their presentation order, but the nonmoving harmonic average of all intervals is 27.1 seconds. This is earlier than the arithmetic average (30 seconds), which is in line with the trend observed in the data by Matell et al. (2014).

Figure 13 (top left panel) compares the response strength averaged over peak trials in the VI and in a regular peak procedure with FI 30. The VI peak is higher and slightly earlier (at roughly 29.68 sec) than the FI peak, matching roughly with the data (bottom row). When normalized both by peak height and time the curves show the superimposition (top right panel) also seen in the data.

### 1422 **3.9.2** Discussion

The model predicts a harmonic mean value for the position of the response peak, 1423 which is always less than the arithmetic mean, but because it is a weighted 1424 moving average the actual value may vary. As we saw in the simulations, the VI 1425 response curve peaked at a value (29.68 sec) very near the arithmetic mean of the 1426 intervals (30 sec). This may explain the trend observed in the data by Matell 1427 et al. (2014). However, because that trend was not statistically significant, 1428 further experiments would be needed to establish if the response peak during 1429 VIs is nearer to the harmonic or the arithmetic mean. 1430

Taken together, these results are more easily accommodated by theories that can store an average of CS durations like RWDDM. Modular Theory is such an example, since it also stores an average of intervals in its pattern memory. Other models such as LeT and MS or CSC-TD would struggle with this result. The CS representation in these models break down the CS into a sequence of



Figure 13: VI and FI. Top row: simulated average response strength during peak trials (left), and the same data plotted after both axes are normalized (right). Bottom row: average response strength data from an experiment in VI and FI, redrawn from figure 1 in Matell et al. (2014). Model parameters:  $\alpha_t = 0.1, \alpha_V = 0.1, \mu = 1, \sigma = 0.3, m = 0.2, H = 40.$ 

units activated serially in time. With a uniform distribution of CS durations
associative strength would likely be spread broadly over the weights that cover
the interval, generating a broader pattern of responses that would not be centred
on the mean.

# 1440 3.10 Temporal Averaging

<sup>1441</sup> Although animals are able to time different durations simultaneously, as seen <sup>1442</sup> in mixed FIs, paradoxically under certain circumstances a type of temporal averaging can be observed. This is a relatively new and important phenomenon,
which challenges in particular theories of timing to propose a mechanism that
can explain such averaging.

When rats are trained using two distinct stimulus modalities, a visual stimu-1446 lus (a light) and an auditory (a tone), each signalling reinforcement at a different 1447 time, responding during compound presentations of both stimuli peaks roughly 1448 in the middle of both durations (Swanton et al., 2009). This intermediate re-1449 sponse curve to the compound superimposes with the two other single stimulus 1450 curves when normalized, suggesting that the animal is timing only one aver-1451 age duration. The type of average being computed appears to be modulated by 1452 the reinforcement probabilities associated with each stimulus duration, with the 1453 weighted geometric average fitting the data better than a weighted arithmetic 1454 average or a non-weighted average (Swanton and Matell, 2011; Matell and Hen-1455 ning, 2013; Matell and Kurti, 2014). Significantly, temporal averaging in rats 1456 is only consistently observed when the auditory stimulus signals the short in-1457 terval and the visual stimulus signals the long interval (Swanton and Matell, 1458 2011; Delamater and Nicolas, 2015). Even when each stimulus is associated 1459 with a different response option (light reinforced with a left nosepoke, tone with 1460 a right) rats still tend to mix the temporal information during compound trials 1461 (De Corte and Matell, 2016). 1462

We do not make a strong claim about RWDDM's ability to explain this data. Rather, we show that it has the necessary elements from which an account can begin to be formulated. MoT also has similar elements from which an account can be built. CSC-TD, MS-TD and LeT do not appear to be equipped to deal with this phenomenon.

### 1468 3.10.1 Simulations

In RWDDM the accumulator is the mechanism that marks the passage of time. 1469 The temporal proximity to an event is determined by how close the level of 1470 accumulation is to a fixed threshold value. A CS that signals reward later than 1471 another CS, will have a lower rate  $(A_{low})$  of accumulation than the shorter CS 1472  $(A_{high})$ . Because in RWDDM associative strength is set by time to reward, the 1473 two CSs will also have different associative strengths,  $V_{\rm low}$  and  $V_{\rm high}$  respectively. 1474 We may assume that under temporal averaging circumstances the stimuli are 1475 of such nature that they cause the subject to integrate their information. At 1476 the start of the compound trials, the ambiguity presented by the compound 1477 stimulus may cause the representations of the two component stimuli to be only 1478 partially retrieved. If the subject fails to represent the two stimuli separately, 1479 the result may be the formation of a single representation composed by only 1480 a fraction of the timing rate A and associative strength V of each individual 1481 stimulus. The fractions are then added into one single rate and one single 1482 associative strength, and processed as if they were the components of a single 1483 stimulus representation. For the simulation below, we assume that the fractions 1484 added are exactly half of their individual values:  $A_{\text{compound}} = A_{\text{low}}/2 + A_{\text{high}}/2$ , 1485 and  $V_{\text{compound}} = V_{\text{low}}/2 + V_{\text{high}}/2$ . 1486

We used a long CS of duration 20 seconds and a short CS of duration 10. We simulated a peak procedure with each CS and with the compound. A plot of the response strength averaged over peak trials is shown in the top left panel of figure 14. The three peaks scale when normalized (top right panel).

The peak of the compound is roughly at 13.33 sec, which would be the expected value for an averaged rate A = (1/10 + 1/20)/2, the harmonic average of the intervals. The height of the compound peak is also at an intermediate level between the two end peaks. The simulations match roughly with the data



Figure 14: Temporal averaging. Top row: simulated response strength averaged over peak trials in temporal averaging (left), and the same data normalized by maximum response strength and peak time (right). Bottom row: peak trial response strength data from an experiment in temporal averaging, redrawn from figure 1 in Swanton et al. (2009). Model parameters:  $\alpha_t = 0.2$ ,  $\alpha_V = 0.1$ ,  $\mu = 1$ ,  $\sigma = 0.35$ , m = 0.2, H = 30.

1495 (bottom row of figure 14)

### 1496 **3.10.2** Discussion

<sup>1497</sup> The assumption we made here, that temporal averaging is the result of only <sup>1498</sup> one accumulator being active during the compounds and fed with half the rate <sup>1499</sup> for each of the stimuli, is plausible and can accommodate the main features of <sup>1500</sup> the data. However, given the evidence from mixed FIs it seems animals are

capable of keeping multiple timers running in parallel, without averaging their 1501 rates. Also, if averaging of rates always happened during compounds, then 1502 the explanation provided by RWDDM for the left shift in the response curve 1503 in the compound peak procedure would not hold. We suggest one possible 1504 way of interpreting these three phenomena based on a failure of representation 1505 selection caused by the ambiguity of the signal. In mixed FIs there is one 1506 single CS that signals two rewards at very different times. There is not much 1507 ambiguity in how to interpret the signal, so the subject keeps two timers running 1508 in parallel. In the case of compounds formed by individual CSs that signal 1509 reward at the same time, as in the compound peak procedure, there is also 1510 not much ambiguity. There's very little difference between the time memories 1511 evoked by the CSs, so choosing only one, the faster one, leaves no ambiguity as 1512 to which CS is signalling reward. In the case of compounds formed by individual 1513 CSs of different modalities that signal reward at different times, the ambiguity 1514 might be such that cannot be resolved easily. The information from each CS 1515 may then be only partially retrieved and added into one representation, resulting 1516 in temporal averaging. 1517

As mentioned previously, this is not a strong account of the conditions that 1518 generate temporal averaging. But whatever the final word on this may be, 1519 RWDDM has components that allow it to generate averaging and timescale 1520 invariance. However, RWDDM predicts this average to be the harmonic mean, 1521 and not the geometric mean weighted by reinforcement probabilities that has 1522 been frequently found (Swanton and Matell, 2011; Matell and Henning, 2013; 1523 Matell and Kurti, 2014). Also, Matell and Henning (2013) reported evidence 1524 of summation of response rates during the compound trials. In our simulations 1525 here we assumed that equal fractions were taken of the rates of each CS, resulting 1526 in a combined non-weighted harmonic average of rates, but different fractions 1527

(or weights) may be taken. In particular, the data indicates that the weights are set by the reinforcement probabilities of each individual stimulus. Since this information is stored in the associative strength V, we could assume the subject integrates the two timer rates as follows:

$$A_{\text{compound}} = \left(\frac{V_{\text{low}}}{V_{\text{low}} + V_{\text{high}}}\right) A_{\text{low}} + \left(\frac{V_{\text{high}}}{V_{\text{low}} + V_{\text{high}}}\right) A_{\text{high}}$$

Although this would produce a weighted average, it is still a weighted harmonic average of the intervals and not a weighted geometric average found in the data, so the account given by RWDDM would still be partial. As for the summation of response rates observed in the compound trials, this could be explained by RWDDM if instead of taking a fraction of the V values for each stimulus to form the V compound, the subject simply summed, or partially summed, both V values.

Another model that is equipped to deal with averaging is Modular Theory. 1539 If we allow for one single accumulator fed by one half of each time memory, 1540 then MoT would predict a peak of responding at the arithmetic mean of the two 1541 intervals. A weighted average could also be obtained following the procedure we 1542 sketched above for RWDDM. However, this would yield a weighted arithmetic 1543 mean, and not the weighted geometric mean obtained in the data. As for 1544 timescale invariance, MoT relies on a noisy timer threshold whose mean is always 1545 a fixed proportion of the time memory, with a standard deviation proportional to 1546 this mean. Therefore, timescale invariance is guaranteed for all time memories, 1547 averaged or not. 1548

LeT would not be able to explain temporal averaging without modifications. It cannot change its average transition rate between states without compromising timescale invariance. Without changing the transition rate it is difficult to see how else LeT could account for a different timing in the presence of the
compound. CSC-TD and MS-TD also lack any mechanism that could be usedto account for temporal averaging.

## 1555 3.11 Summary of Results and Analysis

Table 4 summarizes the results from the simulations. RWDDM was able to reproduce the main features of the data in 8 out of the 10 experiments. In the other 2 the model was able to partially account for the data.

To allow for comparison we have offered qualitative predictions for the other 1559 4 models in table 4. It is important to note that for most of the 10 phenomena 1560 analysed here simulations using these models are not available in the literature. 1561 Although we have tried our best to provide predictions based on our under-1562 standing of these models, we have not actually simulated them. Therefore it is 1563 possible that in some cases a model may produce results that we did not foresee 1564 if the right set of parameters is found or some of the assumptions are relaxed. 1565 It is also possible that some simple modifications might allow the models to 1566 explain the data. We endeavoured to point out some such modifications that 1567 seem likely to work when discussing the simulation results above, but we do not 1568 make predictions based on them because the purpose here is only to provide a 1569 comparison of the current mechanisms of each model and therefore encourage 1570 future work on model improvement. With that in mind, Modular Theory has 1571 fared best after RWDDM, being able to account for 7 out of the 10 experiments. 1572 MS-TD and CSC-TD shared the second place with 3 out of 10. LeT came in 1573 last, able to account for 2 experiments. The last column of table 4 identifies the 1574 main mechanisms responsible for successfully accounting for each phenomenon. 1575

## 1576 4 General Discussion

RWDDM was able to reproduce faster reacquisition due to its memory for time 1577 being conserved during extinction. This memory is used to activate the stimulus 1578 representation. Learning is slower in acquisition because RWDDM increases the 1579 activation in the stimulus representation gradually over the trials. The stimulus 1580 representation needs to be 'built up' first, and this process depends on learning 1581 the timing of the US. Extinction eliminates associative strength but leaves the 1582 time memory, hence the stimulus representation, intact. Reacquisition proceeds 1583 faster because the stimulus representation does not need to be built up again. 1584 Other models explain this by allowing the associative strength learning rate to 1585 be faster in reacquisition. 1586

Time change in extinction was accounted for because of RWDDM's ability to time CS duration independently from US associations. Time is learned entirely by time markers. The TD models and LeT do not make this separation. These models do not have a mechanism to time stimuli without the US stamping in the changes.

Improved timing in latent inhibition was also accounted by RWDDM's abil-1592 ity to learn timing independently of associations. Preexposure allows the model 1593 to build its time representation, which is later expressed by behaviour during 1594 the acquisition phase. The only other model that learns to time independently 1595 of associations is MoT, but it does not have a mechanism to explain the latent 1596 inhibition effect. The latent inhibition effect alone, i.e. the initial decrement in 1597 the acquisition curve of a preexposed stimulus, was made possible in RWDDM 1598 by using the P-H rule to change the learning rate for associative strength. The 1599 use of the P-H rule instead of the RW would certainly have other theoretical 1600 implications for the general theory we are introducing in this paper, but we 1601 have used it only in this case. We will make further comments in the conclu-1602

sion. Blocking with different durations was easily accounted in one condition, 1603 the short blocked and long blocking CS. The blocking effect in this condition 1604 followed from the summation term in the RW rule. For the other condition, long 1605 blocked and short blocking CS, a straight application of the model did not yield 1606 the results expected. But the experimental results leave open the possibility 1607 that this might be a case of second-order conditioning, where the summation 1608 term in RW does not play a role. In this case, RWDDM is well placed to ex-1600 plain the results, since it can time the whole sequence of stimuli. The only other 1610 models capable of explaining these results were the TD models. 1611

The time specificity in conditioned inhibition was very well accounted for by the combination of the summation term in the RW rule, which allowed for inhibition to develop, and the independent timing mechanism in RWDDM that allowed it to time US omission. However, the alternative account provided by the different time representation in the TD models was also successful. The other theories failed here for the same reason as in blocking, they lack a rule like RW that can deal with compound stimuli effects.

The response curves centred at the mean of intervals in the VI procedure was well accounted by the ability of RWDDM to learn the average of intervals. This ability is only present in Modular Theory, making it the only other model able to account for the results here.

In the case of temporal averaging, RWDDM was able to account for the general features of the phenomenon, namely a response curve that peaks at the average of the intervals signalled by the compound stimulus. However, RWDDM predicts the peak to be at the harmonic mean, whilst some experimental results suggest it happens at the geometric mean. RWDDM's account of temporal averaging was hypothesised as the result of ambiguity in the signal. In trying to resolve whether the compound should be treated as a single stimulus or as two separate stimuli, the subject settles on using one accumulator that is fed partial timing information from both stimuli. Other hypothesis might turn out to be more adequate, but this is one possibility that fits well with the RWDDM framework. The only other model that would produce averaging under the same hypothesis is MoT.

The classic ISI effect followed from two mechanisms in RWDDM. The lower 1635 response curves during longer stimuli were explained by time setting the asymp-1636 tote of associative learning by hyperbolic delay discounting. The larger spread 1637 of response curves during longer stimuli and the superimposition of normalised 1638 curves follows from RWDDM's timescale invariant time representation. The 1639 noise in RWDDM's accumulator decreases with the interval being timed in such 1640 a way that it results in timescale invariance of the response curves. Modular 1641 Theory can also reproduce all features in the data. This is because it relies on 1642 a timescale invariant response rule function that generates less responding in 1643 longer intervals. LeT can account for superimposition, but it does not have a 1644 mechanism to account for the lower curves in longer stimuli. MS-TD can ac-1645 count for both elements because of the form of its microstimuli representation. 1646

The double peaks observed in the response curves during mixed FIs is ex-1647 plained by RWDDM using simultaneous timing. It generates two different repre-1648 sentations, one for each reward. Thus, it can account for mixed FIs by the same 1649 principles used to account for the ISI effect and simple FI schedules. Modular 1650 Theory takes the same approach of simultaneous timing and is also successful. 1651 The TD models and LeT can provide a partial account due to their distributed 1652 time representation. But timescale invariance of the peaks is not observed in 1653 CSC-TD and only approximately in MS-TD. LeT produces the timescale invari-1654 ance but not the decrease in peak height with time. 1655

1656

The left shift of response curves seen in compound peak procedure and dis-

inhibition of delay was well accounted for by RWDDM. It did so because of
intertrial variability in noise estimation. By choosing in every compound trial
the time memory that predicts reward sooner, RWDDM produces the left shift
in response. The only other models that can appeal to the same principle to
explain it are LeT and MoT.

The superiority of RWDDM and MoT in explaining the majority of the phe-1662 nomena analysed highlights the importance of some of their shared mechanisms. 1663 Both models have separate rules for updating time and associative strength. 1664 This makes them capable of timing any stimuli, independent of changes in asso-1665 ciative strength. Both models represent psychological time as linearly related to 1666 physical time through the theoretical construct of the accumulator. Their mem-1667 ory for time stores a moving average of the experienced intervals. They both 1668 allow for intertrial variability in time estimation. Among their differences, only 1669 one proved crucial in discriminating the two models in the experiments anal-1670 ysed here: the lack of a mechanism in MoT to account for stimulus compounds. 1671 RWDDM uses the RW rule, which was developed to deal with phenomena such 1672 as blocking and inhibition, whilst MoT uses the linear operator, a historically 1673 earlier association rule that cannot handle compounds. This was the single dif-1674 ference that caused the difference between MoT and RWDDM in number of 1675 phenomena explained. 1676

MS-TD came in third place in number of phenomena successfully explained, but the gap between it and MoT was comparatively high, with MoT being almost twice more successful than MS-TD. CSC-TD came just half a point below MS-TD. This is certainly a result of their similarities. The only difference between these two TD models is in their time representation. However, this different representation allowed MS-TD to explain only one more phenomenon than CSC-TD, the ISI effect. Therefore, in the set of experiments analysed

here MS-TD did not show a significant improvement on CSC-TD. This does 1684 not mean that MS-TD is not a significant improvement on CSC-TD overall. 1685 Its superior account of timing is significant. But the set of experiments chosen 1686 here are particularly challenging even for a dedicated timing theory, so they 1687 raise the bar even higher. The strength of the TD models was in account-1688 ing for compound phenomena of blocking and inhibition, due to their RW rule 1689 for association. Their weaknesses was that they rely on changes in associative 1690 strength to express changes in timing. This prevented them from explaining 1691 time change in extinction and improved timing in latent inhibition. They both 1692 lack a memory to store the average of intervals, so they could not explain be-1693 haviour in VI schedules. Finally, their lack of trial to trial variability in time 1694 estimation prevented them from accounting for the left-shift in the compound 1695 peak procedure. 1696

With respect to the number of successes only, LeT came in last. The results 1697 allowed us to identify at least four limitations in LeT's current formulation. The 1698 first is that it ties its time representation to changes in associative strength. 1699 This prevented it to explain time change in extinction and improved timing in 1700 latent inhibition. The second limitation is that it relies on the linear operator 1701 rule for associative strength, which prevented it from accounting for blocking 1702 and time specificity in conditioned inhibition. Thirdly, its distributed memory 1703 for time does not store the average of the intervals seen. This prevented it 1704 from accounting for the behaviour in VI. Lastly, it doesn't have a mechanism 1705 to explain the decrease in peak height of the response curves with longer ISIs. 1706 However, as a timing model, LeT's strength is in explaining timescale invariance. 1707 If it can be made to overcome at least the weakness of its associative learning 1708 rule, for example by also adopting the RW to update associative strength, LeT 1709 could be on a par with the TD models. 1710

RWDDM faced a few problems in explaining the set of phenomena analysed 1711 here. In latent inhibition the model was able to learn the timing for the pre-1712 exposed CS, but our choice of CS representation translates this into a response 1713 curve that does not fully match the data. A better solution might involve a 1714 two-state CS representation, one state for the early stages of training and the 1715 other for the latter stages. RWDDM could not account for the lack of blocking 1716 with a long blocked CS and a short blocking CS. One possible solution that 1717 does not require changing the model is to treat the blocking CS as a secondary 1718 reinforcer. A more difficult problem related to asynchronous co-terminating 1719 CSs such as the ones used in the blocking experiment analysed here, is that 1720 in its current formulation RWDDM cannot produce a stable solution. Because 1721 RWDDM assigns a different learning asymptote for each CS in the compound, 1722 it generates an inconsistent system of equations for V. How to fix this remains 1723 an open problem. Finally, in temporal averaging RWDDM predicts a peak in 1724 CR at the harmonic mean of the intervals, not at the geometric mean as has 1725 been observed in the data. More experiments might help to determine if the 1726 harmonic average should indeed be ruled out as an explanation. 1727

One relevant phenomenon that we did not explore here is the peak procedure. 1728 In particular, Balci et al. (2009) have produced evidence that in the long peak 1729 trials animals don't stop responding immediately after the expected reward 1730 time, but instead take a number of peak trials to learn to stop. The Gaussian 1731 function  $x_i(\Psi_i)$  used as the CS representation in RWDDM ensures that CR 1732 levels will begin to decrease after  $\Psi_i(t)$  crosses threshold  $\theta$  without any learning. 1733 To address the findings in Balci et al. (2009) the RWDDM CS representation 1734 could be changed to a sigmoid, saturating after the timer  $\Psi(t)$  crosses a first 1735 threshold. A second threshold could then be introduced to mark the time to stop 1736 responding. When the timer crosses this stop threshold the saturation process 1737

in the CS representation would stop and a decay process would begin. This 1738 however would still be an incomplete account, as a mechanism would be needed 1739 to explain the learning of the second threshold. But if such a CS representation 1740 was used, the model would also fit a larger body of data coming from studies that 1741 analyse responding during individual trials of the peak procedure. Schneider 1742 (1969) and subsequently Gibbon and Church (1990) and others (Cheng and 1743 Westwood, 1993; Matell et al., 2006) have argued that the pattern of responding 1744 is better characterized not by a Gaussian but instead by an approximate square-1745 wave function, with a low-high-low response frequency pattern. It can be shown 1746 that by introducing a stop threshold to the timer  $\Psi_i(t)$ , the TDDM timer (used 1747 in RWDDM) can fit the data on times of start and stop responding (Luzardo 1748 et al., 2017). Alternatively, the accumulator  $\Psi_i(t)$  itself could be used as the 1749 CS representation, replacing  $x_i$  in equations (9) and (10). In this case, an 1750 upper absorbing boundary would need to be set on the accumulator to prevent 1751 response strength increasing considerably in the first few trials following a CS 1752 duration increase for example. Also, such a choice of CS representation would 1753 cause within-trial responding to become linear, rather than the more commonly 1754 observed sigmoidal pattern. If a sigmoidal response curve is to be preserved, a 1755 different choice of response function would be required. 1756

Another phenomenon that we did not address but deserves mention is the 1757 timescale invariance of the acquisition process (Gallistel and Gibbon, 2000). It 1758 refers to the general finding that the number of trials required until an acqui-1759 sition criterion is met depends on the ratio of intertrial (or context) and trial 1760 durations, the I/T ratio (Gibbon, 1977; Lattal, 1999; Holland, 2000). Gibbon 176 and Balsam (1981) provided an account for this that postulates a decision pro-1762 cess based on the reward expectancy signalled by the stimulus versus the one 1763 signalled by the context. A ratio between the two expectancies is calculated, and 1764

once the ratio exceeds a certain value, acquisition starts. If the same postulate 1765 of a decision ratio of reward expectancies is made, RWDDM may account for 1766 the I/T ratio in a similar manner. If we assume that animals time the interval 1767 between USs (the context or I duration) with rate  $A_I(n)$  and also the CS dura-1768 tion as usual with rate  $A_T(n)$ , then we can form the ratio  $r(n) = A_T(n)/A_I(n)$ . 1769 As the number of trials n increases, the A rates converge to their asymptotic 1770 values, and the ratio r will converge to  $A_T/A_I = (1/T)/(1/I) = I/T$ . This 1771 is essentially the same account given by Gibbon and Balsam (1981), with the 1772 timer rates  $A_T$  and  $A_I$  substituting Gibbon and Balsam's expectancies H/T1773 and H/C. 1774

At least three testable RWDDM predictions came out from the simulations 1775 reported here. The first concerns blocking with different durations. A long 1776 blocked CS will not be blocked by a short co-terminating blocking CS, and two 1777 peaks in responding will be observed during test trials with the blocked CS: one 1778 at the time the short blocking CS would normally start, and another at the end of 1779 the blocked CS. The second prediction is that conditioned inhibition is the exact 1780 opposite of excitation. This means that the behaviour produced by inhibition is 1781 timed in the same manner as in excitation. Finally, in temporal averaging the 1782 response peak in the compound stimulus should be at the harmonic average, 1783 or weighted harmonic average. One prediction that did not come out of the 1784 simulations but that is worth mentioning concerns time estimation during very 1785 early trials. Our assumption of a low initial value for the accumulator rate 1786 A implies that in the initial trials durations will be overestimated. A new 1787 experiment testing this prediction could help validate, or invalidate, the model. 1788 RWDDM is, to the best of our knowledge, the first time the RW associative 1789 learning rule is coupled with a accumulator-based timing theory. An important 1790 implication of this effort for associative learning is that it allows for a richer 1791

analysis of the effects of timing in compound stimuli experiments. Here we have 1792 analysed blocking and conditioned inhibition, but there is evidence suggesting 1793 time may have important effects in other cue-competition phenomena such as 1794 overshadowing (Kehoe and James, 1983; Jennings et al., 2007). Timing effects 1795 in compounds has until now received somewhat little attention, with many pub-1796 lished experimental studies reporting only aggregate response measures. This is 1797 perhaps to be expected, since most associative learning models that can handle 1798 compounds do not have any, or a rich enough, time representation. RWDDM 1799 is an attempt at filling this theoretical gap. 1800

Another limitation of associative learning models is that they tend to simply 1801 postulate the timing features of the stimulus representation, without a detailed 1802 account of how these can mechanistically arise and evolve. This is the case with 1803 the CS representations of CSC-TD, MS-TD and others like C-SOP (Brandon 1804 et al., 2003). RWDDM's adaptive timer and time-adaptive CS representation 1805 provide a fuller account of the timing mechanism and its dynamics. Another 1806 recent model that provides this level of detail is the Timing from Inverse Laplace 1807 Transform (TILT, Shankar and Howard, 2012; Howard et al., 2015). It can 1808 dynamically develop a timescale invariant representation of stimulus history 1809 using a two-layer neural network. It can also reproduce the important I/T ratio 1810 conditioning phenomenon, but so far it has only been implemented with the 1811 linear operator rule for associative learning, which precludes it from accounting 1812 for cue competition phenomena. 1813

The RWDDM architecture suggests that timing is largely independent of the process of association formation and maintenance. Associations however, according to RWDDM, depend on timing both to set the asymptote of associative strength and to build the CS representation so that it can enter into association with the US. Thus, RWDDM implies that interactions between timing and associative learning are mainly one-directional. This appears to match roughly with experimental findings. In a review Kirkpatrick (2013) found that prediction error influenced measures of time estimation only through changes in reward magnitude and devaluation, whilst effects in the other direction included the appropriate timing of CRs from start of conditioning, trial and intertrial durations affecting strength and probability of CR occurrence, and cues with different temporal information affecting cue competition.

## 1826 5 Conclusion

In this paper we introduced a new real-time model for classical conditioning and
timing. The model combines elements from two theories, the Rescorla-Wagner
conditioning model and the TDDM interval timing theory.

We have simulated the model on 10 conditioning phenomena selected from 1830 the literature, which collectively represent a particular challenge for any single 1831 model to explain. The model was successful in accounting for 9, and can be 1832 made to account for the rest if simple modifications are made. The mechanisms 1833 used by other models of similar scope were evaluated to see if they could also 1834 account for the data. The model that got closer to this level of success in this set 1835 of phenomena was Modular Theory. This was due to MoT and RWDDM having 1836 a significant overlap in terms of mechanisms. Both models use an accumulator 1837 to mark the passage of time. Both models require only a single associative 1838 unit per stimulus that adapts to the temporal information conveyed by the 1839 stimulus. Their main difference is that MoT still uses the linear operator rule 1840 which precludes it from explaining blocking and other compound phenomena, 1841 whilst RWDDM uses the RW which can account for those phenomena. The 1842 same limitation is faced by TILT, a recent model that we did not analyse but 1843 that shows promising results and has desirable timing properties. 1844

RWDDM may be improved in several ways. It is quite likely that the asymp-1845 tote of learning may not be described by the simple inverse relationship to re-1846 inforcement time that we assumed. In some of the experiments modelled here, 1847 response peak seemed to decrease slower with ISI than our inverse relation-1848 ship predicted. Functions other than Gaussians might be used to represent the 1849 CS, which could better fit the data in the case of latent inhibition for example. 1850 These and other theoretical issues may be better elucidated by new experiments 1851 involving compound stimuli and a manipulation of their durations, such as the 1852 experiments with blocking, compound peak procedure and temporal averaging 1853 analysed here. 1854

We have also adopted the P-H rule in one experiment, but have not explored 1855 its application in the others. Making the P-H rule an integral part of RWDDM 1856 would add one more parameter but it would also allow RWDDM to account for 1857 other preexposure and attentional effects that the rule is designed to account. 1858 This is not a difficult modification, and we have already shown it to be feasible. 1859 RWDDM may be regarded, like TD, as a real-time extension of RW. Unlike 1860 TD and LeT, it does not require a number of associative units that grows linearly 1861 with time. It adds to RW the powerful timing mechanism of TDDM. But also, 1862 by making a link with a version of DDM, it shows that it may be possible to 1863 arrive at a unified account of timing, conditioning and decision making. 1864

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Simulation	Group	Phase 1	Phase 2	Phase 3
Acquisition, extinction and reacquisition	1 FI 5	80 CS+	100 CS-	80 CS+
The state of the s	FI 20-40	150  CS(20) +	150 CS(40)-	
Extinction with diff. duration	FI 20-10	150 CS(20) +	150 CS(10)-	
	FI 5			
ISI effect	FI 10	150  CS+		
	FI 20			
	VI 30	mixed $1500 \text{ CS}+$ , $375 \text{ peak}$		
T 1 S 1 T 1	FI 30	mixed $500 \text{ CS+}, 125 \text{ peak}$		
Mixed FI	MFI 15-75	mixed 200 A(15)+, 200 $\overline{A}(75)+$		
	Preexposed A	80 A-	120  A+	
Latent inmibition	Control C		120 C+	-
	Blocking A	120 A(10  or  15)+	60 A (10) D1 (16) 1 60 A (16) D1 (10) 1	
Blocking diff. durations	Blocked B1, B2		$(10) \times 100 \times 1000 \times 100 \times 10$	-
	Control C	I	60 C(10)B2(15)+ or 60 C(15)B2(10)+	
Disinhibition of delay	FI 30	100  A+, 100  B-	mixed 300 A+, 300 B+, 100 AB+	
Compound peak	FI 50	100  A+, 100  B+	mixed 300 A+, 300 B+, 100 AB peak	
Conditioned inhibition	1 group	mixed 300 each $E1(10)+$ , $E2(30)+$ , $E1(10)I1(10)-$ , $E2(30)I2(30)-$	mixed $300 \text{ each } \text{E3}(10)+, \text{E3}(30)+$	100 each peak E3I1, E3I2, E3
Temporal averaging	FI 10-20	700 L(20)+, 700 S(10)+, 154 L peak, 154 S peak, 154 SL peak		

Table 3: Simulation designs.

phenomenon	RWDDM	CSC-TD	MS-TD	LeT	MoT	explaining mechanism
faster reac- quisition	yes	yes <sup>1</sup>	yes <sup>1</sup>	yes <sup>1</sup>	yes <sup>1</sup>	time-adaptive stimu- lus representation or changes in learning rate
time change in extinction	yes	no	no	no	yes	separate rules for time adaptation and asso- ciative strength
latent inhibi- tion and tim- ing	part.	no	no	no	no	PH rule and separate rules for time adap- tation and associative strength
blocking with diff. dura- tions	part.	yes	yes	no	no	RW rule and ability to time any stimulus or distributed time rep- resentation
time spec. of conditioned inhibition	yes	yes	yes	no	no	RW rule and con- centrated memory for time or distributed time representation
compound peak proce- dure	yes	no	no	yes	yes	intertrial variability in time estimation
ISI effect and superimposi- tion	yes	no	part.	part.	yes	asymptote of assoc. strength set by time and accumulator noise or time representation that gets diffuse with longer time
mixed FI	yes	part.	part.	part.	yes	ability to generate multiple time rep- resentations or a single distributed time representation
VI and FI	yes	no	no	no	yes	memory that stores average of intervals
temporal av- eraging	yes	no	no	no	yes	memory that stores average of intervals and the accumulator

Table 4: Summary of main simulation results and comparison with other models. Notes: (1) if learning rate is allowed to vary.