1	Encoding time in neural dynamic regimes with distinct
2	computational tradeoffs
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24 Abstract

25 Converging evidence suggests the brain encodes time in dynamic patterns of neural 26 activity, including neural sequences, ramping activity, and complex dynamics. Most temporal 27 tasks, however, require more than just encoding time, and can have distinct computational 28 requirements including the need to exhibit temporal scaling, generalize to novel contexts, or 29 robustness to noise. It is not known how neural circuits can encode time and satisfy distinct 30 computational requirements, nor is it known whether similar patterns of neural activity at the 31 population level can exhibit dramatically different computational or generalization properties. To 32 begin to answer these questions, we trained RNNs on two timing tasks based on behavioral studies. 33 The tasks had different input structures but required producing identically timed output patterns. Using a novel framework we quantified whether RNNs encoded two intervals using either of three 34 35 different timing strategies: scaling, absolute, or stimulus-specific dynamics. We found that similar 36 neural dynamic patterns at the level of single intervals, could exhibit fundamentally different 37 properties, including, generalization, the connectivity structure of the trained networks, and the contribution of excitatory and inhibitory neurons. Critically, depending on the task structure RNNs 38 39 were better suited for generalization or robustness to noise. Further analysis revealed different 40 connection patterns underlying the different regimes. Our results predict that apparently similar neural dynamic patterns at the population level (e.g., neural sequences) can exhibit fundamentally 41 42 different computational properties in regards to their ability to generalize to novel stimuli and their robustness to noise—and that these differences are associated with differences in network 43 connectivity and distinct contributions of excitatory and inhibitory neurons. We also predict that 44 45 the task structure used in different experimental studies accounts for some of the experimentally observed variability in how networks encode time. 46

47 Author summary

48 The ability to tell time and anticipate when external events will occur are among the most 49 fundamental computations the brain performs. Converging evidence suggests the brain encodes 50 time through changing patterns of neural activity. Different temporal tasks, however, have distinct 51 computational requirements, such as the need to flexibly scale temporal patterns or generalize to 52 novel inputs. To understand how networks can encode time and satisfy different computational 53 requirements we trained recurrent neural networks (RNNs) on two timing tasks that have 54 previously been used in behavioral studies. Both tasks required producing identically timed output patterns. Using a novel framework to quantify how networks encode different intervals, we found 55 56 that similar patterns of neural activity-neural sequences-were associated with fundamentally 57 different underlying mechanisms, including the connectivity patterns of the RNNs. Critically, 58 depending on the task the RNNs were trained on, they were better suited for generalization or robustness to noise. Our results predict that similar patterns of neural activity can be produced by 59 60 distinct RNN configurations, which in turn have fundamentally different computational tradeoffs. Our results also predict that differences in task structure account for some of the experimentally 61 62 observed variability in how networks encode time.

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65 Introduction

66 The ability to predict when external events will occur, and to detect temporal regularities 67 in the environment, are among the most fundamental computations the brain performs [1-5]. Thus, 68 the brain must have a rich repertoire of mechanisms to tell time and perform temporal 69 computations. Indeed, converging experimental and computational evidence indicates that a wide 70 range of different brain areas encode time through dynamically changing patterns of neural activity 71 [1, 6-10]. These patterns can take the form of monotonic ramping of the firing rates of neurons, or 72 so-called population clocks that can take the form of neural sequences or complex patterns of 73 neural activity [1, 11].

Experimental and computational analyses of the different neural encoding schemes for the 74 representation of time have focused primarily on the discrimination and production of isolated 75 76 intervals or durations. However, the computational requirements for processing temporal 77 information go far beyond merely requiring a timer to discriminate or produce a single duration or 78 interval. Some forms of temporal processing require the ability to smoothly scale a time-varying 79 motor pattern. For example, the ability to play a song on the piano at different tempos, or catch a 80 ball thrown at different speeds, requires that the underlying patterns of neural activity unfold at 81 different speeds [12-15]. Indeed, some tasks in animal studies explicitly require animals to exhibit 82 temporal scaling: depending on context cues or training blocks animals must temporally scale their 83 motor response [14, 16-18]. In contrast, other timing tasks are categorical in nature, for example in the language domain phrasal boundaries are based in part on a categorical boundary of the pause 84 between phonemes—e.g., great eves x gray ties [19, 20], similarly, in the motor domain, the 85 86 distinction between a double-click and two single clicks of a computer mouse is categorical. Furthermore, in both the human and animal literature standard temporal bisection tasks require 87

subjects to make a two-alternative forced-choice categorical judgment regarding whether a
stimulus was short or long [21, 22].

90 It remains unclear if different computational requirements, such as the need to exhibit 91 temporal scaling or categorical timing, rely on similar or fundamentally different underlying neural 92 mechanisms to encode time. Consider a task in which an animal has to produce two intervalse.g., in response to two different sensory cues. Generally speaking, three encoding schemes could 93 94 allow the same network to produce these two different intervals: absolute timing, temporal scaling, 95 and stimulus-specific timing. Under *absolute* timing the neurons would respond at the same 96 moments in time during both the production of short and long intervals but additional neurons would be active during the long interval; in a *temporal scaling* scheme neurons encode the same 97 relative time during both short and long intervals; and in a stimulus-specific code, there would be 98 99 unrelated patterns for each interval (e.g., entirely different neural sequences for the short and long 100 interval). These different schemes possess specific computational tradeoffs regarding their 101 suitability for temporal scaling versus categorical timing.

102 To date, a large diversity of neural signatures for the encoding of time—including scaling, absolute timing, and stimulus-specific timing-have been observed during tasks that require 103 animals to discriminate or produce multiple intervals [14, 16-18, 23-30]. Here we propose that 104 105 some of this diversity is driven by task structure, and examine whether task structure influences 106 the way recurrent neural networks may encode time. To address this hypothesis we trained RNNs 107 on two tasks with identical output motor requirements and characterized how the networks encode 108 time and generalize to novel stimuli. Our results establish that subtle differences in task structure lead to neural dynamic regimes that are better suited for temporal scaling or categorical timing. 109

111 **Results**

To begin to understand how task structure might shape how time is encoded in neural networks, we trained recurrent neural network models (RNNs) on one of two tasks inspired by previous experimental studies[14, 18, 23]. The RNNs were based on firing rate units with distinct populations of excitatory (80%) and inhibitory (20%) units. We will refer to the tasks as the 2-Context (Fig 1A) and 2-Stimulus (Fig 1B) tasks—critically, the timed motor outputs were identical



Fig 1. RNNs were trained on one of two timing tasks, both of which required producing the same timed output patterns. (A) Schematic of the 2-Context task. Each RNN was composed of 200 units—80% excitatory units (purple) and 20% inhibitory units (dark red)—and received a *go* and a *context* input. The context level signals the interval length to be produced: high = long (6 s, blue), low = short (3 s, green). (B) Schematic of the 2-Stimulus task. The same RNN was used in both tasks, except that the short- and long-interval was cued by two different inputs that were transiently activated. (C) Learning curve for the performance of 20 RNNs trained on the 2-Context task. Percentage of trials in which the timing of the output unit met criteria (left) and the error between the output and target (right). Gray traces represent results of each RNN, red dots denote the end of training for a given RNN, and the black trace represents the mean performance. (D) Same as in (C) but for the 2-Stimulus task. (left). Mean crossing times for long interval is significantly higher than that for short interval (right, n = 20 simulations, paired t test, $t_{19} = 77.70$, P < 0.0001). Dashed lines denote the targets and threshold. (F) Same as (E) but for 2-Stimulus task (n = 20 simulations, paired t test, $t_{19} = 45.79$, P < 0.0001).

in both tasks, requiring the production of either a short or long response. In the 2-Context task [e.g.,
14, 18], the *Go* cue (500 ms) indicated the onset of the trial (t=0), and the analog level of a
continuous context input signaled whether a trial is short or long. In the 2-Stimulus task, the short
and long interval trials were cued by two distinct transient inputs [23]. In both cases, the short and
long intervals consisted of a ramp-up of the output unit starting at the interval midpoint—a function
that approximates the behavioral response rate of animals trained to correctly time their
movements [23].

Performance was quantified by the ratio of correctly timed trials (see Methods) and the error between the actual output and the target. RNNs trained on both tasks learned to produce the same appropriately timed motor output (Fig 1C-F), although the RNNs trained on the 2-Context task required fewer training trials to reach the same performance level (n = 20 simulations, twosample two-sided t-test, $t_{38} = 9.75$, P < 0.0001).

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131 Generalization to novel intervals

Having shown that RNNs can produce the same temporal output patterns when trained on 132 two similar tasks, we next asked a key question: are there significant functional differences 133 134 between how the RNNs trained on the different tasks perform in response to novel input conditions? 135 To answer this question we examined generalization to untrained input conditions. To test the generalization in the 2-Context task we varied the amplitude of the context cue between the range 136 of the trained values (0.75=short; 0.25=long). Interestingly the network exhibited fairly smooth 137 138 generalization-i.e., in response to intermediate context levels it produced intermediate motor 139 intervals (Fig 2A)—a finding consistent with previous computational studies [12, 14]. To test 140 generalization in the 2-Stimulus task we mixed the ratio of activation of the two stimulus cues-



Fig 2. RNNs trained on the 2-Context task exhibited smooth generalization to novel intervals, while RNNs trained on the 2-Stimulus task exhibited categorical timing. (A) Output traces of an RNN trained on the 2-Context task across different context input levels. Dashed-black lines denote the output threshold used to quantify timing. Pink squares denote the trained conditions. (B) Similar to (A) but for the 2-Stimulus task. The blue and green squares represent the ratio of activation of the two input units. (C) Plots of the mean crossing time for each RNN across input conditions for the 2-Context (top) and 2-Stimulus (bottom) tasks. Insets, examples of the sigmoid-function fits for a single RNN (black). (D) Left, mean slope of the sigmoid fits for 2-Stimulus task is significantly higher than that for the 2-Context task (n = 20 simulations for each, two-sided t test, $t_{38} = 9.69$, P < 0.0001). Right, correlation coefficient between mean crossing times and input conditions for 2-Context task is significantly higher than that for the 2-Stimulus task (n = 20 simulations for each, two-sided t test on Fisher-transformed values, $t_{38} = 17.39$, P < 0.0001). The absolute correlation coefficient values are shown because in the 2-Context task the correlations are negative. (E) Standard deviations of the crossing times for each RNN in the 2-Context (top) and 2-Stimulus (bottom) tasks, as a function of input conditions.

during training [1, 0] corresponded to short and [0, 1] to long, during testing an intermediary 50/50
mixed input corresponded to [0.5 0.5]. In contrast to the 2-Context task, the RNNs trained on the
2-Stimulus task did not generalize, but the RNNs did not exhibit catastrophic degradation or
behave randomly. Rather, the RNNs expressed categorical timing: the output intervals clustered
near the short or long intervals (Fig 2B), essentially exhibiting a winner-take-all behavior.

147 To quantify these generalization patterns we measured the slope of a sigmoid fit between input levels and output intervals, as well as the correlation between them (Fig 2C, D, see Methods). 148 149 The slope of the sigmoid was significantly less in the 2-Context fits—indicating a quasi-linear relationship between context input level and produced intervals. The sigmoid slope was 150 151 significantly higher in the 2-Stimulus task, consistent with the prototypical signoidal signature of 152 categorical discrimination (Fig 2D, left panel). Similarly, the Pearson correlation coefficients 153 further supported the observation that the input-interval relationship was much more linear in the 154 2-Context task compared to the 2-Stimulus task (Fig 2D, right panel).

155 In addition to the above accuracy measures, we also quantified the precision of timing 156 across the different generalization conditions, as the standard deviation of the crossing time of 157 each trial (Fig 2E). The precision for the 2-Context task was high (low standard deviation) for all 158 the stimulus conditions. In contrast, in the middle range for the 2-Stimulus task precision was very 159 low. This was mainly due to categorical timing, i.e., in some stimulus conditions, the motor output 160 would randomly be attracted towards the short or long interval. Taken together, RNNs trained on the 2-Context task were far superior at generalizing to novel intervals in terms of both timing 161 162 accuracy and precision, however, the RNNs trained on the 2-Stimulus task exhibited categorical timing. 163

164 By design, the key difference in the tasks is that in the 2-Context task there is a continuous 165 input signaling the target interval throughout the task, whereas in the 2-Stimulus task two different input weight vectors signal the desired interval, and each of these inputs is only active for a brief 166 167 period. To further determine whether the difference of the generalization patterns is robust to the 168 input parameters, we manipulated the 'similarity of the inputs corresponding to the short and long 169 intervals in both tasks. Specifically, for the 2-Context task, different analog pairs of context level 170 were used, ranging from (0.95, 0.05) to (0.55,0.45). For the 2-Stimulus task, we gradually 171 increased the similarity by increasing the overlap ratio between the two inputs—proportions of the same elements in the two input weights (S1 Fig A, B). In all five conditions, the generalization 172 173 performance for the 2-Context task was better than that for the 2-Stimulus task (S1 Fig C, D, and 174 E). While the tasks were designed to capture features of those used in behavioral experiments [14, 175 18, 23], in the 2-Context task the onset of the Go and Context stimuli redundantly signal trial onset 176 (t=0). Thus to understand the influence of the Go stimulus we also performed simulations without 177 the Go stimulus in the 2-Context task (S2 Fig A). As expected, omitting the Go stimulus left the 178 generalization performance largely unchanged compared to the standard 2-Context task with Go 179 stimulus, and still significantly better than that for the 2-Stimulus task (S2 Fig B). Finally, to confirm that it is the presence of the continuous context input that plays a critical role in the 180 181 differential generalization patterns, we performed "2-Context" simulations in which the short and 182 long intervals were cued by a transient "context" stimulus rather than a persistent context input. Consistent with our expectations based on previous results [12, 14, 30], in the absence of a 183 184 continuous context input the generalization was more consistent with categorical timing (S3 Fig) Additional simulations confirmed that the difference of the generalization performance 185 186 between the 2-Context task and 2-Stimulus were robust to the change of several hyperparameters

including the initial gain (S4 Fig A-C) and connection probability (S5 Fig A-C) of the recurrent
weights.

189 Potential dynamic regimes underlying the encoding of multiple 190 intervals

191 Converging experimental and theoretical evidence indicates that a broad range of neural 192 dynamic regimes encode time. But to date, these different regimes have not been contrasted in 193 terms of their ability to encode multiple intervals and lead to generalization or categorical timing, 194 or robustness to noise. Here we examine three broad potential strategies for the encoding of two 195 intervals: scaling, absolute, and stimulus-specific codes. To illustrate these three strategies we 196 consider how a network of neurons could encode both a short (3 s) and long (6 s) intervals (Fig 197 3)—note that while we use neural sequences to contrast the three encoding schemes, the same 198 classification applies to other codes for time, including ramping activity. In a temporal scaling 199 strategy (Fig 3A), the dynamics of each unit for the short interval is linearly scaled in time to 200 produce the long interval (Fig 3B), which at the level of single units leads to two overlapping 201 curves (Fig 3C). Similarly, when the neural trajectories of the entire population are projected into 202 a low-dimensional space by principal component analysis the trajectories are also overlapping (Fig. 203 3D). Under an absolute encoding strategy (Fig 3, middle panels) the temporal profile of each unit 204 during the short interval does not change during the long interval. The long interval simply relies 205 on recruiting additional neurons that have later temporal fields. Thus in PCA space, the curves for 206 the short interval matched the first half of that for the long interval. In a stimulus-specific strategy 207 (Fig 3, right panels), the temporal profile of each neuron is essentially uncorrelated during the 208 short and long intervals. Thus in PCA space, the trajectories of the neural patterns of activity 209 produced during the short and long intervals are distinct from one another.

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Fig 3. Three strategies for the encoding of two intervals by the same group of neurons. (A) Schematic of three potential strategies for timing two intervals: scaling, absolute, and stimulus-specific from left to right. (B) Prototypical dynamics for each of the encoding schemes for a population of units during production of the short (top) and long (bottom) intervals. (C) Activity traces of the units denoted by the red arrows in (B) for short (blue) and long (green) intervals. (D) Trajectories of three PCA components for short (cyan-blue) and long (yellow-green) interval for the corresponding population dynamics. The gradient colors (from the light to the dark) denote the flow of time. Circles denote the time points of the 1st, 2nd, 3rd, 4th, 5th, and 6th seconds.

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also possible that the dynamics of a given unit exhibit an absolute code early in a trial followed by

are best described as scaling from one interval to another, while others encode absolute time. It is

scaling later in the trial. Note, however, that it would not make sense to consider a case in whicha unit undergoes scaling early in a trial and then exhibits absolute timing.

We next describe how to quantify these three schemes both at the level of the neural
population and of individual neurons in RNNs trained on either 2-Context or 2-Stimulus tasks.

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Task structure differentially shapes the time encoding strategies at the population level

222 In order to visualize the internal dynamics of the RNNs we first plotted the normalized activity observed during the short and long intervals sorted according to the latency of peak activity 223 224 for each unit during the short interval (Fig 4A-B, left panels), and sorted by the long intervals (Fig 225 4A-B, right panels). Interestingly, although the target output was a ramping pattern, relatively few 226 RNN units appeared to be ramping. Rather, the global activity patterns in both tasks might be best 227 conceptualized as neural sequences. Yet, while the self-sorted sequences appeared to be visually 228 similar for both tasks, the cross-sorted sequences were dramatically different. Specifically, in the 229 2-Context task it appeared that neurons fired in the same order for both the short and long 230 intervals-suggestive of a scaling encoding strategy. However, in the 2-Stimulus task the cross-231 sorted PSTHgrams revealed a more complex relationship between the spatio-temporal patterns of 232 activity during the short and long intervals—suggestive of a more stimulus-specific encoding 233 strategy.

To quantify if the neural dynamics observed in the 2-Context and 2-Stimulus tasks were more consistent with a scaling, absolute, or stimulus-specific code, we first developed as stimulusspecific index (SSI_{pop}) based on previously described geometric approaches [12, 30, 31]. We started with the cross-Euclidean distance matrix between population dynamics for short and long



Fig 4. Distinct population dynamics in RNNs trained on the 2-Context and 2-Stimulus task. (A) Population activity for short (top) and long (bottom) intervals sorted according to the peak activity latency during short (left) and long (right) intervals for RNNs trained on the 2-Context task. (B) Same as A for the 2-Stimulus task. (C), (D), (E) Schematic of the calculation of the stimulus-specific index (SSI_{pop}). A prototypical neural sequence that undergoes pure temporal scaling from the short (top) to long (bottom) intervals is used as an example (C). The vectors of the pairwise time points from the short and long dynamics are used to calculate all pairwise Euclidean distances, and these pairwise distances comprise the cross-distance matrix (**D**), in which a row (e.g., blue rectangle) represents the distances between one column vector of short dynamics and all column vectors during the long dynamics. The minimal index vector (red vectors in (D) and (E)) represents the indices along the x-axis that corresponds to the minimum distances for each row of the cross-distance matrix (red squares). A series of reference vectors that vary from pure scaling to pure absolute timing (black vectors) are compared to the minimal index vector, and a value τ_{min} is defined as the τ at which the pairwise distance reaches the minimum. Finally, the correlation coefficient between the minimal index vector and the absolute-scaling reference vector at τ_{min} is used to calculate SSI_{pop}. (F) Cross distance matrices for an example simulation of the 2-Context (left) and 2-Stimulus tasks (right). Red lines denote the indices of the minimum values for each row. (G) SSIpop for RNNs trained on the 2-Stimulus task is significantly higher than that for 2-Context task (n = 20 simulations for each, two-sided Wilcoxon rank-sum test P < 0.0001). Boxplot: central lines, median; bottom and top edges, lower and upper quartiles; bottom and top whiskers: extremes.

239 intervals (see Methods), which compares the similarity of the activity across all time pairs during 240 the short and long intervals (Fig 4C-D, example based on a case of perfect scaling of the entire population). We then extracted the index (time bin of the long interval) corresponding to the 241 242 minimum value along each row of the cross-time distance matrix (red square in Fig 4D), which 243 results in a vector of the time points that in the long-interval that are closest to each of the time points in the short-interval: the minimal index vector (red row vector in Fig 4D and column vector 244 245 in Fig 4E). This minimal index vector was then matched to all possible reference vectors 246 representing perfect scaling codes to a perfect absolute code (black column vectors in Fig 4E) by computing the distances d_{τ} between each pair (Fig 4E). The reference vector with the minimum 247 distance $(d_{\tau min})$ to the minimal index vector denoted the best absolute-scaling vector. The 248 249 correlation (c_{tmin}) between the best absolute-scaling vector and the minimal index vector determines how good the match is: 1.0 reflects perfect scaling, absolute timing, or a perfect mixture 250 251 of absolute and scaling code. However, the correlation will be low or even negative in the case of 252 a stimulus-specific code. Therefore, SSIpop was defined by 1-c_{tmin} (Fig 4E), meaning that both perfect scaling and absolute timing would result in an SSIpop=0, and the stimulus-specific code 253 254 would be proportional to SSI_{pop}.

We calculated SSI_{pop} for all 20 RNNs in both the 2-Context and 2-Stimulus tasks. SSI_{pop} was significantly higher during the neural dynamics of the 2-Stimulus task compared to the 2-Context task (Fig 4G), indicating that dynamics observed during the 2-Stimulus task reflected a stimulus-specific encoding strategy more so than the 2-Context task. However, consistent with the visual inspection of the dynamics and distance matrices (Fig 4A, F), it is clear that the 2-Stimulus task was not entirely accounted for by a stimulus-specific strategy, suggesting a mixed code. Thus we next examined the three encoding strategies from the perspective of the individual units in thenetwork.

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Task structure shapes timing encoding strategy at the level of single

265 units

To understand whether the encoding of the short and long intervals was most consistent 266 267 with a scaling, absolute, or stimulus-specific code at the level of single units, we used a previously 268 described measure of absolute-versus-scaling index (ASI) [23], and incorporated a novel stimulus-269 specific index (SSI_{unit}) into the framework. Much as SSI_{pop} quantifies how different the dynamics 270 of two neural populations are, SSI_{unit} quantifies how different the firing-rate profiles of a unit are 271 during a short versus long trial (see Methods). More specifically, for a given unit, a high SSIunit 272 implies the temporal profiles during two trials are not related to each other through scaling, 273 absolute timing, or a mixture of both with the absolute part followed by the scaling part. A low 274 SSI_{unit} implies that the temporal profiles are related through scaling, absolute timing, or a mixture 275 of both, thus justifying the use of the ASI to further quantify scaling versus absolute timing. To 276 calculate the SSI_{unit} we first time-warped the temporal profile of a unit during the long interval into 277 a series of reference absolute-scaling traces spanning from pure scaling to pure absolute timing 278 with a mixture of both in between (Fig 5A). These reference traces were defined by a "breaking 279 point" τ marking the transition from absolute timing to scaling (τ =0 reflects perfect scaling and τ =T_{short} reflect absolute timing). All reference traces were compared with the short dynamics by 280 computing the Euclidean distance at each τ (d_{τ}). The reference trace with the minimum distance 281 $(d_{\tau min})$ denoted the best match with the actual temporal profile of the unit. Finally, as with SSI_{pop}. 282 283 the SSI_{unit} was defined as 1.0 minus the correlation between the temporal profile during the short

intervals and the reference trace at τ_{min} ($c_{\tau min}$). For a given unit with a low SSI_{unit} (≤ 0.5), we went on to calculate its ASI which is also based on τ_{min} (see Methods). With the SSI_{unit} and ASI in hand, we classified a given unit as either a stimulus-specific unit (SSI_{unit}>0.5), a scaling unit (SSI_{unit} ≤ 0.5 , ASI ≤ 0.5), or an absolute unit (SSI ≤ 0.5 , ASI>0.5) (Fig 5B).

288 This approach allowed us to classify each unit of the network and contrast the distribution 289 of temporal classifications between the 2-Context and 2-Stimulus tasks. These analyses revealed 290 that RNNs exhibit a mixed encoding strategy, exhibiting a broad range of scaling, absolute, and 291 stimulus-specific units (Fig 5C). However, there were highly significant differences in the 292 distributions of temporal classes between the RNNs trained on the 2-Context and 2-Stimulus tasks 293 (Fig 5D). The 2-Context RNNs were dominated by scaling units, while 2-Stimulus RNNs had 294 more stimulus-specific units. The results partially explain why 2-Context RNNs were better at 295 generalizing to novel intervals. Because our RNN structure obeyed Dale's law it was possible to 296 contrast the encoding strategies of excitatory and inhibitory neurons. Interestingly the distribution 297 of scaling, absolute, and stimulus-specific cells appeared similar between excitatory and inhibitory 298 neurons (Fig 5D).

299 To establish a causal relationship between the distribution of temporal classes to the functional properties of the RNNs we selectively deleted units of different classes from the RNNs 300 301 trained on both tasks (S6 Fig A). We then investigated how the performance changed in response 302 to these deletions. Performance and error across six deletion manipulations (stimulus-specific, 303 scaling, and absolute temporal-classes for the excitatory and inhibitory populations) revealed 304 inhibitory scaling units more severely impaired RNN function (S6 Fig **B**, **C**) for the 2-Context task. 305 In contrast, no single manipulation condition more severely affected both performance and error 306 in the 2-Stimulus task (S6 Fig **D**, **E**). Somewhat surprisingly these results reveal that in the case of 307 the 2-Context task a single subtype of inhibitory neurons—those that were classified as scaling 308 units—are the most critical for network dynamics and encoding time. Whereas in the 2-stimulus 309 task the coding strategy can be considered to be truly mixed, in the sense that all temporal classes 310 and excitatory-inhibitory neurons seem to contribute more or less equally to the underlying 311 dynamics and the encoding of time.



Fig 5. Different distribution of stimulus-specific, scaling, and absolute units between the 2-Context and 2-Stimulus tasks. (A) Schematic of the definitions of the stimulus-specific index (SSI_{unit}) and absolute vs. scaling index (ASI) at the single unit level. Consider a hypothetical firing rate profile of a unit during a short (blue, x(t)) and long (green, y(t)) trial. As described in Methods, a series of time-warped long dynamics are generated at breaking point τ_x : before τ_x the dynamics are the same during both the short and long intervals (absolute timing, $y^{abs}(t)$; after τ_x the dynamics is the scaled version of the corresponding original long dynamics (> τ_x , scaling timing, yscale(t')). Pairwise Euclidean distance between short dynamics and all time-warped long dynamics are computed at each τ_x . The point at which the distance is minimal defines τ_{min} and is used to compute the SSI_{unit} as in SSI_{pop}. To compute the ASI, a normalized measure of the distance before and after τ_{min} is calculated (AbsR) as in described in Methods to quantify the weighting factor for the absolute part (before τ_{min}) and the scaling part (after τ_{min}). ASI is defined by τ_{min} and the weighting factor based on AbsR(τ_{min}). (B) For a given unit, the SSI_{unit} is computed first, and if the SSI_{unit} is higher than 0.5, it is classified as stimulus-specific unit. If the SSI_{unit} is lower than 0.5, its ASI is computed, and it is classified as scaling unit if its ASI is lower than 0.5, otherwise as an absolute unit. (C) Dynamics of five example unit traces for short (blue) and long (green) intervals for the 2-Context (top) and 2-Stimulus (bottom) tasks, the corresponding SSI and ASI values are shown on top. Notice that for a given unit, ASI is only computed only when its SSI_{unit} is lower than 0.5. (**D**) For the 2-Context task (left), most units are classified as scaling units—for both excitatory and inhibitory units (n = 20 simulations, two-way ANOVA with 0.0001). For the 2-Stimulus task (right), stimulus-specific units are the most common (n = 20 simulations, twoway ANOVA with repeated measures, $F_{(2, 38)} = 181.5$ and P < 0.0001, posthoc Tukey tests P < 0.0001).

313 Task structure differentially shapes the relationship between 314 recurrent dynamics and input/output space

After quantifying how the different task structures shaped the encoding strategies, we 315 316 sought to determine if the differences can be understood in terms of the relationship between RNN 317 dynamics and the input/output subspaces. Generally, recurrent dynamics is driven by two sources: the interaction between the inputs and input weights, and between recurrent activity and recurrent 318 319 weights. To start to understand how the inputs affected the recurrent dynamics and how the 320 recurrent dynamics would lead to the output through the output weights, we first performed the 321 principal component analysis on the concatenated dynamics of both intervals for each task (S7 Fig 322 A, B)—the first three PCs for the 2-Context task explained more variance than that for the 2-323 Stimulus task (88.15±0.75% vs 69.72±0.73%, S7 Fig C). We then projected the recurrent dynamics into the low dimensional space spanned by the first three PCs (S7 Fig A, B). Visually in 324 325 PC space, the dynamics of the two intervals for 2-Context task orbited close to each other, while 326 that for the 2-Stimulus task formed two distinct trajectories—consistent with our findings that 2-327 Context task tended to use an absolute-scaling strategy while 2-Stimulus, a stimulus-specific 328 strategy. These observations were further established by plotting the dynamics in response to 329 generalization conditions (Fig 2). In the 2-Context task the dynamics across different inputs 330 smoothly transitioned to nearby trajectories, while in the 2-Stimulus task the trajectories clustered 331 around the two trained (short and long) trajectories (S8 Fig).

To directly compare the relationship between the recurrent dynamics across time and the input/output weights, we projected the input weights—Input_{Go} and Input_{Context} for the 2-Context task, Input_{Short} and Input_{Long} for the 2-Stimulus task—and the output weights into the same PC space. We then computed the pairwise angles between the projected input/output vectors and each

336 segment vector of recurrent dynamics across time (see Methods) (S7 Fig A) for both tasks. 337 Interestingly, for the 2-Context task the dynamics of both intervals first evolved in the Input_{Go} input direction as revealed by the small angle for the first 2 segments. After that, both trajectories 338 339 stayed in a plane almost orthogonal to the Go input till the end of the trial. The dynamics were 340 almost orthogonal to the Input_{Context} at the beginning (with angles close to 90 degrees) and then the 341 angle decreased in the middle period and increased again to about 90 degrees at the later period. 342 Finally, for output weights, the angle stayed close to 90 degrees at the beginning then it decreased 343 to a low level till the end of the trial indicating that the dynamics followed the output weights 344 directions in the later period of the trials to better generate the target ramp staring at the middle 345 point of each trial (S7 Fig **D**).

For the 2-Stimulus task, the dynamics of short and long intervals started to follow their corresponding input directions and then went to the opposite directions after input offset and stayed almost orthogonal thereafter. While for the output weights, the angle started at around 90 degrees and then decreased around the start point of the target ramp then it increased at the end of the trials to the opposite direction (S7 Fig E).

351

Task structure differentially shapes the learned recurrent synaptic connectivity

Ultimately the task-specific differences in RNN dynamics must be attributed to differences in input structure and the recurrent weight matrix. Thus we next characterized the relationship between the recurrent weight matrices and performance. Since our RNNs respected Dale's law, we grouped weights into the four standard subtypes: all excitatory to excitatory unit connections ($E \rightarrow E$), all excitatory to inhibitory unit connections ($E \rightarrow I$), all inhibitory to excitatory unit connection (I \rightarrow E), and all inhibitory to inhibitory unit connections (I \rightarrow I). We then completely deleted each group of synapses and quantified the change in output performance (Fig 6A).



Fig 6. Differential connectivity patterns in RNNs trained on the 2-Context and 2-Stimulus tasks (A) Example of the effects of deleting entire subgroups of synapses on performance in the 2-Context (top) and 2-Stimulus (bottom) tasks. From left to right, example output traces of the short (blue) and long (green) intervals for the control condition, and after deleting all excitatory unit to excitatory unit connections (Delete $E \rightarrow E$), all excitatory unit to inhibitory unit connections (Delete $E \rightarrow I$), all inhibitory to excitatory unit connections (Delete $I \rightarrow E$), and all inhibitory unit to inhibitory unit connections (Delete $I \rightarrow I$). (B) Mean performance (left) and error (right) of the outputs corresponding to the conditions in panel A. The performance for the Delete $E \rightarrow E$ condition is significantly lower than the control but significantly higher than the other conditions in 2-Context task. For the 2-Stimulus task performance for Delete $E \rightarrow E$ was not significantly worse than the control, but significantly higher than the other conditions (two-way ANOVA with mixed-effect design, $F_{4,152} = 823.9, P < 1000$ 0.0001, posthoc Tukey tests P < 0.0001). The error for Delete I \rightarrow E condition is significantly higher than the other conditions in both 2-Context and 2-Stimulus task (two-way ANOVA with mixed-effect design, $F_{4,152}$ = 39.8, P < 0.0001, posthoc Tukey tests P < 0.0001). (C) Left, connection probability in the 2-Context task was significantly higher than in the 2-Stimulus task(two-way ANOVA with mixed-effect design, $F_{1,38} = 338.3$, P < 5000.0001 for the task factor). Probability for the I \rightarrow E connections is significantly higher than that for the other three conditions: $E \rightarrow E$, $E \rightarrow I$, $I \rightarrow I$ in both 2-Context and 2-Stimulus task ($F_{3,1/4} = 2884$, P < 0.0001 for the connection factor, posthoc Tukey tests P < 0.0001). Right, the mean weight in the 2-Context task is significantly lower than that in the 2-Stimulus task (two-way ANOVA with mixed-effect design, $F_{1.38} = 219.1$, P < 0.0001 for the task factor). Probability for the I \rightarrow E connection is significantly higher than that for the other three conditions: $E \rightarrow E$, $E \rightarrow I$, $I \rightarrow I$ in both 2-Context and 2-Stimulus task ($F_{3,1/4} = 183.7, P < 0.0001$ for the connection factor, posthoc Tukey tests). **** = P < 0.0001, and ** = P = 0.002.

362 Interestingly, deleting all $E \rightarrow E$ connections only slightly affected the performance and error for 363 both tasks, while deleting all other three groups decreased the performance or increased the error. Deleting the $I \rightarrow E$ connections produced the largest change in error (Fig 6B). We next quantified 364 365 the connection probability and mean weights of each group (Fig 6C). Consistent with the 366 performance and error results, $I \rightarrow E$ connections exhibited the highest connection probability and 367 mean weights for both tasks. Interestingly, to achieve similar output performance, the two tasks 368 seemed to rely on different strategies in the structural level: 2-Context task favored higher 369 connection probability, while 2-Stimulus task preferred higher mean weights (Fig 6C).

370

371 RNNs trained for the 2-Stimulus task are more robust to noise

372 We have seen that RNNs trained for the 2-Context task are better suited for generalization 373 to novel intervals and this feature is related to the underlying dynamics being governed by a 374 absolute-scaling encoding scheme. A question that emerges from these results is whether there is 375 a computational tradeoff between the distinct dynamic regimes observed in both tasks? For 376 example, while the RNNs trained on the 2-Context task exhibit better generalization, do they 377 perform worse on any other measures? As a first step to address this question we analyzed the 378 robustness of both tasks in response to noise. In the brain, of course, neural networks are 379 continuously subject to extraneous noise, and thus robustness to noise imposes an important 380 constraint on biologically functional dynamic regimes [32]

As above we first trained RNNs on either the 2-Context and 2-Stimulus tasks with the standard settings, namely noise level of 0.45 (σ in Eq. 1), then we tested the networks by applying different values of σ . Example output traces for the 2-Stimulus task under all noise levels tested were less scattered than that for the 2-Context (Fig 7A). This was supported by the fact that the

385 mean error for the 2-Stimulus task was lower than that for the 2-Context (Fig 7B). For both tasks, at high noise levels, there were some incorrect trials (< 10% and no significant difference between 386 387 the two tasks) in which either the output never crossed the threshold during the trial or crossed the 388 threshold outside of the acceptance windows We then directly contrasted the temporal precision 389 of the correct trials and found that the standard deviations for the 2-Stimulus task were lower than that for the 2-Context task (Fig 7C). Taken together, we conclude that the dynamic regimes 390 underlying timing in the predominately stimulus-specific dynamics that emerged in the 2-Stimulus 391 task provided a computational benefit in terms of robustness to noise suggesting computational 392 393 tradeoffs between different dynamic regimes for the encoding of time.



Fig 7. RNNs trained on the 2-Stimulus task were less sensitive to noise perturbations. (A) Output traces for short (blue) and long (green) intervals from an example RNN trained on the 2-Context (left) and 2-Stimulus (right) across different levels of noise (σ) during testing. (B) Mean error (across 50 trials) for 2-Context task (cyan) is higher than that for 2-Stimulus task (orange) (n = 20 simulations, two-way ANOVA with mixed-effect design, $F_{1,38} = 9.35$, P = 0.004). (C) Mean standard deviation of the time of threshold-crossing across all correct trials for 2-Context task (cyan) is higher than that for 2-Stimulus task (orange) ($F_{1,38} = 341$, P < 0.0001). Data are presented as mean \pm SEM.

Similar to the generalization performance, the difference of the robustness to noise between
the 2-Context and 2-Stimulus tasks was consistent across different input parameters (S1 Fig F),

initial gain (S4 Fig **D**), and connection probability (S5 Fig **D**) of the recurrent weights.

398

399

400 **Discussion**

401 Here we trained supervised RNNs on two simple temporal tasks that required the production of 402 identical temporal output patterns based on previous behavioral results [14, 18, 23]: a ramping 403 increase in output firing rate that peaked after either a short (3 s) or long (6 s) interval. The tasks differed only in how the short and long intervals were cued: either by a continuously presented 404 405 context input (2-Context task) or by two distinct brief inputs (2-Stimulus task). In principle the 406 same dynamic regimes could have emerged and solved both tasks, yet, significantly different 407 dynamic regimes emerged in the different tasks. Thus depending on the task RNNs encoded time 408 in different ways, and exhibited fundamentally different computational properties, particularly 409 regarding how the networks generalized to novel stimuli.

410

411 Neural dynamic regimes of population clocks

A converging body of experimental and computational evidence suggests that neural circuits encode time in spatiotemporal patterns of neural activity. Two experimentally relevant neural dynamics regimes by which neurons can encode time include ramping activity and population clocks. Ramping codes generally refer to monotonically increasing (or decreasing) firing rates throughout an interval [24, 33-40]—in ramping codes firing rate often peaks at the time of the target interval, and in principle, a single neuron can encode time throughout the entire duration. Population clocks refer to time-varying patterns of activity in which time is encoded in
the population activity of neurons, which generally exhibit nonmonotonic changes in firing rate,
and importantly these dynamics are generated by the recurrent connectivity within a neural circuit
[1, 11, 41, 42]. Population clocks can include simple sparse neural sequences as well as complex
spatiotemporal patterns in which a given neuron can exhibit multiple time fields [28, 43-51].

423 In the current simulations, the target output patterns were a simple ramping pattern, yet 424 most of the units in the RNNs were not well described as ramping units—even though it seems 425 that this would be the simplest and most direct solution to solve the tasks. Rather, the neural dynamics observed in the RNNs studied here, are most consistent with the notion of population 426 427 clocks in general and neural sequences in particular (Fig 4). These results are in line with other 428 computational models in which neural sequences encode time [52-55]. The reason RNNs trained 429 with supervised learning rules seem to converge to neural sequences rather than ramping activity 430 are not well understood, but it has been recently proposed that neural sequences represent a fairly 431 optimal encoding scheme for downstream neuron (the output unit in our case) to read out time 432 [23].

433

434 Absolute, scaling, and stimulus-specific codes

We outlined three general temporal encoding strategies by which a population of neurons could solve temporal tasks that require producing multiple intervals (Fig 3)—such as the two tasks examined here. The scaling strategy is perhaps the most intuitive because it essentially exploits the same neural dynamics to produce both a short or long interval by altering the speed at which the dynamics unfold. Indeed, such scaling has been observed experimentally [14, 16, 23, 26, 38, 56, 57]. Neurons that exhibit absolute timing have also been experimentally observed, along with 441 neurons that categorically detect the midpoint boundary between short and long intervals [14, 23, 442 26, 27, 56-61]. Stimulus-specific codes in which the same or different intervals can be encoded in 443 different neural trajectories have also been described [17, 47, 58, 62-64]. To date, however, these 444 different encoding strategies have not been carefully analyzed or quantified. To this end, we 445 described two general purpose quantitative measures—the ASI and SSI_{unit}—that can be applied 446 across a wide range of single-unit data and used to classify neural responses.

447 These measures revealed a different distribution of unit types across the RNNs trained on the 2-Context and 2-Stimulus tasks (Fig 5). Specifically, over 50% of the units in the 2-Context 448 RNNs were classified as scaling units, whereas in the 2-Stimulus RNNs over 50% were classified 449 450 as stimulus-specific units-that is, their temporal profiles between the short and long interval were 451 not consistent with either absolute or scaling coding strategies. This differential distribution is 452 consistent with the intuition that because in the 2-Context task the context input is active during 453 both the short and long intervals, and a stimulus-specific encoding strategy is more difficult to 454 implement compared to the 2-Stimulus task—i.e., the input space of the 2-Context task is smaller. 455 Put another way, in the 2-Stimulus task RNNs are likely to begin their trajectories at the beginning 456 of each trial (t=0) in more distant regions of neural state space than in the 2-Stimulus task.

The differential distribution of scaling, absolute, and stimulus-specific neurons accounts in part for the distinct computational features of both types of networks. Specifically, the classification of units into different temporal coding strategies allowed us to demonstrate that selectively deleting some classes impaired RNN performance more than others. Deleting a few inhibitory scaling units impaired RNN performance in the 2-Context task significantly more than deleting absolute or stimulus-selective units. In contrast in the 2-Stimulus task, all classes 463 contributed to performance with an approximately equal weighting—reflecting a much more464 mixed encoding strategy [65, 66].

465

466 **Computational trade-offs between time-encoding dynamic regimes**

The 2-Context and 2-Stimulus tasks required producing the same temporal output patterns 467 468 but generated dramatically different behaviors when challenged with novel inputs. Of particular 469 relevance was that in response to novel levels of activation of the inputs, the 2-Context RNN 470 exhibited a smooth scaling of the temporal profile of the output. In this task, in response to the go 471 stimulus, RNN's generated a neural trajectory that resembled a neural sequence. Depending on the 472 analog value of the context input this trajectory unfolded at either a slow or fast speed to produce 473 the short or long interval, respectively. Critically, in response to novel levels of activation of the 474 tonic context input the velocity of the neural trajectory varied smoothly-thus generating smooth 475 temporal scaling of the output pattern. This same property has been observed in numerous other 476 models of timing [12, 30, 34, 67-69]. Specifically, a single input or variable is able to modulate 477 the velocity of the RNN dynamics in an approximately linear fashion.

478 In contrast to the temporal scaling behavior observed in the RNN trained on the 2-Context 479 task, when the 2-Stimulus RNNs were tested with inputs they were not trained on (e.g., 50% Input 480 1 + 50% Input 2) they did not exhibit smooth generalization. Importantly, they also did not exhibit 481 catastrophic degradation-i.e., the internal dynamics was robust to very different initial states. Rather they exhibited categorical timing-essentially a winner-take-all competition between two 482 483 distinct trajectories. This result is consistent with the notion that RNNs can encode multiple neural 484 trajectories in regimes that have been referred to as dynamic attractors [31, 70], locally stable 485 transient trajectories [71, 72], or stable heteroclinic channels [73, 74]. Here, two trajectories

possess their own basins of attraction (or "rivers-of-attraction") which lead the activity of thenetwork into one or the other of the two dynamic attractors.

Both temporal scaling and categorical timing are behaviorally relevant forms of timing. Specifically, some tasks require smoothly scaling the temporal output patterns, while others require categorically discriminating or producing one of distinct two intervals [12-15, 21, 22]. Thus, we have shown that the population clocks that emerge in RNNs can account for both temporal scaling and categorical timing and that it is possible to distinguish between both regimes based on the percentage of units that undergo scaling or stimulus-specific timing.

It is also relevant to note that RNNs learned to solve the 2-Context task in fewer training trials than the 2-Stimulus task. This may be because it is easier to adjust weights to generate a single trajectory at two different speeds than to generate largely distinct trajectories. Furthermore, during training the 2-Context task the RNN is always subject to tonic external input which in effect might facilitate learning by suppressing the potential emergence of chaotic regimes [75, 76].

499

500 **Experimental predictions**

501 As is evident from the behavioral data, a wide range of distinct neural regimes, from 502 ramping activity to a diverse range of neural population clocks, have been observed experimentally 503 across different brain areas and behavioral tasks [for reviews see: 1, 6, 7, 10]. Here we show that the same is true even in RNNs trained on two tasks that require the production of the same temporal 504 505 output patterns. Our results thus suggest that much of the experimentally observed variability 506 might be accounted for by relatively subtle differences in task structure. Furthermore, because 507 most timing tasks used in laboratories tap into ecologically relevant behaviors, different tasks may 508 encourage generalization patterns that best approximate their ecological relevance. These distinct generalization patterns will, in turn, result in time being encoded in different dynamic regimes—
e.g., regimes that are well-suited for temporal scaling or categorical timing.

A number of strong experimental predictions emerge from our results. First, at the 511 512 behavioral level, we predict that whether rodents are trained on the 2-Context or 2-Stimulus will 513 lead to different generalization patterns to novel stimuli. For example, a single odor along with a 514 tone context stimulus could be used for the 2-Context task, and two brief odors as the stimuli in 515 the 2-Stimulus task. We predict that changing the loudness of the tone in the 2-Context task will 516 scale the output pattern, but mixing the odors will result in categorical timing rather than the production of an intermediary interval. Second, we predict that neural recordings from animals 517 518 trained on these tasks will exhibit specific neural dynamic signatures, i.e., in the 2-Context task 519 more neurons will be categorized as scaling units compared to the 2-Stimulus task. Of course, one 520 must take into account that results may be dependent on the brain areas being recorded. However, 521 based on the current literature we expect this prediction to hold in those areas that have been 522 implicated in timing across many tasks, including the striatum, supplementary/secondary motor 523 areas, and prefrontal cortical areas.

525 Materials and Methods

526 Firing-rate RNN model

527 RNNs were based on firing-rate units that obeyed Dale's law (N = 200, 80/20%528 excitatory/inhibitory). RNN dynamics was described by the following equations:

529

530
$$\tau \frac{dx}{dt} = -x + W^{rec} * r + W^{in} * I + \sigma * N(0,1) * \sqrt{2 * \tau}$$
(1)

531

 $o = W^{out} * r \qquad (2)$

533

534
$$r = \min(ln(1 + e^x), 20)$$
 (3)

535

where $\mathbf{x} \in \mathbb{R}^{N \times 1}$ represents the input currents of RNN units, and firing rate vector **r** is obtained by 536 537 applying a Softplus function constrained by an upper bound of 20. The time constant τ was equal to 100 ms for all units. $\mathbf{W}^{in} \in \mathbb{R}^{N \times 2}$ and I are the input weights and external inputs, which are task-538 539 specific as described below. Each unit received independent Gaussian noise N(0,1) with the standard deviation of $\sigma\sqrt{2\tau}$. Unless otherwise specified, $\sigma = 0.45$. $\mathbf{W}^{\text{rec}} \in \mathbb{R}^{N \times N}$ is the recurrent 540 541 weight matrix. Self-connections were absent in the network. The output (o) of the network is computed linearly from the output weights Wout and r. RNNs were implemented and trained in 542 543 Tensorflow starting from the code of Kim et al [77, 78].

544 *Training.* Networks were trained using adaptive moment estimation stochastic gradient 545 descent algorithm (Adam) to minimize the error between network output o and target z:

547
$$Error = \sqrt{\sum_{t=0}^{T} [o(t) - z(t)]^2}$$
(4)

548

where T is the total length of a given trial. The target and mask are task-dependent as describedbelow. The learning rate was 0.01, and other TensorFlow default values were used.

Only recurrent weights W^{rec} and output weights W^{out} were trained. Unless otherwise 551 552 specified, W^{rec} was initialized as a random sparse matrix with a connection probability of 0.2 from 553 a normal distribution with zero mean and standard deviation (gain) of 1 and transformed to 554 absolute values. To begin in an approximately balanced regime the inhibitory weights were 555 multiplied by 4 for the initialization but not for training. To respect Dale's law during training a rectified linear operation was applied on W^{rec} to clip the weights down to zero and then excitation 556 557 and inhibition were implemented by multiplying the clipped W^{rec} with a diagonal matrix of 1 and -1 representing excitatory and inhibitory units, respectively [78, 79]. Wⁱⁿ was drawn from a 558 559 standard normal distribution and was fixed during training.

560 During training, a discretization step of 20 ms was used. After training, RNNs were ported 561 to Matlab using the trained parameters and a discretization step of 1ms was used to get the 562 dynamics for analyses.

Parameters were updated every trial. After every 100 trials of training, the network was tested for 100 trials to compute the task performance (see below) and mean error. When task performance was higher than 97% and the mean error is lower than 2, the training was considered a success and stopped

568 Interval tasks

569 2-Context task. unless otherwise specified, inspired by the timing task used by previous 570 works [14, 18] in which context cues indicated the lengths of intervals, we designed a 2-Context 571 two-interval task. In this task, the output of the RNN needs to generate either a short (3 s) or long 572 (6 s) interval in each trial. For a given training trial with length *T*, two external inputs I_{go}^1 and 573 $I_{context}^2$ were applied at stim_{onset} after a baseline with random durations between 0.2 and 0.6 s. 574 Specifically,

575
$$I_{go}(t) = \begin{cases} 1 & stim_{onset} < t \le (stim_{onset} + 0.5) \\ 0 & otherwise \end{cases}$$
for both short and long trials.

576

577 $I_{context}(t) = \begin{cases} 0.75 \text{ or } 0.25 \\ 0 \end{cases} \quad stim_{onset} < t \le T, \text{ for short or long trials repectively} \\ otherwise \end{cases}$

578

579 The output targets were defined as:

$$z(t) = \begin{cases} \frac{t - stim_{onset} - 0.5 * Int_{target}}{0.5 * Int_{target}} & (stim_{onset} + 0.5 * Int_{target}) < t \le (stim_{onset} + Int_{target}) \\ 1 & (stim_{onset} + Int_{target}) < t \le (stim_{onset} + Int_{target} + 0.2) \\ 0 & otherwise \end{cases}$$

where the target intervals (Int_{target}) were 3 and 6 seconds for the short and long trials, respectively. *2-Stimulus task.* unless otherwise specified, the 2-Stimulus task was based on a twointerval odor discrimination task [23], which required the production of the identical output patterns as the 2-Context task. However, the short and long intervals were cued by two different inputs I_{short} and I_{long} which like the I_{go} in the 2-Context task stepped up from 0 to 1 for a brief 0.5 s period.

Task performance. Response time for a given trial was defined as the time when the output
crosses a threshold of 0.6. The correct trials were defined as those in which the output crossed the

threshold within an acceptance window between stim_{onset} + 0.5 Int_{target} and stim_{onset} + Int_{target}. Task
performance was defined as the ratio of correct trials among all testing trials.

591 Unless otherwise specified, the "delay" epoch (stim_{onset} to stim_{onset} + Int_{target}) was used for
592 analysis.

593

594 Generalization to novel inputs

To test how the RNN trained on the 2-Context task would generalize to novel intervals as in Fig 2, we first trained the RNN using the normal setting for the 2-Context task, namely I_{context} of 0.75 and 0.25 for the short and long trials, respectively. Then we tested the trained RNNs by gradually varying the context level from 0.75 to 0.25 with steps of 0.05. Fifty trials of each level were obtained for analyses.

600 After training in the 2-Stimulus task generalization to novel inputs was tested by gradually 601 varying the ratio of I_{short} and I_{long} with steps of 0.1 so that the sum of both inputs was always 1.

602 *Correlation measure* To quantify changes in the temporal profile of the output units across 603 different inputs during generalization tests we first computed the correlation coefficient between 604 the mean response times (when the output crosses the threshold) and the generalization conditions 605 for both the 2-Context and 2-Stimulus tasks (the absolute values of the correlations were used due 606 to the negative correlation for the 2-Context task).

607 *Sigmoid slope measure.* To further quantify generalization to novel inputs in both tasks we608 also fitted the mean response times to the input conditions with a sigmoid function as follow:

609
$$y = b + \frac{a-b}{1+e^{g*(m-x)}}$$

610 Standard nonlinear least square methods implemented in Matlab were used to optimize the fits. 611 We then compared the slope g for both tasks. Higher g values reflect more categorical 612 generalization.

613

614 **Prototypical dynamical regimes for timing two intervals**

To illustrate the possible neural dynamical strategies used for timing two intervals scaling, absolute, and stimulus-specific, we generated three pairs of prototypical dynamics for the short (3 s) and long (6 s) intervals composed of 100 units with the time step of 0.001 s (Fig 3). In such settings, the dynamics for the short and long interval were represented as 100×3000 and 100×6000 matrices respectively, with the row being units and column being time points.

620 The dynamics for long interval were the same for all three strategies, which was described621 as:

622
$$x_i(t) = e^{\frac{-(t - \frac{l}{100} * 6)^2}{2 * 0.8^2}} \text{ for } i = 1, 2, ..., 100$$

where dynamics of all units were Gaussian functions with the same variance but different means
uniformly spanned the whole 6 s. The dynamics for the short interval were different for the three
strategies and were defined as follows:

626 *Scaling*. The dynamics for the short interval in the scaling strategy was simply a matrix of627 uniform subsampling of the time dimension of the long dynamics.

628 *Absolute.* For the absolute strategy, the dynamics of the first 50 units for the short interval629 were the same as that for the long interval.

630 *Stimulus-specific*. For stimulus-specific example, we first uniformly subsampled the time
631 dimension of the long dynamics matrix to 3 s. Then we randomized the order of the unit indices.

633 The stimulus-specific index at the population-level (SSIpop)

634	As in Fig 4, to quantify how well the short and long neural trajectories can be explained by
635	the stimulus-specific strategy at the population level, we developed a novel stimulus-specific index
636	in population-level (SSIpop), which is largely based on establishing that the trajectories are not
637	consistent with temporal scaling or absolute timing. We first obtained the mean population
638	dynamics ($\Delta t=1$ ms) for two intervals by averaging across 25 trials, which led to two matrices,
639	X _{short} (200×3000) and X _{long} (200×6000). We computed the pairwise Euclidean distance between
640	X_{short} and X_{long} , which led to the distance matrix D (3000×6000). We then obtained the index of
641	the minimum values across each row of D , which led to the minimal distance vector I_{min} (3000×1),
642	which partially captures the relationship between the population dynamics for the short and long
643	intervals.

644 Next I_{min} was contrasted with a reference matrix **R** (3000×3000) with τ indexing the 645 column:

	г 1	1	1	1				1	1	ן 1
	3	2	2	2				2	2	2
	5	4	3	3				3	3	3
	7	6	5	4				4	4	4
646						•				
		•	•	•	•	•	·	•	•	
		•	•		•		•	•		
	5998	5998	5998	5998				4499	2999	2999
	L_{6000}	6000	6000	6000				6000	6000	3000]

647 Specifically, a given column vector corresponding to τ in **R** is defined as:

648
$$[1,2,3,...,\tau, \tau + \alpha, \tau + 2\alpha, \tau + 3\alpha, ..., \tau + (3000 - \tau)\alpha]$$

649
$$\alpha = \frac{6000 - \tau}{3000 - \tau}, \qquad \tau = 1,2,3,\dots,2999$$

650 Each column vector (3000×1) in **R** represents one absolute-scaling reference profile spanning from 651 pure scaling ($\tau = 1$) to pure absolute ($\tau = 3000$), with mixed profiles in between in which absolute 652 timing transitions to scaling at τ with the scaling factor α varied to keep the length of each vector 653 the same. We then computed the Euclidian distances between I_{min} and all the column vectors of **R** 654 and extracted the vector with the minimum distance at τ_{min} , which indicates the best reference 655 vector that can be used to explain the I_{min}. Note that the construction of the R matrix accounts for 656 units that fire throughout the entire trial—thus capturing the properties of a neuron that always 657 fired at the end of the trial (e.g., a potential motor neuron). It is also possible to build R by fixing 658 the scaling factor at 2 after each point τ , in which case the last element of each column in R would 659 progressively change from 6000 to 3000. We have run analyses with this partial scaling approach 660 as well with qualitatively similar results.

661 Finally, the SSI_{pop} was defined as:

$$SSI_{pop} = 1 - c(\tau_{min})$$

where the $c(\tau_{min})$ is the correlation between the I_{min} and the reference vector at τ_{min} . For pure scaling dynamics for the two intervals as an example, I_{min} should be the main diagonal of distance matrix **D**, [1, 3, 5, 7, ..., 6000], which makes τ_{min} = 1, corresponding to the pure scaling reference vector. Consequently, the $c(\tau_{min})$ is 1 and SSI_{pop} is 0. That indicates that the pure scaling dynamics can not be explained by stimulus-specific strategy but by absolute-scaling strategies, in which the dynamics of the short and long interval relate to each other in a way of absolute or scaling or mixed of both (see below for the absolute-scaling index at the single-unit level)

670

671 Stimulus-specific index and absolute-scaling index (ASI) for single

672 units

673 We extended a previous description of an absolute vs. scaling index (ASI) for single units 674 [23], by including a novel measure of the stimulus-specific profile: the stimulus-specific index at the single-cell level (SSI_{unit} as in Fig 5A). As described previously we searched for the best 675 transformation of dynamics for the long interval (y(t)) to that for the short interval (x(t)), by 676 concatenating an absolute portion of the long response $(y^{abs}(t))$ and a temporally scaled portion of 677 the long response scaled by a factor α (y^{scale}(t')). More specifically, we searched for a breakpoint 678 τ to divide y(t) into an absolute and scaled segment, that best matches x(t), as measured by the 679 Euclidean distance (Dist(τ)). Specifically, 680

$$\alpha = (T_{long} - \tau)/(T_{short} - \tau)$$

682
$$Dist(\tau) = \sqrt{\sum_{t=0}^{\tau} (x(t) - y(t))^2 + \sum_{t=\tau}^{T_{short}} (x(t) - y(\tau + \alpha(t - \tau))^2)}$$

$$\tau_{min} = argmin_{\tau}(Dist(\tau))$$

 $684 \quad Corr(\tau_{min})$

$$685 \qquad = \frac{\sum_{t=0}^{\tau_{min}} (x(t) - \overline{x}) (y(t) - \overline{y}) + \sum_{t=\tau_{min}}^{T_{short}} (x(t) - \overline{x}) [y(\tau_{min} + \alpha(t - \tau_{min})) - \overline{y}]}{\sqrt{\sum_{t=0}^{T_{short}} (x(t) - \overline{x})^2} \sqrt{\sum_{t=0}^{\tau_{min}} (y(t) - \overline{y})^2 + \sum_{t=\tau_{min}}^{T_{short}} (y(\tau_{min} + \alpha(t - \tau_{min})) - \overline{y})^2}}$$

$$SSI_{unit} = 1 - Corr(\tau_{min})$$

687
$$W^{abs}(\tau_{min}) = 1/N_{1:\tau_{min}} \sum_{t=0}^{\tau_{min}} |[x(t) - x(0)][y(t) - y(0)]|$$

688
$$W^{scale}(\tau_{min}) = 1/N_{\tau_{min}:T_{short}} \sum_{t=\tau_{min}}^{T_{short}} |[x(t) - x(\tau_{min})][y(\tau_{min} + \alpha(t - \tau_{min})) - y(\tau_{min})]|$$

689
$$AbsR(\tau_{min}) = \frac{W^{abs}(\tau_{min})}{W^{scale}(\tau_{min}) + W^{abs}(\tau_{min})}$$

691
$$ASI = \left(\frac{\tau_{min}}{T_{short}} + AbsR(\tau_{min})\right)/2$$

692 τ spans all possible breakpoints from 0 to T_{short} (for the short interval and T_{long} for the long interval). 693 The segment before τ denotes the absolute period and the period after τ denotes the segment scaled 694 by α for the long response. τ_{min} corresponds to the breakpoint with the minimal Euclidian distance Dist(τ_{min}). Different from previous work [23], we also computed the correlation coefficient 695 between x(t) and transformed y(t), Corr (τ_{min}). Then the SSI_{unit} is defined as that 1 minus Corr(τ_{min}). 696 697 In the following steps, the absolute and scaling weights are calculated between dynamics for the 698 short interval and the time-warped dynamics for the long interval at τ_{\min} with $N_{a:b}$ being the number 699 of time points between a and b, and absolute ratio $AbsR(\tau_{min})$ was also calculated. The absolute 700 temporal factor corresponds to τ_{min} /T_{short}, and ASI was defined as the average of the absolute 701 temporal factor and the AbsR (τ_{min}).

To classify each unit as a stimulus-specific, scaling, or absolute unit we first calculated SSI_{unit} for each unit. We then classified a unit as stimulus-specific if SSI_{unit} was > 0.5; if the SSI_{unit} was \leq 0.5 then looked at its ASI and classified it as an absolute unit if ASI > 0.5, or as a scaling unit if ASI \leq 0.5.

706

707 Unit-deletion and weight deletions experiments

Based on the classification of units being stimulus-specific, scaling, or absolute, we ran deletion experiments to start to understand the causal role of each type of unit (S6 Fig). For a given unit to be deleted, we removed all the connections attached to that unit in connection matrix W^{rec} as in Eq. 1 and then ran the RNN with the rest parameters fixed. We tested various numbers of deleted units in each type. For a given condition, we randomly selected the deleted cells from the pool 10 times and repeated each deletion experiment for 20 trials for each interval. Thenperformance and error were averaged across all selections and trials.

To quantify how much each class of connection types— $E \rightarrow E$, $E \rightarrow I$, $I \rightarrow E$ and $I \rightarrow I$ connections—contributed to the recurrent dynamics and output performance, we performed synapse deletion experiments. Similar to the unit deletions, for a specific class of connections, we set all the weights of that group to be zeros while leaving the other weights unchanged. Performance and error were then computed for each condition (Fig 6).

720

721 **Pairwise angle analysis**

722 To understand the relationships between the RNNs trained on 2-Context and 2-Stimulus 723 task and the input/output subspace (S7 Fig) defined by the inputs weights and output weights, we 724 first performed principal component analysis (PCA) on the concatenated mean dynamics for the 725 short and long intervals. We then projected the original dynamics into the first three PCs. We then 726 binned the projected dynamics into segments of 250 ms. For a given segment, a vector was 727 obtained by subtracting its start point from its end point. Finally, we computed the pairwise angles 728 between all such segment vectors across time and projections of the input/output weight vectors 729 in the same PC space.

730

731 Noise perturbation experiments

As in Fig 7, to test the robustness of the outputs of the RNN trained on the 2-Context and 2-Stimulus tasks, we first trained the two tasks with noise level $\sigma = 0.45$ as in equation (1). We then tested the trained RNNs with various levels σ from 0.1 to 0.8 for 50 trials for each interval. We then compared the error between the outputs and targets for all trials and the standard deviation of the crossing times for the correct trials. Note that for all conditions tested, the incorrect trials
were less than 10% for both tasks, and there was no significant difference for that between the two
tasks.

739

740 Statistical analyses

- 741 Statistical analyses were carried out with standard functions in MATLAB (MathWorks)
- and Prism (GraphPad Software). The sample size, type of test, P values, and the F values for
- ANOVA are indicated in the figure legends. All data and error bars represent the mean and SEM
- except for the boxplot in Fig 4, where median and quartiles were presented. In all figures, the
- 745 convention is *: P < 0.05, **: P < 0.01, ***: P < 0.001, ***: P < 0.001, ****: P < 0.0001.

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959 Supporting information:

S1 Fig. Generalization difference between the 2-Context and 2-Stimulus tasks are robust across different input 960 961 parameters. (A) Training on the 2-Context task with different analog context level pairs to signal the short (blue) and 962 long (green) intervals (top), produced similarly timed short (blue) and long (green) intervals (bottom). Dashed lines 963 denote the threshold used to measure the crossing time. (B) Training on 2-Stimulus task across different levels of 964 overlap between the two input weight vectors (overlap ratio), quantified by the angle between the two weight vectors 965 (top), and the corresponding learned output traces for short (blue) and long (green) intervals (bottom). (C) The mean 966 (top) and standard deviations (bottom) of the crossing times across 10 simulations for the generalization experiments 967 corresponding to the five conditions as in (A). (D) same as (C) but for 2-Stimulus task. (E) The sigmoid fit slopes of 968 the generalization experiments in the five conditions of 2-Context task (cyan, as in A) were significantly lower than 969 that for 2-Stimulus task (orange, as in **B**): two-way ANOVA, $F_{1,90} = 123.1$, P < 0.0001 (Left). The absolute correlation 970 coefficients of the generalization experiments in the five conditions for 2-Context task (cyan, as in A) were 971 significantly higher than that of the 2-Stimulus task (orange, as in B): two-way ANOVA on the Fisher-transformed 972 data, $F_{1,90} = 374.2$, P < 0.0001 (right). (F) The mean error across all tested levels of the noise perturbation experiments 973 in the five conditions for 2-Context task (cyan, as in A) is significantly higher than that for 2-Stimulus task (orange, 974 as in **B**): two-way ANOVA, $F_{1,90} = 106.1$, P < 0.0001 (Left). Right, the same as the left but for standard deviations of 975 the crossing times: two-way ANOVA, $F_{1.90} = 625.7$, P < 0.0001. 976

977 S2 Fig. Superior generalization in the RNNs trained on the 2-Context task is maintained in the absence of a Go 978 stimulus (A) Schematic of the 2-Context task without the Go stimulus. (B) Left, the sigmoid fit slopes in the 979 generalization experiments for the 2-Context task without Go stimulus are not significantly different from the original 980 2-Context task, and still significantly lower than that for the standard 2-Stimulus task (one-way ANOVA with posthoc 981 Tukey test, $F_{2,27} = 53.4$, ns: P = 0.669, ****: P < 0.0001). Right, the absolute correlation coefficients in the 982 generalization experiments for the 2-Context task without Go stimulus not significantly different from the 2-Context 983 task but significantly higher than that for the standard 2-Stimulus task (one-way ANOVA on the Fisher-transformed 984 data with posthoc Tukey test, $F_{2,27} = 112.9$, ns: P = 0.957, ****: P < 0.0001). 985

986 S3 Fig. Generalization in the 2-Context task relies on continuous input. (A) Schematic of the standard 2-Cotnext 987 task with persistent context input. (B) Schematic of a task in which the two intervals are signaled by the same brief 988 input, but with different analog values. (C) Plots of the mean crossing time for each RNN across input conditions for 989 the persistent (top) and transient (bottom) tasks. (D) Left, mean slope of the sigmoid fits for transient input task is 990 significantly higher than that for the persistent 2-Context task (n = 10 simulations for each, two-sided t test, $t_{18} = 9.98$, 991 P < 0.0001). Right, correlation coefficient between mean crossing times and input conditions for transient 2-Context 992 is significantly lower than that for the persistent 2-Context task (n = 10 simulations for each, two-sided t test on Fisher-993 transformed values, $t_{18} = 7.52$, P < 0.0001). (E) Standard deviations of the crossing times for each RNN in the 994 persistent 2-Context (top) and transient 2-Context (bottom) tasks, as a function of input conditions. 995

996 S4 Fig. Changing the initial gain of the recurrent weight matrix to 1.5 does not alter the generalization and 997 robustness to noise differences between the 2-Context and 2-Stimulus tasks. (A) Plots of the mean crossing time 998 for each RNN across input conditions for the 2-Context (top) and 2-Stimulus (bottom) tasks. Insets, examples of the 999 sigmoid-function fits for a single RNN (black). (B) Left, mean slope of the sigmoid fits for 2-Stimulus task is 1000 significantly higher than that for the 2-Context task (n = 20 simulations for each, two-sided t test, $t_{18} = 6.91$, $P < 10^{-10}$ 1001 0.0001). Right, correlation coefficient between mean crossing times and input conditions for 2-Context task is 1002 significantly higher than that for the 2-Stimulus task (n = 20 simulations for each, two-sided t test on Fisher-1003 transformed values, $t_{18} = 16.56$, P < 0.0001). (C) Standard deviations of the crossing times for each RNN in the 2-1004 Context (top) and 2-Stimulus (bottom) tasks, as a function of input conditions. (D) Left, mean error (across 50 trials) 1005 for 2-Context task (cyan) is higher than that for 2-Stimulus task (orange) (n = 10 simulations, two-way ANOVA with 1006 mixed-effect design, $F_{1,18} = 32.48$, P < 0.0001). Right, mean standard deviation of the time of threshold-crossing 1007 across all correct trials for 2-Context task (cyan) is higher than that for 2-Stimulus task (orange) ($F_{1,18}$ = 128.50, P <1008 0.0001). Data are presented as mean \pm SEM.

1010 S5 Fig. Full initial connectivity of the weight matrix does not alter the generalization and robustness to noise 1011 differences between the 2-Context and 2-Stimulus tasks. (A) Plots of the mean crossing time for each RNN across input conditions for the 2-Context (top) and 2-Stimulus (bottom) tasks. Insets, examples of the sigmoid-function fits 1012 1013 for a single RNN (black). (B) Left, mean slope of the sigmoid fits for 2-Stimulus task is significantly higher than that 1014 for the 2-Context task (n = 20 simulations for each, two-sided t test, $t_{18} = 4.35$, P = 0.00039). Right, correlation 1015 coefficient between mean crossing times and input conditions for 2-Context task is significantly higher than that for 1016 the 2-Stimulus task (n = 20 simulations for each, two-sided t test on Fisher-transformed values, $t_{18} = 6.48$, P < 0.0001). 1017 (C) Standard deviations of the crossing times for each RNN in the 2-Context (top) and 2-Stimulus (bottom) tasks, as 1018 a function of input conditions. (D) Left, mean error (across 50 trials) for 2-Context task (cyan) is higher than that for 1019 2-Stimulus task (orange) (n = 10 simulations, two-way ANOVA with mixed-effect design, $F_{1,18} = 5.78$, P = 0.027). 1020 Right, mean standard deviation of the time of threshold-crossing across all correct trials for 2-Context task (cyan) is 1021 higher than that for 2-Stimulus task (orange) ($F_{L18} = 86.03$, P < 0.0001). Data are presented as mean \pm SEM. 1022

1023 S6 Fig. Differential functional effects of deleting specific classes of units. (A) Schematic of the deletion 1024 experiments. To delete a given unit denoted by the red arrow (bottom), all in and out weights of the recurrent weight 1025 matrix of that units were set to zero. (B) Performance of RNNs trained on the 2-Context task after progressively 1026 deleting units from specific temporal classes: stimulus-specific, scaling, and absolute temporal classes for both 1027 excitatory (left) and inhibitory (right) units. For each data point, units were randomly selected 10 times, and 10 test 1028 trials were obtained. A three-way ANOVA revealed highly significant effects of main temporal-class ($F_{2,619}$ = 31, P < 1029 10⁻¹²) and Ex-Inh ($F_{2,619}$ = 390, $P < 10^{-66}$) factors. Additionally, there was a highly significant interaction between 1030 temporal-class and Ex-Inh class ($F_{2,619} = 27$, $P < 10^{-10}$) and multi-comparison analyses showed that performance for 1031 inhibitory scaling cells was significantly lower than all other 5 deletion manipulations (P < 0.0001 for all comparisons). 1032 (C) Similar to (B) but for error. As in (B), there were highly significant main effects ($F_{2,619} = 34$, $P < 10^{-14}$, and $F_{2,619}$ 1033 = 118, $P < 10^{-24}$, for temporal-class and Ex-Inh, respectively), as well as a significant interaction between temporal-1034 class and Ex-Inh ($F_{2,619}$ = 46, $P < 10^{-18}$). And again the inhibitory scaling cells increased the error more than all other 1035 deletion manipulations (P < 0.0001 for all comparisons). (D-E) There were no main effects of temporal-class or Ex-1036 Inh that were consistently significant for both the performance and error measure. The interaction between temporal-1037 class and Ex-Inh was either trending ($F_{2,619} = 2.5$, P = 0.08) or mildly significant ($F_{2,619} = 3.6$, P = 0.027) for the 1038 performance and error analyses, respectively. Data are presented as performance mean ± SEM across 20 RNNs. Notice 1039 that the performance of stimulus-specific units in (D) and (E) (magenta) are very similar to, and mostly obscured by 1040 the absolute traces (red). (F) Mean output traces across 20 simulations when deleting 6 excitatory (left) and inhibitory 1041 (right) units of the three types: stimulus-specific, scaling, and absolute for 2-Context task. (G) Same as F but for 2-1042 Stimulus task. 1043

1044 S7 Fig. Differential subspace dynamics for RNNs trained on 2-Context and 2-Stimulus tasks. (A) For the 2-1045 Context task, recurrent unit dynamics for the short (blue) and long (green) intervals were projected into the first three 1046 PC spaces. Asterisks denote the onset of inputs (t=0), arrows denote the corresponding weights vectors (Input_{Go}, black; 1047 Input_{Context}, cyan; and Output, red) projected onto the same PC space. Color dots denote the 250 ms intervals along 1048 each trajectory. Inset, schematic of angles between segments of the approximate RNN trajectory (orange) and the 1049 three weight vectors. These vectors were used to compute the pairwise angles to the Input_{Go}, Input_{Context} and Output 1050 vectors. (B) Similar to (A) but for 2-Stimulus task, but here the two input vectors represented the Input_{short} (blue) and 1051 Input_{Long} (green) weight vectors. (C) Same number of PCs explained more variance for 2-Context task than that for 2-1052 Stimulus task (Two-way ANOVA, $F_{(1, 38)} = 255.6$ and P < 0.0001). (D) Average pairwise angles between segments of 1053 short (top)/long (bottom) dynamics and inputs/output vectors as in (A) for 2-Context task (20 simulations, data 1054 presents as Mean \pm SEM). Shaded area donted the duration of the transient Input_{Go} (E) Same as in (D) but for 2-Stimulus task. The shaded area denotes the duration of the transient Input_{Short} and Input_{Long}. 1055

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1058 S8 Fig. PCA plots of the recurrent dynamics for generalization to novel intervals (A) Recurrent dynamics corresponding to different context levels (denoted by the color) as in Fig. 2 were projected into the first three PCs in 20 RNNS trained on 2-Context task. The arrows denoted the directions of Input_{Go} (black), Input_{Context} (cyan), and Output (red) weights projected into the same PC space. (B) similar as in (A) but for the 2-Stimulus task.
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1074 Data and codes availability

- 1075 All data are available in the main text or supplementary materials. Codes used for the simulations
- 1076 in this paper are available at (https://github.com/ShanglinZhou/RNN_2Intervals).

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