Optimal information loading into working memory in prefrontal cortex explains dynamic coding

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Abstract

Working memory involves the short-term maintenance of information and is critical in many tasks. The neural circuit dynamics underlying working memory remain poorly understood, with different aspects of prefrontal cortical (PFC) responses explained by different putative mechanisms. By mathematical analysis, numerical simulations, and using recordings from monkey PFC, we investigate a critical but hitherto ignored aspect of working memory dynamics: information loading. We find that, contrary to common assumptions, optimal loading of information into working memory involves inputs that are largely orthogonal, rather than similar, to the persistent activities observed during memory maintenance, naturally leading to the widely observed phenomenon of dynamic coding in PFC. Using a novel, theoretically principled metric, we show that PFC exhibits the hallmarks of optimal information loading. We also find that optimal loading emerges as a general dynamical strategy in task-optimized recurrent neural networks. Our theory unifies previous, seemingly conflicting theories of memory maintenance based on attractor or purely sequential dynamics, and reveals a normative principle underlying dynamic coding.

Working memory requires the ability to temporarily 14 hold information in mind, and it is essential to perform-15 ing cognitively demanding tasks^{1,2}. A widely observed 16 neural correlate of the maintenance of information in working memory is selective persistent activity. For 18 example, in the paradigmatic memory-guided saccade task³⁻¹³, subjects must maintain the location of one 20 out of several cues during a delay period after which they must respond with a saccade to the correct lo-22 cation (Fig. 1a). Cells in the lateral prefrontal cortex (IPFC) show elevated levels of activity that persist during the delay period and that is selective to the location 25 of the now-absent cue^{3-5,9}. However, neurons typically only reach a steady, persistent level of activity late in the delay period of a trial^{6,8,10,11,14-20}. In contrast, during the cue and early delay period, neurons 29 in IPFC often exhibit strong transient dynamics dur-30 ing a variety of working memory tasks^{3,8,10,11,14-24}. It 31 remains unknown what mechanism may underlie this 32 combination of persistent and dynamically changing 33 neural activities. 34

Recent population-level analyses using the technique of 'cross-temporal decoding' place particularly strin-36 gent constraints on any candidate neural mechanism 37 of working memory maintenance. Cross-temporal de-38 coding measures how well information about the cue location can be decoded from neural responses when 40 a decoder is trained and tested on any pair of time 41 points during a trial^{8,10,11,14,15,25} (Fig. 1b). These analyses reveal a consistent but somewhat puzzling set 44 of results. First, when decoder training and testing times are identical, decodability is high (Fig. 1b, black

⁴⁶ along the diagonal), confirming that information about 47 cue location is indeed present in the population at all times^{8,10,11,14,15,25}. Decodability is also high when both training and testing occurs during the late delay 49 period^{8,10,11,14,15,25}, suggesting that even if there are changes in neural responses during this period, the coding of cue location remains stable^{6,8,10} (Fig. 1b, black inside cyan square). However, decoding perfor-54 mance remains low when a decoder is trained during st the cue or early delay period and tested during the ⁵⁶ late delay period, and vice-versa^{8,10,11,14,15,25} (Fig. 1b, ⁵⁷ gray inside pink rectangles). This demonstrates that ⁵⁸ the neural code for cue location undergoes a substantial change between these these two periods-60 a phenomenon that has been called 'dynamic cod-61 ing'^{8,10,14–16,25}.

62 Classically, the neural mechanism of working memory ⁶³ maintenance is thought to rely on attractor network dynamics^{5-7,12,27-31}. In such networks, the stimulus cue acts as a transient external input, driving the network activity (Fig. 1c; left, pale purple line and arrow) into a suitable state (Fig. 1c; left, pale purple circle) from which its intrinsic dynamics (Fig. 1c; left, dark purple 68 ⁶⁹ line), in the absence of the cue, are 'attracted' into a 70 distinct cue-specific steady state maintained by recur-⁷¹ rent interaction between neurons (Fig. 1c; left, black 72 cross). Thus, these models naturally account for selective persistent activity (Fig. 1c, center). However, because the external input drives network activity to a state that already has large overlap with (i.e. it is sim- $_{76}$ ilar to) the desired attractor state 5-7,29,31-35 (Fig. 1c; ⁷⁷ left, gray arrow), the ensuing dynamics then perform



Fig. 1 | Neural network dynamics of working memory maintenance. a, Illustration of the memory-guided saccade task. Time line of task events in a trial (bottom), with the corresponding displays (top). Top: black circle and squares show fixation ring, and the arrangement of visually cued saccade target locations, respectively (not to scale), red dots and line illustrate gaze positions during fixations and saccade, respectively. Bottom: yellow ticks show timing of stimulus cue onset and offset, yellow bar shows interval within which the go cue can occur. b, Schematic pattern of cross-temporal decoding when applied to neural recordings from the IPFC during working memory tasks^{8,10,14–16,25}. Gray scale map shows accuracy of decoding cue identity (one out of 6) when the decoder is trained on neural activities recorded at a particular time in the trial (y-axis) and tested at another time (x-axis). Yellow lines indicate cue onset and offset times. Note poor generalization between time points inside the pink rectangle (i.e. dynamic coding), but good generalization between time points inside the blue rectangle (i.e. stable coding). The gray tick on the color bar indicates chance-level decoding. c, Schematic of neural network dynamics in an attractor network performing the task shown in a ^{5,6} (see also Extended Data Fig. 1a-c). Left: trajectory in neural state space in a single cue condition during the cue period (pale purple line, ending in pale purple circle) and delay period (dark purple line). Purple arrow heads indicate direction of travel along the trajectory, black cross shows attractor state, gray arrow shows overlap between cue input and late delay activity. Center: time course of relative (i.e. mean-centered) firing rates of one neuron (dim 1 from left panel) for two cue conditions (purple vs. blue, see also inset). Yellow lines indicate cue onset and offset times. Right: cross-temporal decoding of neural activity produced by the network across all 6 cue conditions, shown as in **b**. **d**. Same as **c**, but for an effective feedforward network that generates seguential activities^{21,26} (see also Extended Data Fig. 1d). e, Same as c, but for a network optimized to perform the task shown in a (see also Extended Data Fig. 1e).

'pattern completion'³⁵, whereby this overlap is only 78 slightly improved until it becomes perfect and the de-70 sired attractor is reached (Fig. 1c; left). As a result, 80 neurons show limited transient activity during the delay 81 period (Fig. 1c, center), and cross-temporal decoding 82 reveals stable coding throughout the whole trial, lack-83 ing the characteristic dynamic coding seen in experi-84 mental data (compare Fig. 1b to c, right;^{8,10,14–16,25}). 85 This combination of results emerge across several 86 variants of attractor networks, whether they express 87 a continuum of persistent activity patterns ('ring' or 88 'bump' attractor networks, Extended Data Fig. 1a) or 89 a finite number of discrete patterns (Extended Data 90 Fig. 1b). Critically, even when external inputs in attrac-91 tor, or closely related 'integrator', models were chosen 92 such that neural activity showed longer transient dy-93 namics^{6,32} (Extended Data Fig. 1c, center), inputs still 94 ⁹⁵ relied on a large overlap with the desired attractor (Ex-

tended Data Fig. 1c, left). Therefore, these networks
 maintained a stable code over time without dynamic
 coding⁶ (Extended Data Fig. 1c, right).

⁹⁹ To capture transient dynamics more naturally, a very different class of models have been developed based on mechanisms that generate neural activity sequences. These models typically rely either on effectively feedforward network connectivity^{21,26} or chaotic network dynamics^{24,36–38}. The dynamics of such models rapidly transition between orthogonal subspaces over time (Fig. 1d, left), thus cross-temporal decoding is high only between neighbouring time-points (Fig. 1d, right, black along diagonal). Although such models are ideally suited to capturing transient neural responses (Fig. 1d, center) and poor cross-temporal decoding between cue/early delay and late delay periods (Fig. 1d, right, gray inside pink rectangle), they fail to exhibit persistent activities (Fig. 1d, center) and sta182

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ble coding during the late delay period (Fig. 1d, right, gray inside blue square, except for diagonal). Therefore, previous work leaves open two interrelated key 116 questions: how can a neural circuit exhibit strong transient dynamics before its activity ultimately settles in 118 an orthogonal persistent state, and why would it use such a counterintuitive dynamical regime? 120

In order to study the dynamical principles underlying the combination of persistent and dynamic neu-122 ral activities during working memory, we built on recent advances in using task-optimized neural net-124 works to study network mechanisms^{13,17,20,24,36,39-41}. Thus, instead of starting from strong prior assumptions about either attractor or sequential dynamics underlying working memory, we trained networks for the 128 task of working memory maintenance, and analysed their dynamical behaviour. We found that the be-130 haviour of such task-optimized networks unifies attractor and sequential activity models, showing both persistent activities and orthogonal transient dynamics. giving rise to dynamic coding (Fig. 1e).

To understand the mechanism and functional significance of dynamic coding, we focused on a hitherto 136 ignored aspect of the operation of attractor networks: optimal information loading. In particular, we show 138 that inputs that most efficiently drive the network activity into a desired attractor state tend to be orthogo-140 nal to the attractor state itself which results in an initial 141 period of strong transient dynamics. Thus, transient 142 dynamics and dynamic coding are fundamental and functionally useful features of attractor networks. In 144 order to gain an analytical understanding of this phe-145 nomenon, we developed a mathematical theory for the 146 efficiency of information loading in attractor networks 147 by analysing a simplified class of neural network mod-148 els with linear dynamics. Crucially, our theory also 149 suggested a specific neural data analysis approach for 150 assessing whether a network uses optimal information loading. Using this theoretically-principled approach, we demonstrate key signatures of optimal information loading in neural recordings from IPFC. Finally, we as-154 sess how different cost functions affect the dynamics of task-optimized networks. We show that dynamic 156 coding always emerges after training in a wide variety 157 of models including linear integrators, as well as non-158 linear discrete and bump attractor models, unless the 159 cost function explicitly requires stable coding. Our re-160 sults offer a novel, normative perspective on a core 161 component of the operation of attractor networksinformation loading—which has so far received little 163 attention, and challenge long-held assumptions about 164 pattern completion-like mechanisms in neural circuits. 165 166

Results 167

Pattern completion and optimal information load-168 ing in attractor networks

Traditional approaches to studying attractor networks used models in which the connectivity between

172 neurons was constrained to be effectively symmetric^{5,7,31,33–35,42–46}, making the analysis of their dynamics mathematically more convenient^{34,35,44,47,48}. Thus, 174 we first replicated results with such symmetric networks that were optimized to perform the working memory task shown in Fig. 1a. (While here we show results with attractor networks generated by a particu-178 lar optimization procedure, we show below that these results do not depend on the details of this proce-180 dure, only on the presence of suitable attractors in 181 the state space of the resulting networks.) For simplicity, we only modelled the intrinsic dynamics of the network during the delay period and the effect of the cue was captured by cue-specific initial neural activi-185 ties (i.e. neural activities at the beginning of the delay period^{35,42,43}; Fig. 2b). To study optimal information loading, we optimized these initial activities in order to maximize the performance of the network, as determined by how well the cue could be decoded from neural activities at the end of the delay period (Methods 1.3.1). In other words, we asked where (in neural state space) the dynamics of the network need to start ¹⁹⁴ from so as to consequently generate a robustly identi-¹⁹⁵ fiable, cue-specific pattern of persistent activity.

We found that optimal initial activities gave rise to classical pattern completion dynamics in symmetric networks. First, initial activities were noisy versions of (and in fact highly similar to) the desired persistent patterns (Fig. 2b inset, and Fig. 2c). Second, the ensu-²⁰¹ ing dynamics were driven directly into the correspond-²⁰² ing steady state resulting in only small and gradual changes in activities over the delay period (Fig. 2b). Further analysis of these dynamics showed that the optimal initial activities aligned well with directions in neural state space that best distinguished between the desired persistent activities (Fig. 2d, 'persistent PC1' component of pale arrows and circles; Extended Data Fig. 2b), with only a comparably small component in orthogonal directions specific to these initial activities (Fig. 2d, 'initial PC1 (orthogonalized)') which subse-²¹² quently changed little over time (Fig. 2d, dark trajectories). As a result, cross-temporal decoding performance was high for all pairs of times (Fig. 2e), andas a special case-a decoder based on templates of neural activity during the late delay period (i.e. dur-216 ²¹⁷ ing the steady state of the network), generalized well to all times and was able to decode the cue identity from neural activities with high accuracy throughout the delay period (Fig. 2f, black line). We found that the similarity between initial and persistent activities was critical for these networks. When constrained to use initial activities that were orthogonal in neural state space to persistent activities (i.e. lying in the 'persis-224 ²²⁵ tent nullspace'), these networks performed substantially more poorly at all times (Fig. 2f, red line) and activity often did not settle into the correct attractor state (Extended Data Fig. 2d). In contrast, explicitly enforcing these networks to use initial activities that were similar to persistent activities (i.e. lying in the 'persistent subspace') did not compromise their performance (Fig. 2f, green line; Extended Data Fig. 2c).

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Fig. 2 | Pattern completion and optimal information loading in attractor networks. a, A network with symmetric connections. Left: network schematic. Right: the recurrent weight matrix for 10 of the 50 neurons. b-f, Analysis of neural responses in symmetric attractor networks (such as shown in a) with optimized initial conditions. b, Firing rate activity in a representative trial. Inset shows initial vs. final mean-centered firing rates across neurons (gray dots) in this trial and the Pearson correlation (r; p < 0.001) between initial and final firing rates. Gray line is the identity line. c, Distribution of Pearson correlations between initial and final mean-centered neural firing rates across all 6 cue conditions and 10 networks. d, Sub-threshold activity for 2 cue conditions in an example network (color trajectories). Open circles (with arrows pointing to them from the origin) show the optimized initial conditions, black crosses show stable fixed points, dashed gray line is the identity line. Horizontal axis (persistent PC1) shows network activity projected on to the 1st principal component (PC1) of network activities at the end of the delay period (across the 2 conditions shown), vertical axis (initial PC1 (orthogonalized)) shows projection to PC1 of initial network activities orthogonalized to persistent PC1. e, Cross-temporal decoding of neural firing rate activity (cf. Fig. 1b). f, Performance of a delay-trained decoder (black bar indicates decoding training time period) on neural firing rate activity over time starting from optimized initial conditions with full optimization (black), or restricted to the 5-dimensional subspace spanning the 6 cue-specific attractors (persistent subspace, green), or the subspace orthogonal to that (persistent nullspace, red). Solid lines and shading indicate mean±1 s.d. across all 6 cue conditions and 10 networks. Gray dotted line shows chance level decoding. Green and black lines are slightly offset vertically to aid visualization. g, Same as b but for an attractor network with unconstrained connections. h-I, Same as c-f, for attractor networks with unconstrained connections. The Pearson correlation in **h** (inset) is not significant (p > 0.4).

straint exhibited dynamics distinctly unlike simple pattern completion (Fig. 2g-I). First, initial activities re-236 even be negative (Fig. 2h inset). Second, neural activ-238 ities often underwent substantial and non-monotonic changes before ultimately settling into an attractor 240 state (Fig. 2h). This was also reflected in optimal initial

Attractor networks optimized without a symmetry con-244 tended Data Fig. 2f), with this orthogonality decaying over the delay period (Fig. 2j, dark trajectories). Third, 245 a decoder trained on neural activity from the late desembled persistent activity much less than in symmet- 247 lay period generalized poorly to early times (Fig. 2I, ric networks (Fig. 2i), such that their correlation could 248 black line) and vice versa (Fig. 2k), thus exhibiting a ²⁴⁹ fundamental signature of 'dynamic coding'^{10,14–16} (cf. ²⁵⁰ Fig. 1b). We found that the orthogonality of initial con-²⁵¹ ditions in these networks was instrumental for high ²⁵² performance: in a double dissociation from symmetactivities (Fig. 2j, pale arrows) being strongly orthogo- 253 rically constrained networks, restricting initial condinal to persistent activities (Fig. 2j, black crosses; Ex- 254 tions to be in the persistent subspace (Fig. 2l, green

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line; Extended Data Fig. 2g), but not in the persistent 314 between two stimuli depending on which side of the nullspace (Fig. 2l, red line; Extended Data Fig. 2h), di-256 minished decodability at the end of the delay period 257 (cf. Fig. 2f). 258

The above results were obtained with networks storing a small number of discrete attractors, corresponding to 260 the six cue conditions. Previous work found that sev-261 eral aspects of working memory dynamics in IPFC are better captured by networks in which instead a large 263 number (or even a continuum) of attractor states lie on a ring in state space^{5,7,45,46}. Thus, we repeated 265 ur analyses on optimized networks while explicitly 0 encouraging such a ring attractor to form during op-267 timization (Methods 1.3.4). We found a highly similar 268 pattern of results in ring attractor networks as compared with discrete attractor networks (Extended Data Fig. 3).

Dynamical analysis of optimal information loading

understand why optimal information loading in Тο symmetric versus unconstrained attractor networks is so different, and in particular why inputs orthogonal to attractor states are optimal for unconstrained 276 networks, we reduced these networks to a canonical minimal model class consisting of only two neu-278 ons^{43,49,50}. While attractor network dynamics in general rely on the activation functions (f-I curves) of neu-280 rons being nonlinear^{27,33,35,44}, for analytical tractability, 281 we considered networks with linear dynamics (i.e. in which neurons had linear activation functions). Crit-283 ically, with the appropriate set of synaptic connec-28/ tions, even linear networks can exhibit persistent ac-285 tivity^{6,32,34,43,48,51}—the key feature of working memory 286

maintenance in attractor networks. 287

For our analyses, we again distinguished between 288 models with symmetric connectivity between neurons 289 (Fig. 3a; top)^{34,43,50}, and models without this constraint 290 (Fig. 3a; bottom)^{6,32}. In either case, the specific con-291 nection strengths were chosen to create illustrative examples providing intuitions that-as we show below-293 also generalise to large networks with randomly sam-294 pled connection strengths (Fig. 3d-e, Fig. 4). The dy-295 namics of these networks are fully described in a two-296 dimensional state space spanned by the activities of 297 the two neurons (Fig. 3b) and define a flow-field in this 298 space determining how changes in neural activities 299 depend on the network state (Fig. 3b; blue arrows). 300 While the persistent subspace of nonlinear networks 301 an consist of a number of discrete attractor states 302 corresponding to distinct patterns of persistent activ-303 ity (Fig. 2 and Extended Data Fig. 2, Methods 1.3), 304 linear attractor networks (or 'integrators'⁴³) express a 305 continuum of persistent activity patterns^{6,32,34,43}. In ei-306 ther case, attractor networks encode stimulus informa-307 tion in the location of the state of the network within the persistent subspace. In our two-neuron linear network, the persistent subspace simply corresponds to a 310 line onto which the flow field converges (Fig. 3b; green lines). Therefore, the persistent mode of our network is its 'coding direction'³³, which allows it to distinguish

origin the state of the network is. The larger the magnitude of its activity along this mode at the end of the 316 delay period, the more robustly the identity of the stimulus can be decoded (e.g. in the presence of noise).

To understand the mechanisms of information loading, we considered three distinct stimulus directions 321 in which inputs can offset the state of the network ³²² from the origin (i.e. the background state of the net-323 work before stimulus onset). We then analysed the ³²⁴ time course of the network's activity along the persistent mode^{6,32,33} after being initialised in each of these directions. First, we considered inputs aligned with the persistent mode, the input direction studied in classical attractor networks^{6,32,34,43,50} (Fig. 3b; pale green 328 arrows and open circles). Second, we considered the 'most amplifying mode', which is defined as the stim-330 ulus direction that generates the most divergent and thus best discriminable activity over time⁵²⁻⁵⁶ (Methods 1.7.1; Fig. 3b, red lines, and pale red arrows and ³³⁴ open circles). Third, we considered a random input direction (Fig. 3b; gray lines/circles).

In symmetric networks, the most amplifying mode ³³⁷ is aligned with the most persistent mode (Fig. 3b; 338 top)^{57,58}, and thus does not generate activity transients (Fig. 3c; top)-accounting for the simple pattern completion dynamics seen in classical attractor networks with symmetric connectivity 5,7,31,33-35,42-44 (Fig. 2a-f). However, in unconstrained networks, the 342 343 most amplifying mode is typically different from the most persistent mode (Fig. 3b; bottom). Intuitively, this is because effective feedforward connections exist in unconstrained networks^{21,26,49,55,59} (Fig. 3a, bottom; connection from neuron 2 to neuron 1). Feeding neuron 1 (the persistent mode) indirectly through this feedforward connection from neuron 2 can increase its activity more than just feeding it directly^{21,26} (Fig. 3a,b; bottom). This means that activity evolving from the 351 most amplifying mode exhibits a distinct transient behaviour: its overlap with the most persistent mode 353 is initially low and then increases over time (Fig. 3c; bottom, red line), accounting for the richer transients seen in unconstrained attractor networks (Fig. 2g–I). Thus, there is a form of 'speed-accuracy' trade-off be-³⁵⁸ tween whether inputs should use the most amplifying ³⁵⁹ or persistent mode: if information is required immediately following stimulus offset, such as in a percep-1361 tual decision-making task^{13,40,58}, inputs need to use the persistent mode. However, if there is a time delay until the information is needed, as is the case in all working memory tasks^{2,60}, then the most amplifying 365 mode becomes the optimal input direction. Indeed, an analogous trade-off was already apparent between the 366 persistent sub- vs. nullspace inputs in the non-linear attractor networks we analysed earlier (Fig. 2l, red vs. 368 green).

The insights obtained in the simple two-neuron net-370 ³⁷¹ work also generalised to large randomly connected 372 linear integrator networks, with more than two neu-³⁷³ rons (Fig. 3d,e; see Methods 1.4.1). We were able to



Fig. 3 | Dynamical analysis of optimal information loading. a, Architecture of a symmetric (top) and an unconstrained network (bottom; Methods 1.4.1). b, Neural state space of the symmetric (top) and unconstrained network (bottom). Pale blue arrows show flow field dynamics (direction and magnitude of movement in the state space as a function of the momentary state). Thin green and red lines indicate the persistent and most amplifying modes (Methods 1.7.1), respectively (lines are offset slightly in the top panel to aid visualisation). Pale green, red, and gray arrows with open circles at the end indicate persistent, most amplifying, and random initial conditions, respectively. Dark green, red, and black arrows show neural dynamics starting from the corresponding initial condition. (Green arrows, and the red arrow in the top panel cannot be seen, as no movement in state space happens from those initial conditions.) Filled colored circles indicate final (persistent) neural activity. c, Time course of network activity along the persistent mode (i.e. projection onto the green line in b) when started from the persistent (green), most amplifying (red), or random initial conditions (black) for the symmetric (top) and the unconstrained model (bottom). d, Distributions of absolute overlap with the persistent mode for persistent (pale green), most amplifying (pale red), or random initial conditions (gray) across 100 randomly connected 1000-neuron symmetric (top) or unconstrained networks (bottom; Methods 1.4.1). For the symmetric models, the persistent and most amplifying initial conditions produce delta functions at 1 (arrows). Insets show illustration of large networks of neurons with either symmetric (top) or unconstrained (bottom) connections. e, Time course of absolute overlap with the persistent mode when starting network dynamics from persistent (green), most amplifying (red), or random initial conditions (black) for the symmetric (top) and the unconstrained network (bottom). Lines and shaded areas show mean±1 s.d. over the 100 randomly sampled 1000-neuron networks from d.

⁷⁴ show mathematically that optimal information loading,

in the sense of maximizing overlap with the persistent mode at long delays, is always achieved with inputs aligned with the most amplifying mode (Supplementary Math Note S1). Equivalently, the most amplifying 378 mode is the input direction that requires the smallest magnitude initial condition to achieve a desired level of 380 persistent activity (i.e. a desired level of performance). 381 More generally, we could also show both mathemat-382 ically and in simulations (Extended Data Fig. 4) that 383 the most amplifying mode is near-optimal in achiev-384 ing a desired level of performance while minimizing 385 total network activity over time (i.e. the total energy 386 used by the network) for sufficiently long delay lengths. 387 Moreover, as network size grows, in unconstrained 388 (but not in symmetric) networks, the most amplifying 389 direction becomes increasingly orthogonal to the most 390 persistent mode⁶¹, further accentuating the advantage 391 of amplifying over persistent mode inputs⁶¹ (Fig. 3d-392 e, Extended Data Fig. 5a-b; red vs. green). This is 393 because in large unconstrained networks, there are many effectively feed-forward motifs embedded in the 395 full recurrent connectivity of the circuit, which can all contribute to transient amplification²¹. Random ini-397 tial conditions become fully orthogonal in both net-398 works and result in poor overlap with the persistent mode (Fig. 3d-e, Extended Data Fig. 5a-b; black). 400 Numerical simulations confirmed that these results 401 also generalised to networks with noisy dynamics (Extended Data Fig. 5c). Moreover, explicitly optimizing a network's initial condition in such networks so as to generate persistent activity also made it over-

⁴⁰⁶ lap strongly with the most amplifying mode (Extended ⁴⁰⁷ Data Fig. 5d).

As our mathematical analyses only applied to linear dynamics, we used numerical simulations to study 409 how they generalised to nonlinear dynamics. We 410 found that the same principles applied to the dynamics of a canonical 2-dimensional nonlinear attractor system (analogous to the networks in Fig. 3a-c), when the persistent and most amplifying directions were defined locally around its ground state (Methods 1.6; Extended Data Fig. 6, see also Supplementary Math Note S2). Importantly, we also found that large optimized nonlinear neural networks (with discrete or ring attractors) also showed a similar pattern of results (Extended Data Fig. 3e, and Extended Data Fig. 7a-c, see also Supplementary Math Note S3).

Neural signatures of optimal information loading

423 Our dynamical analysis suggested that there should 424 be clearly identifiable neural signatures of a network performing optimal information loading. To demon-425 426 strate this, and to allow a more direct comparison 427 with data, we used the same large, randomly con-⁴²⁸ nected, noisy linear networks that we analysed ear-129 lier (Extended Data Fig. 5c-d) which could be either symmetric or unconstrained, with the cue period (Fig. 4, yellow ticks and lines) modelled using tempo-431 rally extended constant inputs, mimicking typical experiments^{3–5,10}. We studied the three different infor-433 mation loading strategies that we identified earlier: in-434 ⁴³⁵ puts aligned with either the persistent mode, the most

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Fig. 4 | Neural signatures of optimal information loading. a, Performance of a delay-trained decoder (black bar indicates decoder training time period; Methods 1.7.4) on neural activity over time. Two cue conditions were used with inputs that were identical but had opposite signs. Lines show mean across 10 randomly connected 100-neuron linear symmetric networks (top) and unconstrained networks (bottom) (Methods 1.4.1). Yellow ticks on horizontal axis indicate cue onset and offset times and the vertical gray bar indicates the cue epoch. We show results for inputs aligned with the persistent mode (dark and pale green), the most amplifying mode (red), or a random direction (black and gray). Light colors (pale green and gray, 'noise-matched') correspond to networks with the same level of noise as in the reference network (red), while dark colors (dark green and black, 'performance-matched') correspond to networks with the same level of asymptotic decoding performance as that in the reference network. Gray dotted line shows chance level decoding. Green, red, and black lines are slightly offset vertically in the top panel to aid visualization. b, Cross-temporal decoding of neural activity for the 3 different information loading strategies (persistent, most amplifying, and random respectively in left, center, and right panels) for a representative symmetric network (top) and unconstrained network (bottom) for the performance-matched condition from a. Yellow lines indicate cue onset and offset times. Pink rectangles indicate poor generalization between time points (i.e. dynamic coding) and cyan rectangles indicate examples of good generalization between time points (i.e. stable coding). c, Percent variance explained by the subspace spanned by either the 25% most persistent (green) or 25% most amplifying (red) modes as a function of time in the same symmetric (top) and unconstrained networks (bottom) analyzed in a. Lines and error bars show mean±1 s.d. across networks. We show results for inputs aligned with the persistent mode (left), most amplifying mode (center), or a random direction (right). Gray dotted line shows chance level overlap with a randomly chosen subspace occupying 25% of the full space.

⁴³⁶ amplifying mode, or a cue-specific random direction ⁴⁶⁵ ⁴³⁷ (Fig. 4).

We began by conducting a decoding analysis using templates of late delay activity, as is often done for prefrontal cortical recordings^{6,8,10,14,15,25} (and also in 440 Fig. 2f,I). We first verified that for a fixed level of neu-441 ronal noise, the most amplifying inputs were indeed optimal for achieving high decodability at the end of 443 the delay period (Fig. 4a, red lines). More gener-444 ally, we were also able to show mathematically that 445 the most amplifying inputs in noisy linear networks 446 are optimal for maximizing average decodability dur-447 ing the delay period (Supplementary Math Note S1.7). 448 In contrast, random inputs in both symmetric and unconstrained networks performed considerably more 450 poorly (Fig. 4a, gray lines). Remarkably, persistent 451 mode inputs achieved a similarly low level of decod-452 ability at late delay times in unconstrained networks 453 (Fig. 4a, bottom; compare pale green and gray lines)— 454 but not in symmetric networks in which they were iden-455 tical to the most amplifying input (Fig. 4a, top; overlapping green and red lines). 457

⁴⁵⁸ The level of noise in the networks we have studied so ⁴⁵⁹ far was not constrained by data, which typically shows ⁴⁶⁰ high decodability ^{6,8,10,14,15,25}. This is important be-⁴⁶¹ cause the sub-optimal input conditions could achieve ⁴⁶² high decoding performance by appropriately reducing ⁴⁶³ the noise level in our simulations (Fig. 4a, asymptotic ⁴⁶⁴ values of dark green and black lines). Thus, asymp-

totic decoding performance alone cannot be used to identify the information loading strategy employed by a network. To address this, in subsequent analyses, we used networks in which the level of late-delay performance was matched between the three information loading strategies by appropriately changing the level of noise. Nevertheless, a critical difference emerged 471 between the different information loading strategies even in these 'performance-matched' networks-at 474 least in those with realistic, unconstrained connectivity. For both random and most amplifying input directions, the delay-trained decoder only performed well when tested late in the delay period (Fig. 4a, bottom; black and red lines), whereas for inputs aligned with the persistent direction this decoder performed near ceiling at all times after cue onset (Fig. 4a, bottom; dark green 481 line).

⁴³² Next, in order to more fully characterise the differ-⁴³³ ences between persistent versus random or most am-⁴³⁴ plifying inputs, and for a comprehensive comparison ⁴³⁵ with experimental data^{8,10,14,15,25}, we also employed ⁴³⁶ full cross-temporal decoding (Fig. 4b). This analysis ⁴³⁷ showed that all information loading strategies led to ⁴³⁸ dynamics in which stimulus information was present ⁴³⁹ at all times after cue onset (Fig. 4b, diagonals are ⁴³⁰ all black). Moreover, in symmetric networks, and for ⁴³¹ the persistent mode inputs in the unconstrained net-⁴³² work, stimulus information was maintained using a ⁴³³ 'stable code' ^{10,11,14,16} (Fig. 4b, top, and bottom left, all 567

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off-diagonals are black)—similar to previous integra- sss changed in opposite directions, such that by the end tor models of working memory^{6,32,34} (Extended Data 495 ig. 1c). In contrast, in the unconstrained network, F 496 random and most amplifying mode inputs led to poor 497 decodability between early and late time points af-498 ter cue onset (Fig. 4b, bottom; center and right, offdiagonals indicated by pink rectangles are white/gray), 500 suggesting sequential activities during the early-to-501 late delay transition^{21,26,36}, and giving rise to the phe-502 nomenon of 'dynamic coding'^{8,10,11,14-16}. These activ-503 ities then stabilised during the late delay period as the 504 network dynamics converged to a persistent pattern of 505 activity (Fig. 4b, bottom; center and right, off-diagonals inside cyan squares are black). In sum, these decod-507 ing analyses were able to clearly distinguish between 508 persistent mode and random or amplifying inputs, but 509 not between the latter two. 510

To construct a targeted measure for identifying networks using most amplifying inputs, we exploited the fact that in large networks, random inputs are almost certainly guaranteed to have negligible overlap with any other direction in neural state space, including the most amplifying mode. Thus, we directly measured 516 the time courses of the overlap of neural activities with the top 25% most amplifying modes. We quantified 518 this overlap as the fraction of variance of neural activities that these modes collectively explained (Fig. 4c, 520 red lines; Methods 1.7.3). For a comparison, we also 521 measured these time courses for the overlap with the 522 top 25% most persistent modes (Fig. 4c, green lines). 523 We used the top set of amplifying and persistent directions, rather than just a single direction in each case, because large networks can harbor multiple directions that represent similarly persistent activity patterns and similarly efficient amplifying modes, respectively 52,53. In fact, in the more general setting in which more than two cues need to be discriminated (as is the case in 530 typical experiments; Fig. 1a) these higher dimensional subspaces will even be necessary so that they include the amplifying or persistent directions relevant for all cues.

As expected, for symmetric networks, persistent and equivalent amplifying mode inputs resulted in both 536 overlaps being high immediately from stimulus onset 537 (Fig. 4c, top; left and center). Random inputs started 538 with chance overlap which increased over time to ceil-539 ing as the network settled into its persistent (or, equiv-540 alently, most amplifying) mode (Fig. 4c, top right). In 541 unconstrained networks, persistent mode inputs led to constant high and moderate overlaps with the persistent and most amplifying modes, respectively (Fig. 4c, bottom left). Random inputs again started with chance 545 overlap for both modes, which then increased to the 546 same levels that resulted from persistent mode inputs (Fig. 4c, bottom right). In contrast, most amplifying in-548 puts were uniquely characterised by a cross-over between the time courses of the two overlap measures. Initially, neural activities overlapped strongly with the most amplifying mode (by construction), but showed only chance overlap with the persistent mode (Fig. 4c, bottom middle). Over time, these overlap measures

⁵⁵⁶ of the delay period overlap was high with the persistent mode and lower with the most amplifying mode 558 (Fig. 4c, bottom middle). Therefore, the cross-over of these overlap measures can be used as a signature of optimal information loading utilizing inputs aligned with ⁵⁶¹ the most amplifying modes. For example, modifying an earlier integrator model of working memory⁶ (Ex-562 tended Data Fig. 1c) so that inputs lie in a purely ran-563 domly oriented subspace can result in cross-temporal 564 decoding matrices that look similar to that achieved by 565 the most amplifying mode (Extended Data Fig. 8b), but the overlap measures that we developed here clearly reveal the lack of optimal information loading, even in that modified model (Extended Data Fig. 8e). In ad-569 dition, we confirmed in numerical simulations that the 570 same signature of optimal information loading remains detectable even under the practical constraints of experimental data analysis: when the underlying network dynamics is nonlinear, and only accessible indirectly by fitting linear dynamical models to the neural responses they generate (Extended Data Fig. 7d, 576 Methods 1.4.3 and Supplementary Math Note S3.4).

578 Signatures of optimal information loading in monkey IPFC

To study whether the PFC shows the dynamical signatures of optimal information loading that our theoretical analyses identified, we analysed a data set⁶² 582 of multi-channel recordings of the lateral prefrontal cortex (IPFC) in two monkeys during a variable-delay memory-guided saccade task (Fig. 1a). These recordings yielded 438 and 625 neurons (for monkeys K and T, respectively; Extended Data Fig. 9, Methods 1.1). We analysed the population dynamics of all recorded neurons in each monkey and applied the same metrics to this dataset that we applied to our models. Population dynamics appeared to show rich transient dynamics during the cue and early delay period, followed by relatively stable dynamics during the late delay period (Fig. 5a). This was reminiscent of the dynamics we found in unconstrained attractor networks following optimal information loading (Fig. 2h).

To further quantify this behaviour, we conducted decoding analyses. First, in line with previous studies^{10,14,15}, we found that a delay-trained decoder did not generalise to times outside of the delay period and in particular showed near-chance performance during the cue period (Fig. 5b). This was distinct from the pattern completion dynamics seen in classical attractor network models of working memory (Fig. 2f,I green and Fig. 4a green), but similar to that expected from random or optimal inputs in unconstrained networks (Fig. 2l black and red; Fig. 4a bottom, black and red).

Full cross-temporal decoding reinforced these results: decoders trained during the delay epoch did not generalise to the cue or go epochs and vice versa (Fig. 5c 610 and Extended Data Fig. 10a, pink rectangles). Thus, neural activity exhibited dynamic coding^{14,15} rather than the stable coding characteristic of simple pat-



Fig. 5 | Signatures of optimal information loading in monkey IPFC. a, Top: IPFC recording location. Bottom: neural firing rates (relative to the time-dependent but condition-independent mean) for one stimulus cue condition for 50 example neurons. See Fig. 1a (and Methods 1.1) for experimental paradigm. Neurons are ordered according to their firing rate at the end of the period shown. Vertical yellow lines indicate stimulus cue onset and offset. b, Performance of a delay-trained decoder (black bar indicates decoder training time period; Methods 1.7.4) on neural activity over time. Yellow ticks on horizontal axis indicate stimulus cue onset, offset, and go cue times, and the vertical gray bar indicates the stimulus cue epoch. Data is aligned to either stimulus cue onset (first 1.5 s) or to the go cue (final 1.5 s). Gray dotted lines show chance level decoding. c, Cross-temporal decoding of neural activity for monkey T (see Extended Data Fig. 10a for Monkey K). Yellow lines indicate stimulus cue onset, offset, and go cue times. Pink rectangles indicate poor generalization between time points (i.e. dynamic coding) and the cyan rectangle indicates examples of good generalization between time points (i.e. stable coding). The orange arrow indicates good same-time decoding during the cue epoch. The black vertical bar on the right indicates the delay-trained decoder training time period from b. d. Cross-validated quality of fits on 10 different held-out data splits when fitting 20-dimensional linear dynamical systems to neural activity (blue) and time shuffled controls (dark gray) (Methods 1.4.3). We also show quality of fits of the data against itself ('test vs. train'; light gray). e, Neural activity for each of the 6 cue conditions projected onto the top PC (solid lines) for monkey K (left) and monkey T (right). Solid lines show held-out test data from 1 split, dashed lines show predictions of fitted model dynamics. The inset for monkey T shows which color corresponds to each cue condition. f, Percent variance explained by the subspace spanned by either the 25% most persistent (green) or 25% most amplifying (red) modes as a function of time for the 20-dimensional linear dynamical systems fitted to data from monkey K (top) and monkey T (bottom). Gray lines show chance level overlap defined as the expected overlap with a randomly chosen subspace occupying 25% of the full space (median and 95% C.I. across 200 random subspaces). Comparisons shown in d and f use two-sided permutation tests (*, p < 0.05; **, p < 0.01; n.s., not significant; see Methods 1.7.3)

tern completion (Fig. 1c right; Fig. 4b top, and bot- 33 Fig. 10b-c). Further analyses showed that this orthogtom left; and Extended Data Fig. 1a-c right). Impor- and onality was not simply due to distinct sub-populations 615 tantly, same-time decoding performance was close to throughout the cue and delay epochs (Fig. 5c and 1 617 Extended Data Fig. 10a, orange arrow). This confirmed that the poor cross-temporal generalisation between early and late periods of a trial was not because the cue information had not yet reached PFC, or 621 was maintained by activity-silent mechanisms as pre-622 viously suggested^{11,41,46}. At the same time, also in line with previous studies^{8,10,14–16}, we found relatively 624 stable coding during the late delay period (Fig. 5c and 625 Extended Data Fig. 10a, cyan square). This ruled out purely sequential activity-based dynamics^{21,26,37,38,63} 627 Fig. 1d and Extended Data Fig. 1d).

Quantifying the relative alignment of the subspaces occupied by neural dynamics across time^{6,64} using 630 PCA confirmed the orthogonality of neural activities between different task epochs (Extended Data 651 to the cue and early delay epochs of our recordings

of neurons being active in different task epochs, but was instead largely due to changes in population-wide activities patterns^{10,65} (Extended Data Fig. 10d-e). 637

These results, in line with previous findings^{8,10,15,16}. clearly indicated that activities during the cue period were largely orthogonal from those during the delay period. However, these analyses alone were unable to distinguish between two fundamentally different information loading strategies PFC could employ: random input directions, or optimal information loading using specifically amplifying directions. Thus, in order to clearly identify the information loading strategy underlying the combination of dynamic and stable coding that we found, we applied our overlap measure 648 (Fig. 4c) to these PFC recordings. For this, we first fit-649 ted a 20-dimensional linear dynamical system model 650

(0–1 s after cue onset, Methods 1.4.3). We confirmed 710 a network, something that the brain will not have ex-652 that linear dynamics provided a reasonably accurate 653 cross-validated fit to the data compared to a time shuf-654 fled control (which destroyed the lawful dynamics of the data; Fig. 5d, dark gray, see also Methods 1.4.3), 656 and model-free train vs. test performance (which indicated that cross-validated errors were mostly due to 658 sampling noise differences between the train and test data; Fig. 5d, light gray) and recapitulated the most im-660 portant aspects of the trial-average dynamics in each condition (Fig. 5e).

We then performed the same overlap analysis on the fitted linear dynamics of the data that we used on 664 our simulated networks with linear dynamics (Fig. 4c; 665 Methods 1.7.3). As expected from our decoding analyses (Fig. 5b,c), the overlap of neural activities with 667 the most persistent modes was at chance initially and 668 gradually increased (Fig. 5f, green and Extended Data 669 Fig. 10i). Critically however, the overlap of neural ac-670 tivities with the most amplifying modes was high ini-671 tially and decreased with time (Fig. 5f, red and Ex-672 tended Data Fig. 10i).

We also noted that the overlap with the most amplify-674 ing directions became significantly lower than chance 675 over time. This suggests that PFC circuits may be more mathematically 'non-normal' 21,26,55,56,59 than the networks with randomly chosen weights that we used in Fig. 4. For example, Extended Data Fig. 8f shows 679 this phenomenon in a highly non-normal (purely feed-680 forward) network using optimal information loading 681 (see also Discussion).

As a control, repeating the same analyses on timeshuffled data, or on data taken from the late delay 684 period (when the network should already be near an attractor state) did not result in the same cross-over 686 pattern. In particular, there was a high initial over-687 lap with the most persistent modes and a chance (or below chance) initial overlap with the most amplify-689 ing modes (Extended Data Fig. 10f,g,i). Consistent 690 with these results, we found that at early times, stimulus information was at least as well (and even slightly better) decodable within the amplifying subspace than 693 in the persistent subspace (Extended Data Fig. 10h, 694 = 0), but became significantly better decodable in t 695 the persistent subspace at later times (Extended Data 696 Fig. 10h, t > 0). 697

Therefore, these analyses provide strong experimental evidence that PFC circuit dynamics utilize optimal information loading with inputs aligning with the most 700 amplifying modes (compare to Fig. 4c; bottom middle 701 and Extended Data Fig. 10i, third vs. fourth row) rather 702 than simply using random input directions (compare to 703 Fig. 4c; bottom right and Extended Data Fig. 10i, first 704 vs. fourth row). 705

Information loading in task-optimized non-linear 706 networks

⁷¹¹ plicitly when performing a working memory task. In ⁷¹² addition, the formal equivalence of optimal information loading to using the most amplifying direction as an in-714 put could only be demonstrated for networks with lin-715 ear dynamics receiving instantaneous inputs, while fix-716 ing the magnitude of those inputs. Thus, an important question is whether optimizing simple task-relevant cost functions^{13,17,20,39,40,66}, under only a generic en-718 ergy constraint^{13,39,40,66}, without explicitly encouraging optimal information loading, or fixing input mag-720 nitudes, can be sufficient for the underlying circuits to adopt an optimal information loading strategy.

We trained nonlinear recurrent neural networks (with-⁷²⁴ out connectivity constraints Fig. 6a; Methods 1.3.2) on ⁷²⁵ the same memory-guided saccade task as that which 726 our animals performed (Fig. 1a). Specifically, during the cue epoch, these networks received temporally extended input from one of six input channels on any given trial depending on the cue condition (Fig. 6a. left). Similarly, network activity was decoded into one 730 of six possible behavioural responses via six readout channels (Fig. 6a, right). Following previous approaches^{13,39,40}, all recurrent weights in the network, as well as weights associated with the input and readout channels, were optimized, while only penalising the average magnitude of neural responses over the course of the whole trial (Methods 1.3.3).

To study the generality of optimal information loading, 738 we first implemented two standard cost functions that have been widely used in previous work^{13,17,24,39,40}. 740 These cost functions required networks to stably maintain cue information either immediately after cue onset (cue-delay; Fig. 6b, left), or only at response time (after-go; Fig. 6b, center). However, we reasoned that neither of these standard cost functions may be ap-745 746 propriate for understanding PFC function. The cuedelay cost is well justified when stimuli need to be decoded potentially instantaneously after cue onset, and as such it is most relevant for sensory areas⁵⁸. Conversely, the after-go-time cost may be most directly relevant for motor areas, by only requiring stable coding during the short response period 66. Therefore, we also considered a third cost function that required sta-⁷⁵⁴ ble coding just in time before the go cue appeared, ⁷⁵⁵ i.e. during a period that was divorced from the stimu-⁷⁵⁶ lus or response time windows, and as such was more consistent with the putative role of PFC in cognitive flexibility^{2,25,60} (just-in-time; Fig. 6b, right). 758

All trained networks achieved high performance, as measured by a late-delay decoder, in line with what ⁷⁶¹ their respective cost functions required: already by the ⁷⁶² end of the cue epoch for the cue-delay cost (Fig. 6c ⁷⁶³ and Extended Data Fig. 11a, green), only after cue ⁷⁶⁴ offset but for most of the delay period for the just-intime cost (Fig. 6c and Extended Data Fig. 11a, red), 765 or only shortly before go time for the after-go-time cost 766 (Fig. 6c orange and Extended Data Fig. 12b).

Our definition of optimal information loading relied on 768 We then further analyzed the dynamics with which 709 full access to the algebraic form of the dynamics of 709 these networks achieved competent performance.



Fig. 6 | Information loading in task-optimized non-linear networks. a, Illustration of a recurrent neural network model with unconstrained connectivity (middle). During the cue epoch, networks received input from one of six input channels on any given trial depending on the cue condition (left). Network activity was decoded into one of six possible behavioural responses via six readout channels (right). All recurrent weights in the network (50 neurons), as well as weights associated with the input and readout channels, were optimized (Methods 1.3.2). b, Illustration of cost functions used for training. Yellow ticks indicate cue onset and offset times, yellow bars indicate range of go times in the variable delay task. Boxcars show intervals over which stable decoding performance was required in three example trials with different delays for each of the cost functions considered (Methods 1.3.3): cue-delay (left), after-go-time (center), or just-in-time (right). c, Performance of a delay-trained decoder (black bar indicates decoder training time period; Methods 1.7.4) on model neural activity over time in trials with a 1.75 s delay. Yellow ticks show stimulus cue onset, offset, and go times, and the vertical gray bar indicates the cue epoch. Neural activities were generated by networks optimized for the cue-delay (green), after-go-time (orange), or just-in-time (red) costs. Solid lines and shading indicate mean±1 s.d. across 10 networks. Gray dotted line shows chance level decoding. d, Cross-temporal decoding of model neural activity for cue-delay (left), after-go-time (center), and just-in-time (right) trained models. Yellow lines indicate stimulus cue onset, offset, and go times. The black vertical bar on the right indicates the delay-trained decoder training time period from c. e. Percent variance explained by the subspace spanned by either the 25% most persistent (green) or 25% most amplifying (red) modes as a function of time for 20-dimensional linear dynamical systems fitted to the model neural activities of networks optimized for the cue-delay (left), after-go-time (center), or just-in-time cost (right). Gray lines show chance level overlap defined as the expected overlap with a randomly chosen subspace occupying 25% of the full space. Lines and error bars show mean±1 s.d. over 10 networks.

The cue-delay network showed signatures of attrac- 799 e, right). Again, the apparent lack of attractor dynam-770 temporal decoding was high at all times, including between the cue and delay epochs (Fig. 6d, left; cf. Fig. 1c, Extended Data Fig. 1a-c; see also Extended Data Fig. 11d for state-space plots). Although there was a cross-over of our overlap measures, critically, neural activities were already aligned well above chance with the most persistent modes immediately 778 following cue onset (Fig. 6e, left). This was consistent with these networks being explicitly required to exhibit 780 stable coding at all times by the cue-delay cost. We 781 also found similar dynamics for networks that optimize а 'full-delay' cost, in which cue information must be 783 stably maintained only after cue offset (Extended Data 784 Fig. 13, Methods 1.3.3). 785

786 make particular use of attractor dynamics. Instead, 787 it generated largely sequential activities, i.e. pure dy-788 namic coding akin to the dynamics of a feedforward 789 790 the very end of the delay period (Fig. 6d, center; cf. 792 794 795 purely sequential dynamics (Extended Data Fig. 12c- ser to the early decodability requirement of the cue-delay

tor dynamics with simple pattern completion: cross- soo ics was well explained by the cost function not re-⁸⁰¹ quiring any stable coding during the delay period. In ⁸⁰² summary, network dynamics trained for standard cost ⁸⁰³ functions recovered classical network models of work-⁸⁰⁴ ing memory (Fig. 1c,d and Extended Data Fig. 1a-⁸⁰⁵ d), but were different from those seen in experimental ⁸⁰⁶ recordings^{8,10,14–16,25} (Fig. 5b.c.f).

807 In contrast to both standard training costs, just-in-⁸⁰⁸ time networks showed the signatures of attractor dynamics with dynamic coding: cross-temporal decoding was poor between early and late periods of a trial. but high during the late delay period (Fig. 6d, right; Extended Data Fig. 11b; cf. Fig. 4b, bottom center, Fig. 5c, Extended Data Fig. 10a; see also Extended ⁸¹⁴ Data Fig. 11d for state-space plots), and the overlap At the other extreme, the after-go-time network did not as of neural activities with the most amplifying and persistent modes showed the characteristic cross-over 816 that we predicted theoretically and found experimens18 tally (Fig. 6e, right; cf. Fig. 4c, bottom center, Fig. 5f, network: cross-temporal decoding was only high at sty Extended Data Fig. 7d, bottom right). Note that the main difference from the dynamics of cue-delay net-Fig. 1d and Extended Data Fig. 1d, right; see also Ex- works was that the overlap with persistent modes was tended Data Fig. 12e for state space plots), and the sez lower initially, and increased more slowly afterwards. overlap with the most persistent modes never signifi- seas This was consistent with the speed-accuracy trade-off cantly exceeded that with the most amplifying modes and we saw earlier in unconstrained linear integrator net-(Fig. 6e, center). This was particularly the case for s25 works (Fig. 3e), whereby achieving high overlap with a fixed delay task, for which this cost function yielded and the persistent mode early during the trial (analogous

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cost) is achieved by inputs aligned with the persistent was We expect all these weakly nonsymmetric networks to 828 mode, while a late overlap (analogous to the require-829 ments of the just-in-time cost) is optimized by the most 830 amplifying mode. 831

In summary, all task-optimized networks exhibited a 832 key feature of optimal information loading: they made 833 use of most amplifying modes early during the trial 834 (Fig. 6e, all red lines start high at 0 s). The extent to 835 which they showed the complete cross-over of ampli-836 fving and persistent overlaps predicted by our earlier 837 analyses (Fig. 4c, bottom center), and characteristic of the experimental data (Fig. 5f), was consistent with ow much they were required to exhibit stable cod-840 ng^{8,10,11,14-16}. These results suggest that optimal in-841 formation loading emerges naturally as a dynamical strategy in task-optimized networks, without explicit re-843 quirements on their inputs. 844

Discussion 845

While attractor networks have been proposed 846 underlie a number of core cognitive functo 847 tions 12,17,17,27,29,30,33-35,43,44,50,67-69 prominently in-848 cluding working memory^{5-7,27-32}, their operation was 849 almost exclusively analyzed in terms of how their intrinsic connectivity supports information mainte-851 nance^{5,7,12,27,30,33,34,43,70,71} (but see Refs. 6,32, dis-852 cussed below). We instead studied information 853 loading by external inputs in attractor networks and 854 showed that optimal information loading provides a 855 normative account of the widely observed and puz-856 zling phenomenon of dynamic coding^{8,10,14–16}. We 857 predict that these results should also generalise to more cognitively demanding working memory tasks in which, unlike in the simple memory-guided saccade 860 task we studied here, the correct response is unknown 861 during the delay period, thus requiring the mainte-862 nance of stimulus information before a response can 863 be prepared^{14,15,23,72,73}. Indeed, strongly dynamic 864 population activity, similar to those that we identified 865 nere, has been observed in monkey PFC^{10,14–16,23,24,73} ł 866 and in neural networks^{20,24,39} trained on such tasks. 867

or understanding the dynamics of optimal informa-F 868 tion loading, we used networks whose connectiv-869 ity was constrained to be symmetric as a pedagog-870 ical stepping stone. Such networks are among the 871 most influential and best understood models of at-872 tractor dynamics in general^{35,44}, and of working mem-873 ory more specifically 5,7,12,29,31,33,34,42-46, and they continue to form the basis of many recent models in the 875 field^{5,33,45,46}. We showed that such networks show 876 limited (if any) dynamic coding, and essentially none 877 with optimal information loading. There are also mod-878 els whose connectivity is not strictly symmetric at the 879 microscopic level of cell-to-cell connections, but their 880 macroscopic connectivity (at the level of connections between groups of similarly tuned cells) is strongly 882 symmetric^{71,74}. In other models, excitatory cells are 883 connected symmetrically, but total symmetry is bro-884 ken by the introduction of effectively a single inhibitory neuron providing a uniform, global level of inhibition^{5,7}.

exhibit largely stable coding. Instead, we showed that 888 dynamic coding naturally arises in networks whose 889 connectivity is not constrained to be symmetric, and 890 especially so under optimal information loading.

Our dynamical analysis revealed a novel, theoretically-892 grounded aspect of dynamic coding: not only should neural activities during the cue and early delay period ⁸⁹⁵ be orthogonal to those during the late delay period, ⁸⁹⁶ but they should be orthogonal in the specific direc-897 tions that are aligned with the (locally) most amplifying directions. We found strong evidence for these 898 predictions of optimal information loading in PFC dur-899 ing a memory-guided saccade task. These results unify previous, seemingly conflicting models of work-901 ing memory maintenance that typically either use at-902 tractor dynamics^{5,6,27} or rely on sequential activities 903 often generated by non-normal dynamics^{21,26,36,37}. 904 We found that although both classes of models can 905 capture select aspects of neural data (e.g. sequential models can capture early delay activity whereas 907 attractors are better suited to capturing late delay ac-908 tivity), no model could capture the experimentally ob-909 served rich combination of sequential and persistent 910 dynamics⁷² (Fig. 1; see also³⁹). We showed that op-911 timal information loading in attractor models with realistic, unconstrained connectivity, leads to the specific combination of sequential and persistent dynam-914 ⁹¹⁵ ics that has been observed in experiments. We found that this was true across a range of different specific 916 network architectures: either hand-set (Figs. 3 and 4 917 and Extended Data Fig. 5a,b) or optimized stimulus 918 inputs (Extended Data Fig. 5c,d); nonlinear discrete 919 attractors models (Figs. 2 and 6 and Extended Data Figs. 2, 7 and 11–13); and a nonlinear ring/bump at-921 922 tractor model (Extended Data Fig. 3).

In contrast to our optimal information loading-based 923 account, previous attempts to reconcile transient and 924 persistent dynamics specifically proposed that tran-925 sient dynamics do not affect the coding of the stimulus 926 information^{6,32}. These 'stable coding' dynamics are 927 very different from dynamic coding as observed in experiments^{3,8,10,11,14–24}, and as predicted by our theory 929 of optimal information loading. Put simply, in previous 930 models, the stimulus input drives network activity to-⁹³² wards the desired persistent state. In real data, and in models that exhibit optimal information loading, stimu-⁹³⁴ lus inputs drive network activity orthogonal to the desired persistent state (and instead specifically in a direction that is aligned with the most amplifying mode) 936 before activity ultimately settles into the correct state. 937

There are aspects of the data that were not reproduced accurately by any of the specific models we implemented. First, the overlap with the most amplify-940 ing directions became significantly lower than chance 941 over time in the data. This suggests that PFC circuits may be more mathematically 'non-normal' (i.e. in-943 clude stronger feedforward loops^{21,26,56}, or excitatory-944 inhibitory interactions^{52,55}) than the networks with ran-945 domly chosen or initialised weights we used here^{59,61}.

(For example, we found that networks with strong feed- 1007 space¹⁷; the dynamics during the stimulus period had forward connectivity reproduced this phenomenon; 1005 0 correlation with the late delay activity²⁴; and cross-948 Extended Data Fig. 8f.) Second, the time evolution of the temporal decoding of time revealed strongly sequen-949 the overlaps with the most persistent and most am- 1010 tial dynamics in a variety of tasks²⁰ (see also^{39,41}). 950 plifying modes seemed to obey different time con- 1011 In fact, these features of model activities were also stants, with the persistent overlap evolving substan- 1012 shown to be reflected in the corresponding experimen-952 tially slower than the amplifying overlap. This may 1013 tal data in each case 17,20,24. Nevertheless, it remained 953 be a result of high dimensional, graded dynamical 1014 unclear whether these features were epiphenomenal transitions between multiple amplifying and persistent 1015 or an integral part of the functioning of these networks. modes compared to the less complex dynamical tran- 1016 Our results suggest that optimal information loading sitions that we observed in our models. 957

1018 There have been multiple mechanisms proposed to account for some of the features of the data that 959 seem to be at odds with basic attractor network dy-960 1019 namics, most prominently dynamic coding^{14,15}. These 961 hypothetical mechanisms include short-term plastic-962 1020 ity^{11,23,39,41,46,75}, specific changes in the strength of 963 1021 input and recurrent connections^{45,76}, and separate 964 1022 stimulus- and delay-responsive cells^{3,10}. We showed 1023 that the core phenomenon of dynamic coding emerges 966 1024 naturally, without any of these additional mechanisms, 967 1025 from the same ultimate principle that explains per-968 1026 sistent activities (robust memory maintenance imple-969 1027 mented by attractor dynamics). Moreover, the high 1028 initial overlap with the most amplifying modes, which 971 was a core prediction of our theory confirmed by the 1030 data, is not specifically predicted by any of these alter-973 1031 native mechanisms. Nevertheless, these mechanisms 974 1032 are not mutually exclusive with ours. In fact, they might 1033 help explain the more nuanced aspects of the data that 976 1034 our specific network implementations did not capture 977 1035 (see above), as well as aspects of the data that lie 1036 outside the scope of our theory (e.g. activity silent in-1037 formation maintenance during inter-trial intervals⁴⁶). 980

1038 A number of recent studies of neural network dynam-1039 981 ics have analysed the relationship between the direc-982 tion of inputs and the magnitude of responses they 1041 083 evoke^{52,56,61}. However, these studies have typically 1042 focused on networks with transient dynamics, such as 1043 985 those relevant for perception⁵⁸, or motor control^{52,61}. 1044 986 In particular, Ref. 61 found that optimal inputs (result-987 ing in the largest transients) are typically orthogonal to 1046 the activity patterns that the network expresses in re-989 sponse to them, providing a normative account for the 1048 990 experimentally observed orthogonality of preparation 1049 991 and execution subspaces in motor cortex^{64,77}. Our 1050 992 work suggests that the use of optimal inputs to drive 1051 993 network dynamics, and the orthogonality of those in-994 puts to network responses, is a more general principle 1053 of cortical circuits, extending beyond the motor cortex. 1054 particular, our results demonstrate the importance 1055 10. Spaak, E., Watanabe, K., Funahashi, S. & Stokes, In 997 of optimal initialization even when the transients fol- 1056 998 lowing initialization themselves may be irrelevant, as 1057 information is ultimately maintained by stable attractor 1058 1000 states. 1001 1059 1060 In line with our results, previous studies optimizing net-

works on related tasks requiring persistent, rather than 1003 1062 transient, responses also exhibited key features of dy-1004 1063 namic coding: neural activities initially pointed strongly 1005

orthogonal to the ultimate attractor location in state 1065

1017 could provide a unifying principle for these observations.

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Author Contributions

J.P.S., M.L., and M.G.S. conceived the study. K.W. 1367 performed all experimental recordings, T.S. assem-1334 1368 bled the neural recording system, and K.W. and T.S. 1369 performed data pre-processing. J.P.S. and M.L. devel-1336 oped the theoretical framework, performed analytical derivations, and wrote the first draft of the manuscript. 1338 J.P.S. performed all numerical simulations, analysed the data, and produced the figures. J.P.S., M.L., and M.G.S. interpreted the results. All authors revised the 1374 able in the following repository upon peer-reviewed final manuscript.

Competing Interests statement. The authors declare no competing interests.

Reporting Summary. Further information on re-1345 search design is available in the Life Sciences Reporting Summary linked to this article. 1347

Randomization. No new experimental data was gathered for this paper. There is no group allocation in this study. Trial types were randomly determined by a computer program.

1352 Blinding. As data collection had been performed well 1353 before our theory of optimal information loading was developed and our corresponding analyses were per-1355 formed, it was effectively blind to the purposes of our 1356 study. Data analysis was not performed blind to the 1357 conditions of the experiments.

1358 Data exclusion. As described below (Methods 1.1.4), 1 neuron from monkey K's dataset was removed from all analyses because it was recorded in fewer than 10 trials for at least one stimulus cue condition. As also described below (Methods 1.1.4), randomly generated linear networks with unconstrained connectivity in Fig. 4 were excluded if the inner product between the most amplifying mode and persistent mode was 1365 greater than 0.2 (i.e. we only kept networks that were 1366 relatively mathematically non-normal).

All experimental data will be Data availability. made available in the following repository upon peer-1370 reviewed publication: https://github.com/jakepstroud.

1371 Code availability. All code was custom written in ¹³⁷² Python using NumPy, SciPy, Matplotlib, Scikit-learn, 1373 and Tensorflow libraries. All code will be made avail-1375 publication: https://github.com/jakepstroud.

1 Methods

1377 **1.1** Experimental materials and methods

Experimental methods have been described before⁶² and largely followed those used in our previous publications^{10,78,79}. We briefly summarize the methods below.

1380 1.1.1 Subjects and apparatus

We used two female macaques (monkey T, Macaca mulatta, 5 kg; monkey K, Macaca fuscata, 8 kg). Both 1381 monkeys were housed individually. The light/dark cycle was 12/12 hr. (light, from 8:30 a.m. to 8:30 p.m.). The 1382 monkeys sat guietly in a primate chair in a dark, sound-attenuated shield room. During both training and neural recording sessions, we restrained the monkeys' head movement non-invasively using a thermoplastic head cap as described in⁸⁰. This head cap is made of a standard thermoplastic splint material (MT-APU, 3.2 mm thick, 1385 CIVCO Radiotherapy, IA., USA), and was molded out so that it conformed to the contours of the animals' scalp, 1386 cheek bone, and occipital ridge. Visual stimuli were presented on a 17 inch TFT monitor placed 50 cm from 1387 the monkeys' eyes. Eye movements were sampled at 120 Hz using an infrared eye tracking system (ETL-200, 1388 ISCAN, MA.). Eye fixation was controlled within a 6.5° imaginary square window. TEMPO software (Reflective 1389 Computing, WA.) was used to control behavioral tasks. All experimental procedures were approved by the Animal 1390 Research Committee at the Graduate School of Frontier Biosciences, Osaka University, Japan and were in full compliance with the guidelines of the National BioResource Project 'Japanese Macagues'. Experimental work performed in non-human primates that was not funded by Wellcome may not adhere to the principles outlined in 1393 the NC3Rs guidance on Non-human Primate Accommodation, Care and Use. 1394

1395 1.1.2 Behavioral task

The monkeys were trained on a memory-guided saccade task requiring them to remember the location of a visual 1396 stimulus cue on a screen and to make a correct eye movement after a delay period (Fig. 1a). Specifically, this 1397 task required monkeys to fixate on a central ring for a period of 2.6-7.4 s followed by a stimulus cue (a white 1398 square) appearing in one of six pre-determined locations for 0.25 s. After a variable delay period of 1.4-7.5 s. the fixation ring was replaced by placeholders at all six possible stimulus cue locations (go cue). Monkeys were 1400 required to make a saccade within 0.5 s to the placeholder where the original stimulus cue was presented and maintain their gaze for 0.25 s for monkey T and either 0.25 s or 0.6 s for monkey K (these two gaze maintenance 1402 times were switched in different blocks for monkey K) to receive a juice reward. The monkeys were extensively 1403 trained, with close to perfect performance (monkey T, 96.1%; monkey K, 96.3%, mean across sessions). Fixation 1404 break errors were excluded from the calculation of percent correct rate.

1406 1.1.3 Recordings

After training was completed, we conducted an aseptic surgery under general anesthesia. We stereotypically implanted a plastic recording chamber on the lateral surface of the prefrontal cortex, under the guidance of structural MRI images (Extended Data Fig. 9). In monkey T, we implanted a cylindrical chamber (RC-T-S-P, internal diameter 12.7 mm, Gray Matter Research, MT.) in the right hemisphere (AP = 33, ML = 14.5; AP, anteriorposterior; ML, medio-lateral). A 32-channel semi-chronic microdrive system (SC-32, Gray Matter Research) was mounted inside this chamber. In monkey K, we implanted a cuboid chamber (width 12 mm, depth 16 mm, height, 15 mm, S-company Itd., Tokyo, Japan) over the principal sulcus in the left hemisphere.

We collected neural data in a total of 48 daily sessions (21 in monkey T; 27 in monkey K). In monkey T, we used the 32-ch microdrive (SC-32) that housed 32 single-contact tungsten electrodes with inter-electrode spacing of 1.5 mm. In monkey K, we used a 32-ch linear microelectrode array (Plexon U-Probe, Plexon, TX.) with an interelectrode spacing of 150 μ m along a single shaft. We positioned the U-Probe by using a custom-made grid (width 12 mm, depth 16 mm, height, 10 mm) which had a total of 165 holes with 1 mm spacing. We advanced the U-Probe by a custom-made hydraulic microdrive (S-company Itd.).

Raw extracellular neural signals were amplified and recorded in reference to a titanium bone screw at the vertex (in monkey T) or the shaft of the linear array (monkey K) using a neural signal amplifier RZ2 Bioamp Processor (Tucker-Davis Technologies, FL.). Behavioral data (task-event information and eye-movement information) were also sent to the RZ2 Bioamp. Neural data acquisition was performed at a sampling frequency of 24414.08 Hz, and behavioral data acquisition at 1017.25 Hz. For analysis of spiking activity, the raw neural signal was filtered (300 Hz to 6 kHz) for offline sorting (Offline Sorter, Plexon). In monkey T, approximately three hours before each recording session, we took the monkey to the testing room and advanced each electrode in the SC-32

¹⁴²⁷ by a minimum of 62.5 μ m in order to ensure recording of new neurons. We then put the monkey back in the ¹⁴²⁸ home cage until we brought it out again for the recording session. In monkey K, we adopted the method of the ¹⁴²⁹ U-Probe insertion reported in⁸¹. We first punctuated the dura using a guide tube (a shortened 23 gauge needle), ¹⁴³⁰ and inserted the U-Probe array slowly, usually with a step of 500 μ m. We kept monitoring electrocardiogram ¹⁴³¹ (pulsatory fluctuation) on superficial electrodes to identify the point of cortical entry. We usually left 3–5 superficial ¹⁴³² channels outside the cortex. After array insertion, we waited 1–1.5 hours until the recorded single-unit and ¹⁴³³ multiunit activities indicated that the electrode array was stably positioned in the cortex. While waiting, the monkey ¹⁴³⁴ watched nature and animal video clips and received a small snack on a monkey chair.

In monkey T only, to determine location of the frontal eye field (FEF), and confirm that our recording area was outside it, intracortical microstimulations (22 biphasic pulses, 0.2 ms duration at 333 Hz, \leq 150 μ A) were applied through microelectrodes. When eye movements were elicited below 50 μ A, the site was considered to be in the low-threshold FEF. In monkey T, our recording area did not include the low-threshold FEF.

1439 1.1.4 Pre-processing

We excluded neurons that were recorded in fewer than 10 trials for any cue condition. For each monkey, we pooled neurons from all recording sessions to create pseudopopulations of 438 neurons for Monkey K (after we removed 1 neuron from monkey K's dataset due to an insufficient number of trials) and 625 neurons for Monkey T (no neurons were removed from monkey T's dataset). To compute neural firing rates, we convolved spike trains with a Gaussian kernel with a standard deviation of 25 ms. Trial-averaged trajectories of time-varying mean firing rates were computed separately for each neuron and each cue condition. For analysis methods that used crossvalidation (see below), we split trials into separate train and test sets with a 1:1 train:test ratio, and computed trial-averaged trajectories for each training and test set (using 1:1 splits). For non-cross validated analyses, we either computed trial averages based on all the data, or on a subset of the data (see below). We aligned neural activity to either stimulus or go cue onset (see also below in Methods 1.7) and shifted activity by -50 ms to allow for the delay in time for information about these cues to enter PFC. For consistency with our simulations (see below), we subsampled neural firing rates at a 1-ms time resolution.

1452 1.2 Neural network models: overview

All our simulated networks (Figs. 2–4 and 6 and Extended Data Figs. 2–5, 7 and 11–13) evolved according to a canonical model of stochastic recurrent neural circuit dynamics^{40,82}:

$$\tau \frac{\mathrm{d}\mathbf{x}^{(c)}(t)}{\mathrm{d}t} = -\mathbf{x}^{(c)}(t) + \mathbf{W} \,\mathbf{r}^{(c)}(t) + m_{\mathrm{h}}(t) \,\mathbf{h}^{(c)} + m_{\mathrm{g}}(t) \,\mathbf{g} + \mathbf{b} + \sigma \,\boldsymbol{\eta}^{(c)}(t) \tag{1}$$

$$dt = \mathbf{r}(t) + \mathbf{w} + \mathbf{h}(t) + \mathbf{$$

where $\mathbf{x}^{(c)}(t) = (x_1^{(c)}(t), \dots, x_N^{(c)}(t))^{\top}$ corresponds to the vector of (unitless) 'subthreshold', i.e. coarse-grained (lowpass filtered), trial-averaged raw somatic membrane potentials of the N neurons of the network⁸² in cue condition = 1,..., C (initialised at $\mathbf{x}^{(c)}(t_0)$ at the beginning of the simulation t_0 , which could be at or before stimulus onset С 1455 t = 0, $\mathbf{r}^{(c)}(t)$ is their momentary (trial-averaged) firing rates, with $\mathbf{f}(\mathbf{x})$ being the activation function that converts at membrane potentials to firing rates, au is the membrane time constant, W is the recurrent weight matrix (shown 1457 e.g. in Fig. 2a and g), $\mathbf{h}^{(c)}$ is the input given to the network depending on the stimulus cue, g is the stimulus-cueindependent go cue that occurs at the go time t_{go} , $m_h(t)$ and $m_g(t)$ are box car 'masking' kernels such that the stimulus and go cues are only effective within a limited period at the beginning and end of the trial, respectively, b is a cue-independent bias, σ is the standard deviation of the noise process, and $\eta^{(c)}(t)$ is a sample from a standard 1461 (mean 0 and variance 1) Gaussian white (temporally and spatially) noise process. Note that $\eta^{(c)}(t)$ represents the 1462 effective noise corrupting trial-averaged trajectories (rather than the noise corrupting individual trials). 1463

Networks shown in different figures corresponded to different special cases of Eqs. 1 and 2 (see Table 1). Specif-1464 ically, for linear networks (Figs. 3 and 4, Fig. 5d,e,f, Fig. 6e, Extended Data Fig. 7d, Extended Data Fig. 4, 1465 Extended Data Fig. 5, and Extended Data Fig. 10f,g) f(x) = x was the identity function 6,21,26,32,34,43. For nonlinear 1466 networks (Figs. 2 and 6 and Extended Data Figs. 2 and 11–13) $f_i(\mathbf{x}) = [x_i]_+$ was the rectified linear (ReLU) acti-1467 vation function applied element-wise 39,40,61 (except for the ring attractor networks where we used $f_i(\mathbf{x}) = \tanh(x_i)$; 1468 Extended Data Fig. 3). Given that the focus of our study was optimal information loading, stimulus inputs were 1469 either optimized numerically (Fig. 2, Fig. 6, Extended Data Figs. 2 and 3, Extended Data Fig. 5c,d, and Extended Data Figs. 11–13), or set to analytically computed values as dictated by our mathematical analysis (Figs. 3 and 4, Extended Data Fig. 4c,d, and Extended Data Fig. 5a,b), or as a baseline, set to random (or quasi-random, see 1472 below) values (Figs. 3 and 4, Extended Data Fig. 4a,b, and Extended Data Fig. 5a,b). For networks used to study the effects of instantaneous initial conditions (Figs. 2 and 3 and Extended Data Figs. 2-5 and 7), the stimulus masking kernel was zero and instead the initial condition was set to the stimulus input; for other networks (Fig. 4, Fig. 5e-f, Fig. 6 and Extended Data Figs. 11-13) the stimulus masking kernel was a boxcar between 0 1476

Symbol	Fig. 2	Fig. 3a–c	Fig. 3d–e	Fig. 4	Units	Description
N	50	2	1000	100	-	number of neurons
t_0	0	0	0	-0.5	s	simulation start time
t _{go}	-	-	-	-	s	go cue time
t _{max}	2	2	2	2.5	s	simulation end time
au	0.05	0.05	0.05	0.2	s	membrane time constant
$\mathbf{x}^{(c)}(t_0)$	h ^(c)	$\mathbf{h}^{(c)}$	h ^(c)	0	-	initial condition
$f(\cdot)$	nonlinear ^a	linear ^a	linear ^a	linear ^a	Hz	neural activation function
W	optimized ^b	set^d	$\overset{\text{iid.}}{\sim} \mathcal{N}(0, \frac{1}{N})^e$	$\stackrel{\text{iid.}}{\sim} \mathcal{N}(0, \frac{1}{N})^e$	s	weight matrix
С	6	1	1	2	-	number of stimuli
$\mathbf{h}^{(c)}$	optimized ^{c1}	set ^f	set ^b	set ^b	-	stimulus input
g	-	-	-	-	-	go cue
$m_{\rm h}(t)$	0	0	0	$K(0, 0.25)^{g}$	-	stimulus masking kernel
$m_{g}(t)$	0	0	0	0	-	go cue masking kernel
b	optimized ^b	0	0	0	-	cue-independent bias
σ	0.05	0	0	variable ^b	-	noise standard deviation
Symbol	Fig. 5e-f	Fig. 6a–d and	Fig. 6e	Extended Data	Units	Description
	0.00	Extended Data	0.00	Fig. 2		I
		Fig. 1e		U		
Ν	20	50	20	50	-	number of neurons
t_0	0	-0.5	0	0	s	simulation start time
$t_{\rm go}$	-	2	-	-	s	go cue time
$t_{\rm max}$	1	3	1	2	s	simulation end time
au	0.05	0.05	0.05	0.05	s	membrane time constant
$\mathbf{x}^{(c)}(t_0)$	0	0	0	h ^(c)	-	initial condition
$f(\cdot)$	linear ^a	nonlinear ^a	linear ^a	nonlinear ^a	Hz	neural activation function
W	fit ^b	optimized ^b	fit ^b	optimized ^b	s	weight matrix
С	6	6	6	6	-	number of stimuli
$\mathbf{h}^{(c)}$	fit^{b}	optimized ^{c2}	fit^b	optimized ^{c1}	-	stimulus input
g	-	$\sum_{c} \mathbf{h}^{(c)}$	-	-	-	go cue
$m_{\rm h}(t)$	$K(0, 0.25)^{g}$	$\widetilde{K}(0, 0.25)^g$	$K(0, 0.25)^{g}$	0	-	stimulus masking kernel
$m_{\sigma}(t)$	0	$K(t_{ro.}, t_{ro.} + 0.5)^{g}$	0	0	-	go cue masking kernel
b	fit ^b	optimized ^b	fit ^b	optimized ^b	-	cue-independent bias
σ	0	0.05	0	0	-	noise standard deviation
Symbol	Extended	Extended Data	Extended Data	Extended Data	Units	Description
e y me er	Data Fig. 3	Fig. 7b	Fig. 7c	Fig. 7d	omis	Debenphon
N	50	50	50	20	-	number of neurons
t_0	0	0	0	0	s	simulation start time
$t_{\sigma 0}$	-	-	-	-	s	go cue time
$t_{\rm max}$	2	2	2	2	s	simulation end time
au	0.05	0.05	0.05	0.05	s	membrane time constant
$\mathbf{x}^{(c)}(t_0)$	h ^(c)	h ^(c)	h ^(c)	h ^(c)	-	initial condition
$\mathbf{f}(\cdot)$	nonlinear ^a	linear ^a	nonlinear ^a	linear ^a	Hz	neural activation function
Ŵ	optimized ^b	optimized ^b	optimized ^b	fit^{b}	s	weight matrix
С	36 (6) ^h	6	6	6	-	number of stimuli
$\mathbf{h}^{(c)}$	optimized ^{c2}	optimized ^{c1}	optimized ^{c1}	fit^{b}	-	stimulus input
g	-	-	-	-	-	go cue
$m_{\rm h}(t)$	0	0	0	0	-	stimulus masking kernel
$m_{g}(t)$	0	0	0	0	-	go cue masking kernel
b	optimized ^b	optimized ^b	optimized ^b	fit^b	-	cue-independent bias
σ	0.05	0	0.05	0	-	noise standard deviation

Table 1 |Continued on text page.

Symbol	Extended	Data	Extended	Data	Ex	tended	Data	Units	Description
	Fig. 4a,b	Fig. 4c,d			Fig. 5a,b				-
Ν	100 100		10, 100, 1000		-	number of neurons			
t_0	0 0		0	0		s	simulation start time		
t _{go}	-		-		-			S	go cue time
t _{max}	2		2	2			s	simulation end time	
au	0.05		0.05	0.05		S	membrane time constant		
$\mathbf{x}^{(c)}(t_0)$	h ^(c)		h ^(c)	$\mathbf{h}^{(c)}$ $\mathbf{h}^{(c)}$		n ^(c)		-	initial condition
$f(\cdot)$	linear ^a		linear ^a		lin	linear ^a		Hz	neural activation function
W	$\stackrel{\text{iid.}}{\sim} \mathcal{N}(0, \frac{1}{N})^e$		$\overset{\text{iid.}}{\sim} \mathcal{N}(0, \frac{1}{N})^e$		$\stackrel{\text{iid.}}{\sim} \mathcal{N}(0, \frac{1}{N})^e$		s	weight matrix	
С	1		1		1	1		-	number of stimuli
$\mathbf{h}^{(c)}$	$\stackrel{ ext{iid.}}{\sim} \mathcal{N}ig(0, rac{1}{N}ig)$		optimized ^{c4}		set	set ^b		-	stimulus input
g	-		-		-	-		-	go cue
$m_{\rm h}(t)$	0		0	0		0		-	stimulus masking kernel
$m_{\rm g}(t)$	0		0		0		-	go cue masking kernel	
b	0		0		0	0		-	cue-independent bias
σ	0		0 0		0	0		-	noise standard deviation
Symbol	Extended	Exten	ded Data	Extended Extended Data		Units	Description		
	Data Fig. 5c,d	Data Fig. 5c,d Fig. 10f		Data Fig. 10g Figs. 11-13					
Ν	100	20		20		50		-	number of neurons
t_0	0	0		$t_{\rm go} - 1$		-0.5		s	simulation start time
t _{go}	-	-		1		2		S	go cue time
t _{max}	2	1		1		3		S	simulation end time
au	0.05	0.05		0.05		0.05		S	membrane time constant
$\mathbf{x}^{(c)}(t_0)$	h ^(c)	0		set^b		0		-	initial condition
$f(\cdot)$	linear ^a	a linear ^a		linear ^a		nonlinear	a	Hz	neural activation function
W	$\stackrel{ ext{iid.}}{\sim} \mathcal{N}ig(0, rac{1}{N}ig)^e$	$\left(\frac{1}{N}\right)^e$ fit ^b		fit ^b		optimized	l ^b	S	weight matrix
С	1	6		6		6		-	number of stimuli
h ^(c)	optimized ^{c3}	fit ^b		-		optimized	C2	-	stimulus input
g	-	-		-		$\sum_{c} \mathbf{h}^{(c)}$		-	go cue
$m_{\rm h}(t)$	0	K(0,0	.25) ^g	0		$\overline{K(0, 0.25)}^{\varepsilon}$	g	-	stimulus masking kernel
$m_{\rm g}(t)$	0	0		0		$K(t_{\rm go}, t_{\rm go})$	$+0.5)^{g}$	-	go cue masking kernel
b	0	fit ^b		fit^{b}		optimized	l ^b	-	cue-independent bias
σ	0.05	0		0		0.05		-	noise standard deviation

Table 1 | Parameters used in the simulations of our models.

^{*a*} For nonlinear networks, $f_i(\mathbf{x}) = [x_i]_+$ was the rectified linear (ReLU) activation function. For linear networks $f_i(\mathbf{x}) = x_i$. The only exception to this was when we created ring attractor networks (Extended Data Fig. 3) in which we used a tanh nonlinearity $f_i(\mathbf{x}) = \tanh(x_i)$. See also text. ^{*b*} See text for details.

^c Inputs were optimized either with both a norm constraint and an overall energy constraint (c_1); only an overall energy constraint (c_2); only a norm constraint (c_3); or so that the dynamics produced the mathematically minimal overall energy (c_4 , see Supplementary Math Note S1). See text for more details.

^{*d*} For the symmetric network, we used $\begin{pmatrix} 0.375 & 0.625 \\ 0.625 & 0.375 \end{pmatrix}$; for the unconstrained network, we used $\begin{pmatrix} 1 & 50 \\ 0 & -11.5 \end{pmatrix}$.

^{*e*} For the symmetric networks, we enforced $\mathbf{W} \leftarrow \frac{1}{2}(\mathbf{W} + \mathbf{W}^{\top})$. For all networks we also shifted the obtained weight matrix by the identity matrix multiplied by a constant so that the largest real part in the eigenvalues of \mathbf{W} is exactly 1 (i.e., the largest eigenvalue of the associated Jacobian would therefore be 0 due to the leak term), and we rejected any \mathbf{W} 's for which the eigenvalue with largest real part had an imaginary component. For Fig. 4, to provide a slightly better agreement between the model dynamics and the experimental recordings, we rejected any \mathbf{W} 's for which the inner product between the most amplifying mode and persistent mode was greater than 0.2 (i.e. we only kept \mathbf{W} 's that were relatively mathematically non-normal).

^{*f*} We used 3 possible input directions (which all had a Euclidean norm of 1): inputs either aligned with the most persistent mode (\mathbf{x}^p) , the most amplifying mode (\mathbf{x}^a) , or a random direction (\mathbf{x}^r) . For the symmetric model, $\mathbf{x}^p = \mathbf{x}^a = [0.707, 0.707]^\top$ and we used $\mathbf{x}^r = [0.98, 0.18]^\top$. For the unconstrained model, $\mathbf{x}^p = [1, 0]^\top$, $\mathbf{x}^a = [0.25, 0.97]^\top$ and we used $\mathbf{x}^r = [0.9, 0.45]^\top$.

 $g K(t_1, t_2) = \begin{cases} 1 & \text{if } t_1 \leq t \leq t_2 \text{ s}, \\ 0 & \text{otherwise.} \end{cases}$ In the table, t_{go} refers to the timing of the go cue (see text).

^{*h*} For training, we used C = 36 cue conditions. For our subsequent analyses (Extended Data Fig. 3), we used C = 6 cue conditions to be consistent with the other models.

and 0.25 s and the initial conditions were either set to zero (Fig. 4, Fig. 5e-f) or sampled randomly around zero 1477 (Fig. 6 and Extended Data Figs. 11–13). For task-optimized networks (Fig. 6 and Extended Data Figs. 11–13), the go cue masking kernel $m_g(t)$ was a boxcar starting at go cue onset, t_{go} (which could be fixed, Extended Data 1479 Fig. 12, or variable, Fig. 6 and Extended Data Figs. 11–13) and lasting for 0.5 s, for all other networks it was set to 1480 0 everywhere. The networks used to analyse the dynamics of information loading (Fig. 3, Extended Data Fig. 4, 1481 and Extended Data Fig. 5a,b) were deterministic by setting $\sigma = 0$, all other networks used noisy dynamics (see 1482 Table 1). All models used a time constant of τ = 50 ms. We solved the dynamics of Eqs. 1 and 2 using a first-order 1483 Euler–Maruyama approximation between t_0 and the simulation end time, t_{max} , with a discretization time step of 1484 1 ms. 1485

For analysis methods that used cross-validation (see below), for each cue condition, we simulated network dy-1486 namics twice with independent realizations of $\eta^{(c)}(t)$, to serve as (trial-averaged) train and test data. For other 1487 analyses, we used a single set of simulated trajectories. All analyses involving networks with randomly generated (or initialized) connectivities that also did not require re-fitting their responses with other networks (Fig. 2, Fig. 3d,e, Fig. 4, Fig. 6a-d, Extended Data Figs. 2 and 3, Extended Data Fig. 7a-c, Extended Data Fig. 5, and 1490 Extended Data Figs. 11–13) were repeated a total of n = 100 times, consisting of either 10 different networks and 10 different simulations (non-cross-validated) or simulation-pairs (cross-validated), each time with independent samples of $\eta^{(c)}(t)$ (for networks with stochastic dynamics—with one exception, see below; Fig. 2, Fig. 4, Fig. 6a–d, Extended Data Figs. 2 and 3, Extended Data Fig. 7a–c, and Extended Data Figs. 11–13), or (for deterministic networks, Fig. 3d,e and Extended Data Fig. 5a,b, as well as for the stochastic networks of Extended Data Fig. 5c,d) 100 different networks, each with a single simulation (non-cross-validated) or simulation-pair (crossvalidated). For those analyses that did require the re-fitting of nonlinear networks' responses with other (linear deterministic) networks (Extended Data Fig. 7d and Fig. 6e), we used just one simulation of the original (stochas-1498 tic nonlinear) network, so n = 10 simulations in total. Analyses of linear deterministic networks (Fig. 3, Fig. 5f, Fig. 6e, Extended Data Fig. 6, Extended Data Fig. 7b,d, Extended Data Fig. 4, Extended Data Fig. 5a,b, and 1500 Extended Data Fig. 10f,g), used a single simulation per network as their dynamics were deterministic. Note that 1501 for (linear deterministic) networks obtained by fitting simulated neural responses, this meant that these analyses 1502 were repeated a total of n = 10 times, once for each set of original responses that had been fit (Fig. 6e, Extended 1503 Data Fig. 7d, and Extended Data Fig. 10f,g). For (linear deterministic) networks obtained by fitting experimentally 1504 recorded neural responses, this meant that these analyses were performed either only once on the single split of 1505 the data (Fig. 5d; blue, Fig. 5e, f, and Extended Data Fig. 10g), or were repeated n = 100 times on 100 different 1506 random time shuffles of the data (Fig. 5d; dark gray, and Extended Data Fig. 10f).

1508 1.3 Nonlinear networks

For the dynamical equations of nonlinear networks, see Methods 1.2. For nonlinear networks (Figs. 2 and 6, Extended Data Figs. 2 and 3, Extended Data Fig. 7c, and Extended Data Figs. 11–13), we ensured that they performed working memory maintenance competently by optimizing their free parameters, **W**, **b**, and $\mathbf{h}^{(c)}$ for appropriate cost functions (see below, Methods 1.3.3).

1.513 1.3.1 Nonlinear networks with instantaneous inputs

Following classical theoretical approaches to attractor network dynamics, we first used nonlinear neural networks in which stimulus inputs acted instantaneously to determine the initial conditions of the dynamics Fig. 2 and Extended Data Figs. 2, 3 and 7. These networks were optimized using a 'just-in-time' cost function (Methods 1.3.3) under one or two constraints. First, for all these networks, we constrained stimulus inputs to have a Euclidean norm of 3 so that we could compare information loading strategies fairly when inputs were constrained to lie in certain subspaces (see also below): either the persistent subspace, persistent nullspace, locally persistent subspace, locally most amplifying subspace, or a random subspace (Fig. 2f,I, Extended Data Fig. 2, and Extended Data Fig. 7). When initial conditions were optimized without a subspace constraint (Fig. 2a–e,g–k, Fig. 2f,I; black lines, and Extended Data Fig. 2b,f), we obtained similar results without this norm constraint on the initial conditions¹. Importantly, we show that qualitatively similar results can be obtained without this norm constraint, but with only a more general energy-based penalty^{13,39,40,66} (Fig. 6 and Extended Data Figs. 3 and 11–13, see also Methods 1.3.3). Second, for symmetric networks, we enforced W $\leftarrow \frac{1}{2}(W + W^{T})$.

¹⁵²⁶ These networks were trained in two epochs. For the first 1000 training iterations, we optimized all free parameters. ¹⁵²⁷ After this, we confirmed that our trained networks did indeed have attractors (i.e. that they were attractor networks)

¹This may sound surprising, as one might expect that, without the norm constraint, networks should just use the trivial solution of using inputs that initialize the network directly in the attractor state. However, the performance of this trivial solution is essentially indistinguishable from that of a network whose dynamics only approach the attractor state later in the delay period when performance actually matters. (Indeed, all working memory tasks we considered had a delay period and the information only needed to be accessible at the end of the delay period and not directly after stimulus offset.) This explains why we never found a trivial solution during optimization even without the norm constraint.

and determined where these attractors were in state space by finding the stable fixed points of the networks' dynamics (Fig. 2d,j, Extended Data Fig. 2b,f, and Extended Data Fig. 3a)—see below. We then continued for another 1000 training iterations with only optimizing the initial conditions, $\mathbf{h}^{(c)}$, while keeping the other parameters, \mathbf{W} (Fig. 2a,g) and **b**, fixed at the values obtained at the end of the first 1000 iterations. For this, we considered three possible scenarios for introducing additional constraints on the initial conditions (beside the one on their norm, see above): they were either projected and then restricted to the persistent subspace or to the persistent nullspace (see Methods 1.7.1 for how these subspaces were computed), or there was no such constraint applied so that they could utilize any direction in the full state space of the network. In addition, to understand the link between the linearized (Methods 1.4.2) and original (above) forms of the dynamics of these networks, we also considered three more constraints on the initial conditions: constraining them to the most persistent, most amplifying, or a random subspace of the linearized dynamics (Extended Data Fig. 7a–c).

1.3.2 Nonlinear networks with temporally extended inputs

To more closely follow the experimental paradigms which we modelled, we also used nonlinear networks in which stimuli provided temporally extended inputs (Fig. 6 and Extended Data Figs. 11–13). To construct these networks, stimulus inputs and the weight matrix were freely optimized (Methods 1.3.3), without any constraints, and optimization proceeded for a full 2000 iterations, without dividing training into different epochs.

1.544 1.3.3 Cost functions and training for nonlinear networks

We trained networks using one of four cost functions: a 'cue-delay' cost, a 'full-delay', a 'just-in-time' cost, and an 'after-go' cost. These costs only differed in terms of the time period in which we applied the cost function. The general form of the cost function we used was a cross entropy loss plus a regularisation term:

$$\mathcal{L} = \left\langle -\alpha_{\text{nonlin}}^{(1)} \sum_{c=1}^{6} \left(\mathbf{y}^{(c)} \right)^{\top} \int_{T_1}^{T_2} \log \left(\text{Softmax} \left(\mathbf{W}_{\text{out}} \, \mathbf{r}^{(c)}(t) + \mathbf{b}_{\text{out}} \right) \right) \, \mathrm{d}t + \alpha_{\text{nonlin}}^{(2)} \sum_{c=1}^{6} \int_{t_0}^{t_{\text{max}}} \left\| \mathbf{r}^{(c)}(t) \right\|_2^2 \, \mathrm{d}t \right\rangle \tag{3}$$

where T_1 and T_2 determine the time period in which we applied the cost, $\alpha_{\text{nonlin}}^{(1)}$ and $\alpha_{\text{nonlin}}^{(2)}$ control the relative contributions of the cross-entropy loss and firing rate regularisation, $\mathbf{W}_{\text{out}} \in \mathcal{R}^{6 \times N}$ and $\mathbf{b}_{\text{out}} \in \mathcal{R}^6$ include the 6

sets of 'readout' weights and biases, respectively, and $\mathbf{y}^{(c)} \in \mathcal{R}^6$ is a one-hot vector where $y_i^{(c)} = \begin{cases} 1 & \text{if } i = c \\ 0 & \text{otherwise} \end{cases}$

defining the 'target' output for each cue condition. We initialized elements of the network parameters \mathbf{W} , \mathbf{b} , $\mathbf{h}^{(c)}$, as well as the readout parameters \mathbf{W}_{out} and \mathbf{b}_{out} from a Gaussian distribution with mean 0 and variance 1/N, and then optimized using gradient descent with Adam optimization⁸³, where gradients were obtained from backpropagation through time. The angle brackets, $\langle \cdot \rangle$, denote averaging over batch sizes of 50 random realisations of $\mathbf{r}^{(c)}$. We used a learning rate of 0.0005.

See Table 2 for how we set the parameters of Eq. 3 (T_1 , T_2 , t_0 , $\alpha_{nonlin}^{(1)}$ and $\alpha_{nonlin}^{(2)}$) depending on the cost function and the level of regularization. Briefly, the cue-delay cost included both the cue (between stimulus cue onset and offset) and the delay period (between stimulus cue offset and go cue onset), the full-delay cost the included delay period but not the cue period, the just-in-time cost started between stimulus onset and the earliest go time and ended at the onset of the go cue, and the after-go cost started at go cue onset and lasted for the duration of the go cue (0.5 s). For simulating the random delay task (Fig. 6 and Extended Data Figs. 11-13), analogous to what animals need to solve (see below), we sampled the go time uniformly between $t_{go} = 0.75$ s and $t_{go} = 2$ s. For just-in-time (Fig. 2 and Extended Data Fig. 2) and after-go trained networks (Extended Data Fig. 12), we also used a fixed delay task with a simulation end time of $t_{max} = 2$ s or a go time of $t_{go} = 2$ s, respectively. For the other 1561 cost functions, networks trained on the fixed delay task yielded very similar dynamics to their counterparts trained 1562 on the variable delay task (not shown). We set $\alpha_{\text{nonlin}}^{(1)}$ and $\alpha_{\text{nonlin}}^{(2)}$ so that networks could reliably learn the task (at performance levels comparable across different settings) while also exhibiting relatively stable dynamics (i.e. if $\alpha_{\text{nonlin}}^{(1)}/\alpha_{\text{nonlin}}^{(2)}$ is too large, the network dynamics can explode whereas if $\alpha_{\text{nonlin}}^{(1)}/\alpha_{\text{nonlin}}^{(2)}$ is too small, the network is not able to learn the task). Note that vanishing gradients during training impacted the value of $\alpha_{\text{nonlin}}^{(1)}$ that 1565 was required for different networks to exhibit similar performance (Fig. 6c). Nevertheless, $\alpha_{nonlin}^{(2)}$ was varied by 1567 an order of magnitude between Fig. 6 and Extended Data Figs. 11-13 to specifically test the robustness of our 1568 results to this parameter.

1.3.4 Optimized ring/bump attractor networks

When training to create ring/bump attractor networks (Extended Data Fig. 3), we made three modifications to the nonlinear networks described above. First, in line with other approaches for optimizing recurrent neural

Cost function	<i>T</i> ₁ (s)	<i>T</i> ₂ (s)	$\alpha_{\rm nonlin}^{(1)}$ (1/s)	$\alpha_{\text{nonlin}}^{(2)}(1)$	Figures
cue-delay	0	t _{go}	20	0.00005	Fig. 6 and Extended Data Fig. 11
			20	0.0005	Extended Data Fig. 11
full-delay	0.25	t _{go}	20	0.00005	Extended Data Fig. 13
			20	0.0005	Extended Data Fig. 13
just-in-time	0.5	$t_{\rm max}$	33	0.05	Extended Data Fig. 3
	0.5	$t_{\rm max}$	33	0.00005	Fig. 2 and Extended Data Fig. 2
	0.75	t _{go}	33	0.00005	Fig. 6 and Extended Data Fig. 11 and Extended Data
					Fig. 1e
				0.0005	Extended Data Fig. 11
after-go	t _{go}	$t_{\rm go} + 0.5$	10	0.00005	Fig. 6
				0.0005	Extended Data Fig. 12

Table 2 | **Parameters for nonlinear network optimization.** Times T_1 and T_2 are relative to stimulus onset at t = 0. Units are shown in parentheses after the name of the corresponding parameter.

networks ^{17,20,24,66}, we used a hyperbolic tangent nonlinearity because the saturation of this nonlinearity greatly encouraged a continuous attractor to form compared with a ReLu nonlinearity. Second, we trained networks with 36 cue conditions (without enforcing any particular metric relationship between the corresponding inputs, $\mathbf{h}^{(c)}$, as above), and then subsequently restricted our analyses to 6 evenly spaced cue conditions to keep consistency with our other analyses. Third, we used a cost function that measured estimation (or fine, rather than coarse, discrimination) performance across those 36 conditions, thus encouraging a ring attractor to form:

$$\mathcal{L} = \left\langle \alpha_{\text{nonlin}}^{(1)} \sum_{c=1}^{36} \int_{T_1}^{T_2} \left[1 - \cos\left(\hat{\theta} - \theta^{(c)}\right) \right] dt + \alpha_{\text{nonlin}}^{(2)} \sum_{c=1}^{36} \int_{t_0}^{t_{\text{max}}} \left\| \mathbf{r}^{(c)}(t) \right\|_2^2 dt \right\rangle$$
(4)

with
$$\hat{\theta} = \operatorname{atan2}(\mathbf{W}_{\operatorname{out}}^{\operatorname{y}} \mathbf{r}^{(c)}(t), \mathbf{W}_{\operatorname{out}}^{\operatorname{x}} \mathbf{r}^{(c)}(t))$$
 (5)

where $\hat{\theta}$ is the population vector-decoded stimulus angle, such that atan2(y, x) gives the angle that the point [x, y]makes with the x-axis, \mathbf{W}_{out}^x , $\mathbf{W}_{out}^y \in \mathcal{R}^{1 \times N}$ are 2 sets of 'readout' weights defining the plane in which decoded angles are defined, and $\theta^{(c)} = 2\pi \frac{c}{36}$ is the target angle for cue condition *c*. All other terms were the same as those defined in Methods 1.3.3.

We note that the cosine term in the cost function (Eq. 4), quantifying the precision of the decoded angle, is closely related to a cost measuring the population Fisher information about angle. To see this, recall that the Fisher information in this case is

$$\mathcal{I} = -\int \mathcal{P}(\mathbf{r}|\theta^{(c)}) \frac{\partial^2}{\partial\theta^2} \ln \mathcal{P}(\mathbf{r}|\theta) \,\mathrm{d}\mathbf{r}$$
(6)

In the limit of a sufficiently large population, the maximum likelihood estimator, $\hat{\theta}_{ML}$ achieves the same Fisher information as the full population vector, and is distributed as a (circular) Gaussian (von Mises) distribution centered on the true orientation, $\theta^{(c)}$, with some constant (circular) concentration, κ , so we can write

$$\mathcal{I} \simeq -\int \mathcal{P}\left(\hat{\theta}_{\mathsf{ML}}|\theta^{(c)}\right) \,\frac{\partial^2}{\partial\theta^2} \,\ln \mathcal{P}\left(\hat{\theta}_{\mathsf{ML}}|\theta\right) \mathsf{d}\hat{\theta}_{\mathsf{ML}} \tag{7}$$

$$= \kappa \int \mathcal{P}(\hat{\theta}_{\mathsf{ML}}|\theta^{(c)}) \cos\left(\hat{\theta}_{\mathsf{ML}} - \theta^{(c)}\right) d\hat{\theta}_{\mathsf{ML}}$$
(8)

Assuming that our population vector-based decoder is an efficient estimator that approximates the maximum likelihood decoder, $\hat{\theta} \simeq \hat{\theta}_{ML}$, and substituting the integral over the distribution of the estimate with an empirical average over its stochastic realizations, we can further rewrite the Fisher information as

$$\mathcal{I} \simeq \kappa \left\langle \cos\left(\hat{\theta}_{\mathsf{ML}} - \theta^{(c)}\right) \right\rangle \tag{9}$$

¹⁵⁷⁵ (We also checked empirically that, in line with our assumptions above, the empirical distribution of $\hat{\theta}$ was well ¹⁵⁷⁶ approximated by a von Mises distribution centered on |*theta*^(c) with a constant concentration across target angles.) ¹⁵⁷⁷ Eq. 9 is thus identical to the first term in Eq. 4 up to an additive constant (the 1 inside the square bracket in Eq. 4), ¹⁵⁷⁸ which does not matter for optimization, a multiplicative constant (which can be incorporated into $\alpha_{nonlin}^{(1)}$) and a ¹⁵⁷⁹ sign-flip, because we are maximizing Fisher information in Eq. 9 but minimizing the cost in Eq. 4. Therefore, ¹⁵⁸⁰ minimizing (the first term in) Eq. 4 also (approximately) maximizes Eq. 6, and vice versa.

1581 1.4 Linear networks

For the dynamical equations of linear networks, see Methods 1.2. Linear networks were either constructed '*de novo*' (Figs. 3 and 4 and Extended Data Figs. 4 and 5), obtained by a local linearization of canonical nonlinear dynamical systems (Extended Data Fig. 6) or of nonlinear neural network dynamics (Extended Data Fig. 3e and Extended Data Fig. 7a–c), or they were fitted to neural responses obtained from experiments (Fig. 5f and Extended Data Fig. 10f,g) or the simulation of nonlinear networks (Fig. 6e and Extended Data Fig. 7d).

1.587 1.4.1 De novo linear networks

We used *de novo* linear networks to develop an analytical understanding of the dynamics of optimal information loading. These networks included small 2-neuron networks with hand-picked parameters (see Table 1) chosen to illustrate the differences between normal (symmetric) and non-normal (unconstrained) dynamics and the effects of different initial conditions (Fig. 3a–c), as well as large networks (with 10, 100, or 1000 neurons) with randomly generated parameters (Fig. 3d,e, Fig. 4, Extended Data Fig. 4, and Extended Data Fig. 5a,b; see Table 1). We always set the largest eigenvalue of the weight matrix to be exactly 1 (i.e., the largest eigenvalue of the associated Jacobian would therefore be 0 due to the leak term) so that these networks had an integrating or 'persistent' mode^{6,32,34,43} (see Table 1)

Initial conditions (Fig. 3, Extended Data Fig. 4, and Extended Data Fig. 5a,b) or temporally extended inputs 1596 (Fig. 4) were determined by computing the most persistent and amplifying direction(s) based on the Jacobian of the dynamics (Figs. 3 and 4 and Extended Data Fig. 5a,b, see Methods 1.7.1; for how initial conditions were 1598 determined in Extended Data Fig. 4c,d see Supplementary Math Note S1.8). For the networks in Fig. 4, we 1599 also added a small amount of noise to the input to allow for some transient dynamics for all input directions (see Fig. 4c at 0 s). Alternatively, we optimized initial conditions for maximal asymptotic overlap with the most persistent mode (Extended Data Fig. 5c,d; see below). For setting the noise level, σ , in these networks, we 1602 considered two scenarios: noise matched (Fig. 4a, light green and gray) and performance matched (Fig. 4a, dark 1603 green and black). For noise matched simulations, we first determined the highest value of σ that still allowed us to 1604 obtain 100% decodability (using a delay-trained decoder) for all networks when receiving inputs aligned with the most amplifying mode (Fig. 4a, red). This resulted in $\sigma = 0.1$ for symmetric models, and $\sigma = 0.17$ for unconstrained models. We then used the same σ for simulations using inputs aligned with the most persistent and random 1607 directions. For performance matched simulations, we used a different value of σ for each possible input direction so that all models achieved 100% decodability using a delay-trained decoder. For symmetric models, this resulted in σ = 0.1 for inputs aligned with either the persistent or most amplifying modes, and σ = 0.005 for random inputs. 1610 For unconstrained models, this resulted in σ = 0.17 for inputs aligned with the most amplifying mode, σ = 0.02 for 1611 inputs aligned with the persistent mode, and $\sigma = 0.005$ for random inputs. (Note that, consistent with our theory, 1612 smaller noise levels were necessary to achieve the same desired level of performance for input directions that were predicted to be increasingly suboptimal by our analysis.) 1614

To demonstrate that the initial conditions along the most amplifying directions, obtained by control theoretic analyses, were indeed optimal for maximising the overlap with the most persistent mode (the measure of optimality we used for these networks, Fig. 3c,e), we also used a direct numerical optimization approach, analogous to that used to optimize initial conditions in our nonlinear networks (Figs. 2 and 6, see also Methods 1.3.3). Specifically, we optimized $\mathbf{h}^{(c)}$ (constrained to have unit Euclidean norm) with gradient descent using Adam optimization⁸³ with gradients obtained from back-propagation through time using the following cost function

$$\mathcal{L} = \int_{1.5 \text{ s}}^{2 \text{ s}} \left[\tanh\left(\mathbf{v}_{1}^{\mathsf{T}} \mathbf{x}(t)\right) - 1 \right]^{2} \mathrm{d}t$$
(10)

where v₁ is the eigenvector associated with eigenvalue 0 of the Jacobian (i.e. the most persistent mode). We used a learning rate of 0.0001. We performed the above training procedure independently for 100 random noisy networks (either symmetric or unconstrained) and we show averaged results in Extended Data Fig. 5c,d. We also used random initial conditions as controls. These had elements that were either sampled from a standard normal distribution (re-scaled to have unit Euclidean norm) in large networks (Fig. 3d,e, Fig. 4, and Extended Data Fig. 5a,b), or in the case of 2-neuron networks, quasi-randomly chosen (with unit Euclidean norm) for illustrative purposes (Fig. 3a–c).

1622 1.4.2 Local linearization of nonlinear dynamics

To better understand how the dynamics of optimal information loading that we identified in linear networks apply to nonlinear attractor dynamics, we performed a local linearization of our simulated nonlinear networks (Extended Data Fig. 6, Extended Data Fig. 3, and Extended Data Fig. 7a–c). This approach required access to the 'true' dynamical equations of the nonlinear networks—which we had by construction.

We performed local linearizations of the original nonlinear network dynamics in x-space (the space of variables 1627 which the dynamics was defined, Eq. 1) around the origin (we found empirically that initial conditions were in 1628 distributed close to the origin)-which served as the reference point with respect to which the norm of optimized 1629 initial conditions was constrained in the networks we linearized (Methods 1.3; analogous to our analysis of information loading in linear networks, Fig. 3a-e, and see also Methods 1.7.2). As the ReLU firing rate nonlinearity 1631 of these networks is non-differentiable at exactly the origin, we computed the 'average' Jacobian of the system 1632 the immediate vicinity of the origin instead (this allowed us to use the same linearization and the same set in 1633 of amplifying modes for all initial conditions; we obtained highly similar results by linearizing separately for each initial condition). Because the derivative of each ReLU is 0 or 1 in half of the activity space of the network, this 1635 resulted in the Jacobian I = $\frac{1}{2}$ W^{*} - I, where W^{*} is the weight matrix of the original nonlinear network. Note that one obtains the same result even without averaging, by regarding the ReLu nonlinearity as the limiting case of the soft-ReLu nonlinearity: $[x]_{+} = \lim_{\beta \to \infty} \frac{1}{\beta} \ln(1 + e^{\beta x})$, of which the derivative at x = 0 is $\frac{1}{2}$ (at any value of the inverse 1638 temperature, β) and thus results in the same Jacobian as above. We confirmed that the resulting dynamics were 1639

always stable (largest real eigenvalue of J was less than 0). We then used this system to identify the locally (around the origin) most amplifying or most persistent modes (Extended Data Fig. 7a).

For simulating these linearized networks (Extended Data Fig. 7b), we then used the Jacobian we thus obtained 1642 to map the resulting linearized dynamics to a deterministic integrator with the effective weight matrix $\mathbf{W} = (\mathbf{J} + \mathbf{I})$ 1643 $\lambda_{
m max}$ I, where $\lambda_{
m max}$ is the largest real eigenvalue of J. Thus, the resulting dynamics were always marginally stable 1644 (largest real eigenvalue of J was exactly 0). (Note that for subsequent analyses involving most persistent and 1645 amplifying modes, we used the original weight matrix, see more in Methods 1.7.1. Nevertheless, the most 1646 persistent modes of the weight matrices we used for simulation and those we used in subsequent analyses were 1647 identical, as they only relied on the eigenvectors of the weight matrix, or the Jacobian, and the rank order of their associated eigenvalues, which this stabilization did not affect. We also checked numerically that making the 1649 system marginally stable only had very minor effects on the most amplifying modes, with correlations between 1650 the most amplifying modes of the original and simulated dynamics being above 0.9. Thus, in these respects, our 1651 simulations were representative of the dynamics of the original systems.) The bias parameters, b, were the same 1652 as in the original nonlinear networks. The initial conditions, $\mathbf{h}^{(c)}$, were either the ones we originally optimized for 1653 the nonlinear dynamics without any constraints (beside a constraint on their norm), or they were optimized while constraining them to the most persistent, most amplifying, or a randomly chosen subspace of these linearized dynamics (all were of the same dimensionality for a fair comparison, Extended Data Fig. 7). 1656

For ring attractor networks (Extended Data Fig. 3), which used a tanh nonlinearity (Methods 1.3.4), the associated linearized system around the origin is given by the Jacobian J = W - I, which we then used to identify the locally most amplifying and persistent modes (Extended Data Fig. 3e).

Ve used the same approach to linearize the dynamics of the canonical minimal nonlinear attractor dynamics that we used to gain insights into information loading in nonlinear systems (Methods 1.6, see also Supplementary 1661 Math Note S2 and Extended Data Fig. 6). In this case, the Jacobian was well defined at the origin, so there was no need to average it. For consistency with the notation and terminology we use in the rest of this paper, 1663 and without loss of generality (as linear dynamical systems and linear neural networks are isomorphic), we refer 1664 to the resulting linear dynamical system as a 'linear neural network' and define it by its 'effective' weight matrix 1665 (defined via the Jacobian as above). Initial conditions were magnitude-matched and chosen to align with the most persistent or the most amplifying direction extracted from the Jacobian (Methods 1.7.1), or chosen randomly, or 1667 varied systematically to cover the whole range of possible directions. There were no other parameters for these 1668 linearized 'networks'. 1669

1.4.3 Fitting linear neural networks to neural responses

¹⁶⁷¹ In order to be able to apply our theoretically derived measures of optimal information loading without having access to the true dynamics of the system, we also created linear neural networks whose parameters were fitted to experimental data (see below). As a control, we repeated the same fitting procedure with simulated nonlinear networks to validate that our approach provides meaningful results when 1. we do not have access to the true dynamics but only to samples of activities generated by those dynamics, and 2. we also cannot assume that the true dynamics are linear.

¹⁶⁷⁷ We fitted deterministic linear neural networks to 1 s of trial-averaged neural activity (experimentally recorded, or ¹⁶⁷⁸ simulated by a nonlinear neural network model). For the main analyses (Fig. 5d–f, Fig. 6e, Extended Data Fig. 3f, ¹⁶⁷⁹ and Extended Data Fig. 7d), we used data starting from the onset of the stimulus cue. For the control analysis of ¹⁶⁸⁰ late delay experimental recordings (Extended Data Fig. 10g), we used the final 1 s of neural activity just prior to ¹⁶⁸¹ the go cue. For the shuffle control (Fig. 5d; dark gray, and Extended Data Fig. 10f), we again used data starting

from stimulus onset but randomly shuffled neural activity across time and proceeded by fitting this shuffled data 1683 instead.

For fitting high dimensional neural data, we first performed principal components analysis on neural activity (dimensions: neurons, data points: time points, indexed by t, and cue conditions, indexed by c), and projected it through the principal components (PCs): $\mathbf{x}_{*}^{(c)}(t) = \mathbf{P} \mathbf{r}_{*}^{(c)}(t)$, where the columns of **P** are top 20 principal components of the data, and $\mathbf{r}_{*}^{(c)}(t)$ is trial averaged neural responses (mean-centered, see above) at time t in condition c. These top 20 PCs captured approximately 75% and 76% of variance for monkeys K and T, respectively during the cue and early delay period (Fig. 5d-f), 70% and 60% of variance for monkeys K and T, respectively during the late delay period (Extended Data Fig. 10g), and over 95% of the variance for all simulated neural activities (Fig. 6e). The projected neural activity time courses of the neural data, $\mathbf{x}_{*}^{(c)}(t)$, served as the targets that needed to be matched (after a suitable linear transformation with 'read-out' matrix $\mathbf{C} \in \mathcal{R}^{20 \times 20}$) by the neural activity time courses generated by the fitted neural network's dynamics in the corresponding cue conditions, $\mathbf{x}^{(c)}(t)$ (Eqs. 1 and 2). For fitting the parameters of the network (\mathbf{W} , $\mathbf{h}^{(c)}$, \mathbf{b}) and the readout matrix (\mathbf{C}), we used the following cost function:

$$\mathcal{L} = \varepsilon^2 + \alpha_{\text{lin}} \left[\left\| \mathbf{C} \right\|_{\text{F}}^2 + \left\| \mathbf{b} \right\|_2^2 + \sum_{c=1}^6 \left\| \mathbf{h}^{(c)} \right\|_2^2 \right]$$
(11)

(13)

with

and

$$\varepsilon^{2} = \frac{1}{6} \sum_{c=1}^{5} \int_{0}^{1.5} \left(\mathbf{e}^{(c)}(t) \right)^{\top} \mathbf{D} \, \mathbf{e}^{(c)}(t) \, \mathrm{d}t \qquad \text{being the mean squared error of the fit} \qquad (12)$$
$$\mathbf{e}^{(c)}(t) = \mathbf{C} \, \mathbf{x}^{(c)}(t) - \mathbf{x}^{(c)}(t) \qquad \text{the momentary error} \qquad (13)$$

where D is a diagonal matrix with the variances explained by the corresponding PCs in P on the diagonal (en-1684 couraging the optimization procedure to prioritize fitting the top PCs), $\|\cdot\|_{F}^{2}$ is the Frobenius norm of a matrix. 1685

Although including C as an additional parameter (free to be optimized) makes the fitting problem overparametrized 1686 (at least with respect to the fitting error, ε^2 , as for any invertible choice of G, any remapping of the parameters 1687 as $W \rightarrow G W G^{-1}$, $h^{(c)} \rightarrow G h^{(c)}$, $b \rightarrow G b$, and $C \rightarrow C G^{-1}$ achieves the same ε^2), we included C because it allowed the network to develop dynamics that make appropriate use of persistent and amplifying modes without 1689 simultaneously having to match the neural dimensions (which can be easily re-mapped with C). We then used C as the read-out matrix when identifying amplifying modes (see Methods 1.7.1). Indeed, when validating this fitting procedure with simulated responses generated by linear stochastic neural networks with different (instantaneous) 1692 information loading strategies (most persistent, most amplifying, random; see Methods 1.7.2), and using our 1693 standard subspace-overlap-based measures (Methods 1.7.3) to identify the information loading strategy from these simulated responses, we found that the true information loading strategy was recovered more reliably with including C than without it (not shown). Moreover, we found this approach was even able to distinguish between different information loading strategies of nonlinear networks from simulated data (Extended Data Fig. 7d). 1697

Also note that we had no constraints on W to define stable dynamics. Nevertheless, when fitting experimental recordings, and responses generated by nonlinear attractor networks, we found that the largest real eigenvalue 1699 of the fitted W was typically within the 0.95 $\leq \lambda_{max} \leq$ 1.05 range, i.e. the dynamics were near marginal stability, in 1700 line with the dynamics of our *de novo* linear neural networks (Methods 1.4.1), as well as of those that we obtained 1701 by local linearization (Methods 1.4.2). The only exception was when fitting the responses of nonlinear networks 1702 ained on an after-go-time cost (Methods 1.3.3) which resulted in dynamics without attractors and, consequently, the fitted linear dynamics had $\lambda_{max} > 1.05$.

We used Adam⁸³ to perform gradient descent optimization of W, $h^{(c)}$, b, and C with gradients obtained from 1705 back-propagation through time, and a learning rate of 0.0001. We initialized elements of all of these parameters from a Gaussian distribution with mean 0 and variance 1/20. We set the regularisation parameter to α_{lin} = 1/12, 1707 although we found that the results did not change substantially when setting $\alpha_{\text{lin}} = 0$ or using larger values of α_{lin} .

The stimulus-masking kernel ($m_h(t)$, Table 1) was matched to how the responses being fitted were obtained: with temporally extended or instantaneous inputs. Specifically, when fitting responses to temporally extended inputs (experimentally measured, Fig. 5f and Extended Data Fig. 10f, or simulated, Fig. 6e), the masking kernel of the fitted linear network matched the cue period. When fitting responses generated by networks driven by instantaneous inputs (Extended Data Fig. 3f and Extended Data Fig. 7d), or when fitting the late delay period of experimental recordings (during which no stimulus is present, Extended Data Fig. 10g), the stimulus masking ker-1714 nel was set to zero, and instead the initial condition of the fitted linear network was tuned to match the responses (see below).

In most cases (Fig. 5f, Fig. 6e, and Extended Data Fig. 10f), we set the initial condition $\mathbf{x}^{(c)}(t_0) = \mathbf{0}$. There were two exceptions to this. First, when fitting the late delay dynamics in the experimental recordings (Extended Data Fig. 10g), we set $\mathbf{x}^{(c)}(t_0) = \mathbf{C}^{-1} \mathbf{x}^{(c)}_{*}(t_0)$ (i.e. we fixed the initial condition of the latent dynamics to the data; we also observed qualitatively similar results when we included $\mathbf{x}^{(c)}(t_0)$ as a separate optimizable parameter in this case).

Second, when fitting simulated data from models that used instantaneous stimulus inputs (Extended Data Fig. 3f and Extended Data Fig. 7d), we set $\mathbf{x}^{(c)}(t_0) = \mathbf{h}^{(c)}$.

1723 1.5 Previous working memory models

We used the following dynamics for implementing all previous neural network models of working memory:

$$\mathbf{x}^{(c)} = \mathbf{W} \mathbf{r}^{(c)}(t) + m_{\rm h}(t) \mathbf{h}^{(c)} + m_{\rm g}(t) \mathbf{g}$$
(14)

$$\tau_{\rm r} \frac{\mathrm{d}\mathbf{r}^{(c)}(t)}{\mathrm{d}t} = -\mathbf{r}^{(c)}(t) + \mathbf{f}\left(\mathbf{x}^{(c)}(t)\right) + \mathbf{b}_{\rm r} + \boldsymbol{\sigma}_{\rm r} \,\boldsymbol{\eta}_{\rm r}^{(c)}(t) \tag{15}$$

where all symbols refer to the same (or a closely analogous, see below) quantity as in Eqs. 1 and 2. Note that we use this notation to best expose the similarities with and differences from the dynamics of our networks (Eqs. 1 and 2), rather than the original notation used for describing these models ^{5,6,26}, but the dynamics are nevertheless 1726 identical to those previously published. Overall, these dynamics are closely analogous to those that we used earlier for our networks with the following differences. First, for us, dynamics were defined in x-space, with r being an instantaneous function of x. Here, the dynamics are defined instead in r-space (Extended Data Fig. 1ad and Extended Data Fig. 8), with x being an instantaneous function of r. (There are slightly different assumptions 1730 underlying these rate-based formulations of neural network dynamics when deriving them as approximations of the dynamics of spiking neural networks⁴⁸, and the two become identical in the case of linear dynamics.) As a result, time constants, τ_r , biases, \mathbf{b}_r , and the variance of noise, σ_r (as well as noise itself, $\eta_r^{(c)}(t)$), are defined for r rather than x. For nonlinear variants of these networks, there are also differences for the choice of single neuron nonlinearities, f(.). Furthermore, some of these networks distinguish between excitatory and inhibitory cells, with different time constants, and noise standard deviations. Thus, each of these parameters is represented as a diagonal matrix, $\tau_{\rm r}$ and $\sigma_{\rm r}$, respectively, with each element on the diagonal storing one of two possible values of that parameter depending on the type (excitatory or inhibitory) of the corresponding neuron ($\tau_r^{\rm E}$ and $\sigma_r^{\rm E}$, or $\tau_r^{\rm I}$ and 1738 $\sigma_{\rm r}^{\rm I}$, respectively). Most importantly, all of these networks used a set of parameters which were hand-crafted to 1739 produce the required type of dynamics, rather than optimized for a function (or to fit data) as in the case of our 1740 networks. In line with our analyses of experimental data and task-optimized networks (Figs. 5 and 6), simulations started at $t_0 = -0.5$ s, i.e. 0.5 s before stimulus cue onset (defined as t = 0), the stimulus cue lasted for 0.25 s, and the go cue appeared at t_{go} = 2 s and lasted for 0.5 s. (Note that for these networks we considered the fixed-delay variant of the task as that is what these networks were originally constructed to solve.) As with our networks (Methods 1.2), we solved the dynamics of Eqs. 14 and 15 using a first-order Euler-Maruyama approximation between t_0 and the simulation end time with a discretization time step of 1 ms. 1746

For analysis methods that used cross-validation (see below), we simulated network dynamics twice (for each cue condition) with independent realizations of $\eta_r^{(c)}(t)$, to serve as (trial-averaged) train and test data. For other analyses, we used a single set of simulated trajectories. All analyses involving these networks were repeated n = 10 times, using 10 different simulations (non-cross-validated) or simulation-pairs (cross-validated), each time with independent samples of $\eta_r^{(c)}(t)$.

¹⁷⁵² Table 3 provides the values of most network and other parameters used for simulating each model. In the following ¹⁷⁵³ we provide the additional details for each of these models that are not included in Table 3.

1.5.1 Classical bump attractor model

The bump attractor model that we used (Extended Data Fig. 1a) has been described previously (see Ref. 5). The model contained separate excitatory and inhibitory populations. As in the discrete attractors model, the weight matrix was of the form

$$\mathbf{W} = \begin{pmatrix} \mathbf{W}^{\text{EE}} & -\mathbf{W}^{\text{IE}} \\ \mathbf{W}^{\text{EI}} & -\mathbf{W}^{\text{II}} \end{pmatrix}$$
(16)

where the elements of \mathbf{W}^{IE} , \mathbf{W}^{EI} , and \mathbf{W}^{II} were set to 6.8/N, 8/N, and 1.7/N, respectively. The excitatory sub-matrix \mathbf{W}^{EE} had a circulant form:

$$W_{ij}^{\text{EE}} = \frac{6 e^{1.5 \cos\left(\frac{4 \pi (i-j)}{N}\right)}}{\sum_{k=0}^{N/2-1} e^{1.5 \cos\left(\frac{4 \pi k}{N}\right)}}$$
(17)

1755 for cell-pairs i, j = 1, ..., N/2.

Stimulus cue inputs were also analogous to those used in the discrete attractors models and were set to

$$h_i^{(c)} = \frac{200 \ e^{1.5 \ \cos\left(\pi\left(\frac{4i}{N} - \frac{2c-1}{6}\right)\right)}}{\sum_{k=1}^{N/2} e^{1.5 \ \cos\left(\pi\left(\frac{4k}{N} - \frac{2c-1}{6}\right)\right)}}$$
(18)

¹⁷⁵⁶ for cues c = 1, ..., 6 and cells i = 1, ..., N/2 (i.e., as above, inputs were only delivered to the excitatory neurons).

Parameters used in network simulations of previous models								
Symbol	Extended Data	Extended Data	Extended Data	Extended Data	Units	Description		
	Fig. 1a	Fig. 1b	Fig. 1c and	Fig. 1d and				
			Extended Data	Extended Data				
			Fig. 8a,b,d,e	Fig. 8c,f				
Ν	100	108	100	100	-	number of neurons		
t_0	-0.5	-0.5	-0.5	-0.5	S	simulation start time		
t _{go}	2	2	2	2	S	go cue time		
t _{max}	3	3	3	3	S	simulation end time		
au	-	-	0.05	0.01	S	membrane time constant		
$ au_{r}^{\mathrm{E}}$	0.02	0.02	-	-	S	membrane time constant		
						(E neurons)		
$ au_{ m r}^{ m I}$	0.01	0.01	-	-	S	membrane time constant		
						(I neurons)		
$r^{(c)}(t_0)$	0	0	0	0	Hz	initial condition		
$f(\cdot)$	nonlinear ^a	nonlinear ^a	linear ^a	linear ^a	Hz	neural activation function		
W	set ^b	set ^b	set^b	set ^b	S	weight matrix		
С	6	6	6	6	-	number of stimuli		
$\mathbf{h}^{(c)}$	set ^b	set ^b	set^b	set ^b	-	stimulus input		
g	$\sum_{c} \mathbf{h}^{(c)}$	$\sum_{c} \mathbf{h}^{(c)}$	$\sum_{c} \mathbf{h}^{(c)}$	$\sum_{c} \mathbf{h}^{(c)}$	-	go cue		
$m_{\rm h}(t)$	$\overline{K(0, 0.25)}^{c}$	$\overline{K(0, 0.25)}^c$	$\overline{K(0, 0.25)}^c$	$\overline{K(0, 0.25)}^c$	-	stimulus masking kernel		
$m_{\rm g}(t)$	$K(t_{\rm go}, t_{\rm go} + 0.5)^c$	-	go cue masking kernel					
b _r	$b_{\rm r}^{\rm E}$ = 0.2,	$b_{\rm r}^{\rm E} = -1.2$,	0	0	Hz	cue-independent bias		
	$b_{\rm r}^{\rm I} = 0.5$	$b_{\rm r}^{\rm I} = 0.28$						
$\sigma_{\rm r}$	-	-	0.02	0.02	Hz	noise standard deviation		
$\sigma_{ m r}^{ m E}$	1	2	-	-	Hz	noise standard deviation		
						(E neurons)		
$\sigma_{\rm r}^{\rm I}$	3	1	-	-	Hz	noise standard deviation		
						(I neurons)		

Table 3 | Parameters used in previous models.

^{*a*} For nonlinear networks, $f_i(\mathbf{x}) = \begin{cases} [x_i]_+^2 & \text{if } x_i \leq 1, \\ \sqrt{4x_i - 3} & \text{otherwise.} \end{cases}$. For linear networks $f_i(\mathbf{x}) = x_i$. ^{*b*} See text for details.

$$K(t_1, t_2) = \begin{cases} 1 & \text{if } t_1 \leq t \leq t_2 \\ 0 & \text{otherwise.} \end{cases}$$

1757 1.5.2 Discrete attractors model

¹⁷⁵⁸ The discrete attractors model that we used (Extended Data Fig. 1b) has been described previously (see the ¹⁷⁵⁹ methods of Ref. 5). The model contained separate excitatory and inhibitory populations.

The weight matrix was of the form

$$\mathbf{W} = \begin{pmatrix} \mathbf{W}^{\text{EE}} & -\mathbf{W}^{\text{IE}} \\ \mathbf{W}^{\text{EI}} & -\mathbf{W}^{\text{II}} \end{pmatrix}$$
(19)

where the elements of \mathbf{W}^{IE} , \mathbf{W}^{EI} , and \mathbf{W}^{II} were set to 2.4/*N*, 8/*N*, and 2.6/*N*, respectively. The excitatory sub-matrix \mathbf{W}^{EE} was constructed by dividing the population of excitatory cells into six clusters (of 9 neurons each), with each cluster corresponding to one of the stimulus cue conditions. Connections within each cluster were strong, with a value of 30/N. Connections between neurons belonging to clusters that corresponded to adjacent stimulus cues were weaker, with a value of 2.5/N. All other connections were very weak, with a value of 0.02/N. This resulted in a block circulant structure for \mathbf{W}^{EE} .

Stimulus cue inputs were set to

$$h_{i}^{(c)} \propto \frac{350 \ e^{8} \cos\left(\pi\left(\frac{4 \ i}{N} - \frac{2 \ c - 1}{6}\right)\right)}{\sum_{k=1}^{N/2} e^{8} \cos\left(\pi\left(\frac{4 \ k}{N} - \frac{2 \ c - 1}{6}\right)\right)}$$
(20)

for cues c = 1, ..., 6 and cells i = 1, ..., N/2 (i.e. inputs were only delivered to the excitatory neurons).

1767 1.5.3 Linear integrator model

The linear integrator model that we used (Extended Data Fig. 1c and Extended Data Fig. 8a,d) has been described previously (see Ref. 6). There were no separate excitatory and inhibitory populations in this model, and the weight matrix was constructed such that network dynamics were non-normal, non-oscillatory, and stable with a single two-dimensional neutrally stable subspace (i.e. a plane attractor). We achieved this by defining **W** via its eigen-decomposition:

$$\mathbf{W} = \mathbf{V} \mathbf{D} \mathbf{V}^{-1} \tag{21}$$

where the eigenvectors (columns of V, denoted as v_j , for j = 1, ..., N, with elements v_{ij} , for i, j = 1, ..., N) were generated by the following process:

1. Generating a random vector:

for i = 1, ..., N.

$$u_i \stackrel{\text{lid.}}{\sim} \mathcal{N}(0, 1)$$
(22)

1770

2. Making the first 10% of vectors overlapping so that the resulting matrix is non-normal:

$$v_{ik} = \nu_i + \epsilon_{ik} \tag{23}$$

where
$$\epsilon_{ik} \sim \mathcal{N}(0, 0.05^2)$$
 (24)

for i = 1, ..., N and k = 1, ..., K with K = 0.1 N.

3. Making the the other 90% of vectors orthogonal:

$$\mathbf{v}_{K+k}$$
 = the *k*th column of Nullspace($\mathbf{v}_1, \dots \mathbf{v}_K$) (25)

for k = 1, ..., N - K

4. Unit normalizing each vector:

$$\mathbf{v}_k \leftarrow \frac{\mathbf{v}_k}{\|\mathbf{v}_k\|_2^2} \tag{26}$$

and the eigenvalues (λ_i , for i = 1, ..., N, the diagonal elements of the diagonal matrix **D**) were generated by the following process

1. Generating random (real) values:

$$\lambda_i \stackrel{\text{Id.}}{\sim} \text{Uniform}(0, 0.8)$$
 (27)

1775

for i = 1, ..., N - 2.

2. Creating a pair of neutrally stable eigenmodes:

$$\lambda_{N-1} = \lambda_N = 1 \tag{28}$$

The stimulus cue inputs were set to

$$\mathbf{h}^{(c)} = \mathbf{K} \begin{pmatrix} \cos\left(\frac{(c-1)\pi}{3}\right) \\ \sin\left(\frac{(c-1)\pi}{3}\right) \\ 1 \end{pmatrix}$$
(29)

¹⁷⁷⁶ for cues c = 1, ..., 6, and we considered two forms for K: either $\mathbf{K} = [\mathbf{v}_{N-1}, \mathbf{v}_N, \mathbf{v}_{r_1}] + [\mathbf{v}_{r_2}, \mathbf{v}_{r_3}, \mathbf{v}_{r_4}]$ (Extended Data ¹⁷⁷⁷ Fig. 1c and Extended Data Fig. 8a,d; as in the original formulation⁶) or $\mathbf{K} = [\mathbf{v}_{r_1}, \mathbf{v}_{r_2}, \mathbf{v}_{r_3}]$ (Extended Data Fig. 8b,e), ¹⁷⁷⁸ where r_1, r_2, r_3, r_4 were randomly drawn integers over the range 1 to N - 2. The first formulation of K ensured that ¹⁷⁷⁹ stimulus cue inputs partially align with the persistent subspace, whereas the second formulation of K ensured ¹⁷⁸⁰ that stimulus cue inputs align only with random directions.

1781 1.5.4 Feedforward network model

The linear feedforward network model that we used (Extended Data Fig. 1d and Extended Data Fig. 8c,f) has been described previously (see Refs. 21,26). (For pedagogical purposes, we used the simplest set up consisting of a feedforward chain of neurons, see below. However, using a more general network model that contained 'hidden' feedforward chains²¹ did not affect our analyses except for Extended Data Fig. 10e which, in contrast to the simple feedforward chain, could display overlap values greater than 0.5.) There were no separate excitatory and inhibitory populations in this model, and the weight matrix included a single chain running from neuron 1 to neuron *N*:

$$W_{ij} = \delta_{(i-1),j} \tag{30}$$

1782 for cell-pairs i, j = 1, ..., N.

The stimulus cues provided random inputs delivered to only the first 10 neurons so that each input could pass through the feedforward network:

$$h_i^{(c)} \stackrel{\text{iid.}}{\sim} \mathcal{N}(0, 1)$$
 (31)

1783 for cues c = 1, ..., 6 and cells i = 1, ..., 10.

1784 1.6 Canonical nonlinear systems with two stable fixed points

In order to illustrate the applicability of our analysis of optimal information loading in linear dynamical systems to the behaviour of nonlinear dynamical systems, we first studied two variants (either symmetric or non-symmetric) of a canonical nonlinear system that can exhibit two stable fixed points. (These systems are closely related to the damped, unforced Duffing oscillator which is a classic example of a [non-symmetric] system that can exhibit two stable fixed points. Additionally, the analysis of these systems also holds for the Duffing oscillator.)

The dynamics of the first system (which has a symmetric Jacobian matrix) are governed by

$$\frac{dx_{1}(t)}{dt} = x_{1}(t) - x_{1}^{3}(t)$$

$$\frac{dx_{2}(t)}{dt} = -x_{2}(t)$$
(32)

and the dynamics of the second system (which has a non-symmetric Jacobian matrix) are governed by:

$$\frac{dx_1(t)}{dt} = x_1(t) - x_1^3(t) + 3x_2(t)$$

$$\frac{dx_2(t)}{dt} = -x_2(t)$$
(33)

We used a cubic polynomial in Eqs. 32 and 33 because it is the lowest order polynomial that allows a system 1790 to exhibit 2 stable fixed points. Both systems exhibit 3 fixed points: both have a saddle point at the origin and both have 2 asymptotically stable fixed points at $(\pm 1, 0)$ (see Extended Data Fig. 6 for the state space dynamics of these two systems).

We solved the dynamics of Eqs. 32 and 33 using a first-order Euler approximation starting from t = 0 with a discretization time step of 0.02 (note time was unitless for this model).

.7 Analysis methods 1796

Here we describe methods that we used to analyse neural data. Whenever applicable, the same processing and analysis steps were applied to both experimentally recorded and model simulated data. As a first step in all 1798 our analyses, in line with previous work analysing neural population dynamics⁸⁴, we removed the stimulus cueindependent time-varying mean activity from each neuron's firing rate time series (see Fig. 5a for an example). (This was done separately for training and test data for cross-validated analyses, see below.) In most of our 1801 analyses, neural activities were aligned to stimulus cue onset defined to be at t = 0. However, due to the variable 1802 delay duration of the task (Fig. 1a), experimentally recorded neural activities were also aligned to go cue onset 1803 for analyses that required incorporating the late delay and go epochs (i.e. beyond the first 1.65 s after the stimulus cue onset; Fig. 5b-c, Extended Data Fig. 10a-c,g). For simulated neural activities, this was not necessary, as 1805 we always simulated our networks in a fixed-delay task for ease of analysis, even if they were optimized for a 1806 variable-delay task in accordance with how our experimental monkey subjects were trained. 1807

Identifying amplifying, persistent, and other subspaces in network dynamics 71

In order to understand the dynamics of neural networks with potentially complex and high-dimensional dynamics, 1809 and the way these dynamics depend on initial conditions, we identified specific subspaces within the full state space of these networks that were of particular relevance for our analyses. These subspaces served dual roles. 1811 First, as 'intervention tools', to ascertain their causal roles in high dimensional network dynamics, we used them 1812 to constrain the initial conditions of the dynamics of our networks (see also Methods 1.7.2). Second, as 'mea-1813 surement tools', to reveal key aspects of the high-dimensional dynamics of neural networks, we used them to 1814 project high-dimensional neural trajectories into these lower dimensional subspaces (see also Methods 1.7.3). 1815

Our main analyses relied on identifying the most persistent and most amplifying modes of a network. This re-1816 quired dynamics that were linear—either by construction, or by (locally) linearizing or linearly fitting dynamics that were originally nonlinear (see Table 1). We computed the most persistent mode(s) in one of two different ways. First, for networks that were either guaranteed to have stable dynamics by construction (i.e. those constructed 1819 de novo; Figs. 3 and 4 and Extended Data Figs. 4, 5 and 8), or were confirmed to be always stable in practice 1820 (i.e. those constructed by local linearization; Extended Data Fig. 3e,f, Extended Data Fig. 6, and Extended Data 1821 Fig. 7), we simply used the eigenvector(s) of the weight matrix W associated with the eigenvalue(s) that had the 1822 largest real part(s). Second, for networks that were fitted to nonlinear dynamics or recorded data, and whose 1823 dynamics could thus not be guaranteed to be stable (Fig. 5f, Fig. 6e, Extended Data Fig. 3f, Extended Data 1824 Fig. 7d, and Extended Data Fig. 10f,g), we used the eigenvectors of W associated with the largest real eigenvalues that were less than or equal to 1 + δ (with δ = 0.05) (i.e. we find the slowest, or most persistent, modes of 1826 the network—the δ was mostly relevant only for the after-go-time networks of Fig. 6 and Extended Data Fig. 12 1827 which exhibited eigenvalues substantially greater than 1 and setting δ less than 0.05 did not substantially change 1828 our results). (Note that an eigenvalue of W of 1 corresponds to an eigenvalue of 0 of the associated Jacobian of 1829

the dynamics due to the leak term.) 1830

For computing the most amplifying modes, we performed an eigendecomposition of the associated Observability Gramian $\mathbf{Q}^{52,61}$. Specifically, we obtained \mathbf{Q} by solving the following Lyapunov equation:

$$\left(\tilde{\mathbf{W}} - \mathbf{I}\right)^{\top} \mathbf{Q} + \mathbf{Q} \left(\tilde{\mathbf{W}} - \mathbf{I}\right) + \mathbf{C}^{\top} \mathbf{C} = \mathbf{0}$$
(34)

where $\tilde{\mathbf{W}}$ is the 'stabilized' weight matrix of the dynamics (and the $-\mathbf{I}$ terms represent the effect of the leak on the 1831 Jacobian of the dynamics, Eq. 1) and C is the read-out matrix of the network. The most amplifying mode(s) of the 1832 network are given as the eigenvector(s) of Q associated with the largest eigenvalue(s). Again, for networks that 1833 were guaranteed to have stable dynamics by construction (Figs. 3 and 4, Extended Data Fig. 7a-c, Extended 1834 Data Fig. 5, and Extended Data Fig. 8), $\tilde{\mathbf{W}} = \mathbf{W} - \epsilon \mathbf{I}$, where W is the original weight matrix of the dynamics and $\epsilon = 0.01$ (to ensure dynamical stability). For other networks, i.e. either linear networks fitted to experimental 1836 data (Fig. 5f and Extended Data Fig. 10f,g), linear networks fitted to simulated nonlinear dynamics (Fig. 6e, 1837 Extended Data Fig. 3f, and Extended Data Fig. 7d), or local linearizations of nonlinear dynamics (Extended Data Fig. 6, Extended Data Fig. 3e, and Extended Data Fig. 7a–c), we used $\tilde{W} = W$ unless the largest eigenvalue λ_{max} of W was greater than or equal 1, in which case we used $\tilde{W} = W - (\lambda_{\text{max}} - 1 + \epsilon) I$, to ensure that the

linear dynamics with W were stable (which is required for calculating Q). For networks obtained by fitting neural 1841 responses (experimentally recorded or simulated; Fig. 5f, Fig. 6e, Extended Data Fig. 3f, Extended Data Fig. 7d, 1842 and Extended Data Fig. 10f,g), C was obtained by fitting those responses (Methods 1.4.3), as we wanted to 1843 understand how the fitted dynamics taking place in a latent space can generate the most discriminable fluctuations (the principal components of) the neural responses to which they are related by this read-out matrix (although in 1845 using C = I did not change our results substantially). For all other networks (Figs. 3 and 4, Extended Data Fig. 7a– 1846 Extended Data Fig. 3e, Extended Data Fig. 5, and Extended Data Fig. 8), we simply used C = I, as the activity С 1847 of these networks was supposed to be read out in the same space within which their dynamics took place. 1848

We also applied methods which did not rely on the linearization (or linear fitting) of network dynamics. Our goal 1849 was to develop basic intuitions for how much the dynamics of the different simulated nonlinear networks of Fig. 2 1850 and Extended Data Fig. 2 used the persistent subspace of their dynamics. For this, we determined the 'persistent subspace' as the subspace spanned by the 5 principal components of the final 500 ms of neural activities (x) 1852 across all 6 cue conditions, corresponding to 6 distinct attractors, and the 'persistent nullspace' of the network as 1853 the 45-dimensional subspace orthogonal to the persistent subspace. For plots showing the projection of network 1854 activities within the persistent subspace (Extended Data Fig. 2b,f and Extended Data Fig. 2c-d and g-h, bottom) 1855 we used the first two principal components of the full, five-dimensional persistent subspace of the network, as 1856 determined above. For plots showing the projection of network activities to persistent vs. cue-aligned directions 1857 (Fig. 2d,j, and Extended Data Fig. 2c-d and g-h, top right), 'persistent PC1' was determined as the direction 1858 spanning the two persistent states corresponding to the two cue conditions being illustrated (i.e. as above, spanning the final 500 ms of neural activities across the two cue conditions), and 'initial PC1 (orthogonalized) 1860 was determined as the the direction spanning the two initial conditions corresponding to the two cue conditions 1861 being illustrated, orthogonalized with respect to the corresponding persistent PC1. 1862

1.7.2 Subspace-constrained initial conditions

When using the subspaces identified above as 'intervention tools', to constrain the initial conditions of our net-1864 works, we either used the single top most persistent or amplifying mode for linear networks with low-dimensional coding spaces (including the linearized canonical nonlinear attractor dynamical system; Figs. 3 and 4 and Ex-1866 tended Data Figs. 5 and 6), or numerically optimized initial conditions within the corresponding higher-dimensional 1867 subspaces (Fig. 2f,I, Extended Data Fig. 2c,d,g,h, Extended Data Fig. 7; see also Methods 1.3 and Methods 1.4). When the persistent subspace was extracted from neural responses (rather than from the dynamical equations of the network, Methods 1.7.1; Fig. 2f,I, Extended Data Fig. 2c,d,g,h, Extended Data Fig. 7a) we used different 1870 sets of simulations to generate data from which we could estimate the persistent subspace (as explained above). 1871 and to analyse network dynamics when initialized within these subspaces. In all cases, for a fair comparison, the magnitude of initial conditions was fixed (Methods 1.3.1, Methods 1.4.1), and only their direction was affected by constraining them to one of these subspaces. 1874

1875 1.7.3 Measures of subspace overlap

In order to measure the overlap of high dimensional neural dynamics with the subspaces we identified, we used one of two methods. First, for analysing network dynamics across two conditions chosen to correspond to 'opposite' stimulus cues (Fig. 2d,j, Fig. 3c,d,e, Extended Data Fig. 2c,d,g,h, Extended Data Fig. 6c,d, and Extended Data Fig. 5), such that the coding part of the persistent subspace was one-dimensional, we simply measured the projection of neural dynamics onto the first eigenvector (i.e. the eigenvector associated with the largest real eigenvalue) of the corresponding subspace using a dot product:

activity along mode(t) =
$$\mathbf{u}^{\top} \mathbf{x}(t)$$
 (35)

where **u** may correspond to the most persistent, or the most amplifying mode, or the first PC of the persistentorthogonalized cue subspace (as defined above). We also used the same measure for visualising the quality of fit of linear neural network dynamics to experimental data (Methods 1.4.3) with **u** being the first PC of the full state space of neural firings rates (Fig. 5e). In those cases, when **u** had to be estimated from neural responses (Fig. 2d,j, Fig. 5e, Extended Data Fig. 2c,d,g,h), we used a cross-validated approach, using different subsets of the data to determine **u** and **x**(*t*) (from a single split of the data). In other cases, **u** was determined from the truly deterministic dynamics of the system and thus there was no need for cross-validation.

Second, to measure subspace overlaps for *d*-dimensional neural activities across multiple conditions and time points within coarser time bins (Fig. 4c, Fig. 5f, Fig. 6e, Extended Data Fig. 3f, Extended Data Fig. 7d, Extended Data Fig. 8d–e, Extended Data Fig. 11c, Extended Data Fig. 12d, Extended Data Fig. 13d, and Extended Data Fig. 10b,c,f,g), thus corresponding to high-dimensional coding sub-spaces, we used the following properly

normalized measure:

variance explained
$$(t, t') = \frac{\operatorname{Tr}(\mathbf{U}^{\mathsf{T}}(t') \mathbf{\Sigma}(t) \mathbf{U}(t'))}{\operatorname{Tr}(\mathbf{P}^{\mathsf{T}}(t) \mathbf{\Sigma}(t) \mathbf{P}(t))}$$
 (36)

where $\Sigma(t)$ is the covariance matrix of neural activities across conditions and raw (1-ms) time points within time bin t, the columns of P(t) are the first principal components of neural activities within time bin t (i.e. the eigenvectors of $\Sigma(t)$ associated with the largest eigenvalues), and U(t') is the subspace of interest with respect to which overlaps are computed (which itself may or may not depend on time, see below). The time resolution of t and t' (i.e. the duration of time bins within which data was used to compute the corresponding terms at a given t or t'), the choice of U(t'), and the number of vectors used for constructing U(t') and P(t) depended on the analysis (see below).

Specifically, for measuring subspace overlap between neural activity and persistent vs. amplifying modes (Fig. 4c, 1890 Fig. 5f, Fig. 6e, Extended Data Fig. 3f, Extended Data Fig. 7d, Extended Data Fig. 8d-e, and Extended Data 1801 Fig. 10f,g), we set U(t') = U where the columns of U are the first d/4 eigenvectors of the most persistent or 1892 amplifying subspace (orthogonalized using a QR decomposition for the most persistent modes-this was not necessary for most amplifying modes which are orthogonal by construction), or d/4 randomly chosen orthonormal 1894 vectors as a control (shown as 'chance'; computed analytically as 1/4 for 'de novo' linear networks (Fig. 4c and 1895 Extended Data Fig. 8d–e), and numerically for fitted linear networks, also yielding values of approximately 1/4, 1896 Fig. 5f, Fig. 6e, Extended Data Fig. 7d, and Extended Data Fig. 10f). P(t) contained the first d/4 principal 1897 components. In this case, a value of 1 for this metric implies that the d/4 directions of greatest variability in 1898 neural activity overlap exactly with the d/4-dimensional subspace spanned by U. The time resolution of t was 1899 20 ms (for clarity, bins to be plotted were subsampled in the corresponding figures). Note that when this analysis was performed on linear networks fitted to neural data (experimentally recorded or simulated), U, P(t), and $\Sigma(t)$ were all obtained from the same fitted linear network (i.e. no cross-validation). Specifically the parameters of the 1902 network were used to determine U (see Methods 1.4.3), and the neural responses these fitted linear dynamics 1903 generated (rather than the original neural responses that were fit by the linear model) were used to determine 1904 $\Sigma(t)$ and thus P(t). See Methods 1.8 for computing the significance of these overlaps (and their differences). 1905 When analysing optimized ring attractor networks (Extended Data Fig. 3e,f), we used 2-dimensional subspaces 1906 (rather than d/4-dimensional subspaces) because we found empirically that the obtained ring attractors lay in a 1907 2-dimensional subspace.

For analyzing subspace sharing between different task epochs (Extended Data Fig. 11c, Extended Data Fig. 12d, Extended Data Fig. 13d, and Extended Data Fig. 10b), U(t') contained the top *k* principal components (PCs) of neural activity within the time bin indexed by *t'* (we used *k* = 10 for the monkey data and *k* = 4 for our models because the models typically exhibited lower dimensional dynamics), while P(t) included all PCs within the time bin indexed by *t*. For these, we performed principal components analysis with dimensions corresponding to neurons and data points corresponding to time points and cue conditions. The time resolution of both *t* and *t'* was 250 ms, such that the time periods (relative to cue onset) that we used were -500 to -250 ms (spontaneous epoch), 0 to 250 (cue epoch), 1250 to 1500 ms (delay epoch), and the first 250 ms after the go cue, i.e. *t*_{go} to *t*_{go} + 0.25 s (go epoch). In this case, U(t'), P(t) and $\Sigma(t)$ were obtained by fitting all the available neural data (i.e. no cross-validation). See also Ref. 64 for an 'alignment index' metric that is closely analogous to this use of this metric, but uses U(t') and P(t) that contain the same number of eigenvectors, as in our previous case, and are estimated in a cross-validated way, using different sets of trials.

For showing how much variance the top 2 delay epoch PCs capture over time (Extended Data Fig. 10c), we set U(t') = U where the columns of U are the first 2 principal components of neural activities over the time period 750 to 250 ms before the go cue, i.e. $t_{go} - 0.75$ to $t_{go} - 0.25$ s, and P(t) also includes the top 2 principal components. The resolution for t was 10 ms (for clarity, bins to be plotted were subsampled in the corresponding figure). In this case, we estimated U and P(t) in a cross-validated way—we estimated U using training data and P(t) and $\Sigma(t)$ using test data, and we show results averaged over 10 random 1:1 train:test splits of the data. See also Ref. 6 for a measure that is closely related to this use of this metric, but uses the number of neurons in the denominator instead of the total variance.

1929 1.7.4 Linear decoding

We fitted decoders using linear discriminant analysis to decode the stimulus cue identity from neural firing rates
 (Fig. 2e,f,k,l, Fig. 4a,b, Fig. 5b,c, Fig. 6c,d, Extended Data Fig. 7c, Extended Data Fig. 3d, Extended Data Fig. 8a–
 c,Extended Data Fig. 10a,h, Extended Data Fig. 11a,b, Extended Data Fig. 12b,c, and Extended Data Fig. 13b,c).
 We constrained the decoders to be 2-dimensional (in line with previous studies⁶) because this was a sufficient
 dimensionality to decode responses. (We also trained decoders using logistic regression in the full activity space
 and obtained qualitatively similar results; not shown.) We primarily considered two types of decoding analyses:

we either trained decoders on late delay activity and tested on all time points ('delay-trained decoder', e.g. Fig. 4a), or we trained decoders separately at every time point and tested on all times ('full cross-temporal decoding', e.g. 1937 Fig. 4b). In all cases, we measured decoding performance in a cross-validated way, using separate sets of neural 1938 trajectories to train and test the decoder, and we show results averaged over 10 random 1:1 train:test splits of the data. For delay-trained decoders, training data consisted of pooling neural activity over a 500 ms time interval 1940 (the time interval is shown by a horizontal black bar in all relevant figures), and tested the thus-trained decoder 1941 with data in each 1 ms time bins across the trial (for clarity, test bins to be plotted were subsampled every 10 ms 1942 in the corresponding figures). For full cross-temporal decoding, we binned neural responses into 10 ms time bins and trained and tested on all pairs of time bins (specifically, we plotted mean decoding performance across the 10 1-ms raw time bins corresponding to each 10-ms testing bin). We used a shrinkage (inverse regularisation 1945 parameter on the Euclidean norm of decoding coefficients) of either 0.5 or 1 (depending on the ratio of features to number of observations) for all main figures (we also tested various other values and found qualitatively similar 1947 results; not shown). Chance level decoding was defined as 1/C, where C = 2 or 6 is the number of cue conditions that need to be decoded (Tables 1 and 3).

1.7.5 Quality of fit for linear models fitted to neural responses

¹⁹⁵¹ When fitting linear models to neural data (experimentally recorded or simulated; Methods 1.4.3) we used a cross-¹⁹⁵² validated approach for measuring the quality of our fits, with a random 1:1 train:test split of the data (Fig. 5d). For ¹⁹⁵³ this, we first fitted the model on training data ($\mathbf{x}_{*}^{(c)} = \mathbf{x}_{\text{train}}^{(c)}$ in Eq. 13). The quality of fit was then computed on the ¹⁹⁵⁴ test data, $\mathbf{x}_{\text{test}}^{(c)}$, as the fraction of variance of $\mathbf{x}_{\text{test}}^{(c)}(t)$ explained by the simulated responses (after the appropriate ¹⁹⁵⁵ projection, i.e. $C \mathbf{x}^{(c)}(t)$), across all 20 dimensions weighted by D (all parameters, including P, C and D, were set ¹⁹⁵⁶ to their values obtained by fitting the training data). In other words, we computed the Pearson R^2 with respect to ¹⁹⁵⁷ the identity line using the mean squared error, ε^2 in Eq. 12, with the momentary error in Eq. 13 computed using ¹⁹⁵⁸ $\mathbf{x}_{*}^{(c)} = \mathbf{x}_{\text{test}}^{(c)}$. Once the quality of fit for this split was thus established, we conducted all further analysis involving ¹⁹⁵⁹ fitted linear models with the model that was fit to the training half of this split.

As a meaningful lower bound on our quality of fit measure, we also computed the same measure (i.e. fitting a linear dynamical system to training data and calculating the quality of fit using test data) for 100 different time-shuffled controls of the original train:test split of the data (Methods 1.4.3), such that we shuffled time bins coherently between the training and the test data, across neurons and conditions (Fig. 5d, dark gray).

To calibrate how much our fits were limited by the noisiness of the data, we also computed the quality of fit directly between $\mathbf{x}_{\text{train}}^{(c)}(t)$ and $\mathbf{x}_{\text{test}}^{(c)}(t)$ (i.e. using the mean squared error, ε^2 in Eq. 12, with the momentary error redefined as $\mathbf{e}^{(c)}(t) = \mathbf{x}_{\text{train}}^{(c)}(t) - \mathbf{x}_{\text{test}}^{(c)}(t)$) for 100 random 1:1 train:test splits of the data (Fig. 5d, light gray). The extent to which the R^2 computed with this control was below 1 reflected the inherent (sampling) noise of the experimental data that limited the quality of fit obtainable with any parametric model, including ours that was based on linear dynamics. Moreover, a cross-validated R^2 computed with our fits that was higher than the R^2 obtained with this control (Fig. 5d dark and light blue vs. light gray) meant that the inherent assumption of linear dynamics in our model acted as a useful regularizer to prevent the overfitting that this overly flexible control inevitably suffered from. See more in Methods 1.8 on statistical testing for our quality of fit measure.

¹⁹⁷³ When fitting to simulated neural data, we obtained high quality of fits using the same measure ($R^2 > 0.95$, not ¹⁹⁷⁴ shown).

1.7.6 Overlap between the coding populations during the cue and delay epochs

To test whether separate neural populations encode stimulus information during the cue and delay epochs (Extended Data Fig. 10e), we trained (non-cross validated) decoders to decode cue identity using logistic regression on either cue-epoch activity ('cue-trained'; the first 250 ms of activity after cue onset) or delay-epoch activity ('delay-trained'; 1250–1500 ms after cue onset). We used an L2 regularisation penalty of 0.5 (we also tested other regularisation strengths and observed no substantial changes in our results). We took the absolute value of decoder weights as a measure of how strongly neurons contributed to decodability (either positively or negatively). We then binarized the absolute 'cue-trained' and 'delay-trained' weights using their respective median values as the binarization threshold. (This binarization reduces a potential bias effect from large or small weight values in our analysis.) Our measure of overlap between the coding populations during the cue and delay epochs, was then simply the inverse normalized Hamming distance between these two sets of binarized weights:

overlap =
$$1 - \left\langle \left| w_{n,c}^{\text{cue}} - w_{n,c}^{\text{delay}} \right| \right\rangle_{n,c}$$
 (37)

where $w_{n,c}^{\text{cue}}$ ('cue trained') and $w_{n,c}^{\text{delay}}$ ('delay trained') is the binarized weight of neuron *n* in cue condition *c* during the cue and delay epochs, respectively, and $\langle \cdot \rangle_{n,c}$ denotes taking the mean across neurons and cue conditions. For

¹⁹⁷⁸ completely overlapping populations, this measure takes a values of 1, for completely non-overlapping populations, ¹⁹⁷⁹ it takes a values of 0, and for random overlap (shown as 'chance') it takes a values of 0.5.

For the shuffle controls, we randomly permuted the neuron indices of the delay-trained weights (such that using 1980 the median as a threshold thus resulted in values close to 0.5, i.e. chance level; Extended Data Fig. 10e). We show results (for both the original analysis and shuffle control) for 10 random halves of the data (equivalent to 1982 the training halves of 10 different 1:1 train:test splits). We also tested a variety of percentile values other than the 1983 median and our results did not change substantially (choosing a threshold other than the median causes both 1984 the data and shuffle controls to have overlap values lower than those that we obtained with the median as the 1985 threshold, but it does not substantially affect the difference between them). As an additional control, we also 1986 removed neurons that did not contribute to decodability; we removed neurons that had a thresholded weight of 1987 0 for all 6 cue conditions in both the cue and delay epochs. This resulted in removing 13.3 neurons on average for monkey K and 33.5 neurons for Monkey T (when using the median as the threshold) and our results did not 1989 change substantially (not shown). 1990

¹⁹⁹¹ 1.7.7 Finding fixed points

For finding the fixed points of nonlinear network dynamics (Fig. 2d,j, Extended Data Fig. 2b,f, Extended Data Fig. 3a, and Extended Data Fig. 11d), we used a slow-point analysis method¹⁷ that searches for an x for which the L2 norm of the gradient determined by the autonomous dynamics of the network is below a threshold. Note that this was only possible in model neural networks as the method requires access to the equations (and parameters) defining the true (nonlinear) dynamics of a system.

Specifically, for network dynamics governed by (cf. Eqs. 1 and 2)

$$\frac{\mathrm{d}\mathbf{x}(t)}{\mathrm{d}t} = \psi(\mathbf{x}(t)) , \qquad (38)$$

for some function ψ , we sought to find points \mathbf{x}^* such that $\|\psi(\mathbf{x}^*)\|_2$ is small. To achieve this, we drew 1000 x's from a spherical Gaussian distribution with mean 0 and variance 10 (the large variance helps to ensure that we cover a large part of state space) and we optimized each \mathbf{x} to minimize $\|\psi(\mathbf{x})\|_2$ using gradient descent with gradients obtained by back-propagation with an Adam optimizer⁸³. We used an adaptive learning rate (which we found worked substantially better than a fixed learning rate in this scenario) that started at 0.1 and halved every 1000 training iterations (we used 5000 training iterations in total). Finally, we identified the \mathbf{x} 's obtained at the end of optimization as asymptotically stable fixed points, \mathbf{x}^* , if $\|\psi(\mathbf{x})\|_2 < 0.001$ and if the largest real part in the eigenvalues of the linearization of $\psi(\mathbf{x})$ around \mathbf{x}^* was less than 0.

²⁰⁰⁵ 1.7.8 Correlations between initial and final neural firing rates

To measure correlations between initial and final simulated activities, we used the Pearson correlation coefficient (with respect to the identity line) between initial and final mean-centered firing rates across neurons within the same simulation (i.e. no cross-validation; Fig. 2b,h; insets). Histograms show the distribution of this correlation across 6 cue conditions (and the 10 different networks, each simulated 10 times, see above) using a kerneldensity estimate (Fig. 2c,i, Extended Data Fig. 2c,d,g,h, and Extended Data Fig. 3c).

2011 1.8 Statistics

²⁰¹² We performed statistical hypothesis testing in two cases.

First, we tested whether the quality of fit of linear models to experimental data was sufficiently high using permu-2013 tation tests. To construct the distribution of our test statistic (cross-validated R^2 , see also Methods 1.7.5) under the null hypothesis, we used n = 100 different random time shuffles of the data (Fig. 5d, dark gray), such that we 2015 shuffled time bins coherently between the training and the test data, across neurons and conditions, and for each 2016 shuffle used the same random 1:1 train:test split as for the original (unshuffled) data. For additional calibration, 2017 we also constructed the distribution of our test statistic under the alternative hypothesis that all cross-validated 2018 errors were due to sampling noise differences between the train and test data. For this, we used n = 100 random 2019 1:1 train:test splits of the (original, unshuffled) data, and measured the guality of fit directly between the test data and the training data (rather than a model fitted to the training data, see also Methods 1.7.5; Fig. 5d, light gray). In both cases, we computed the two-tailed p-value of the test statistic as computed on the real data (Fig. 5d, blue lines) with respect to the corresponding reference distribution. 2023

Second, we also used a permutation test-based approach to test whether the experimentally observed overlaps with persistent and amplifying modes (or their differences) were significantly different from those expected by chance. For testing the significance of overlaps in a given time step, we constructed the distribution of our

test statistics (the overlap measures; Methods 1.7.3) under the null hypothesis by generating n = 200 random 2027 subspaces within the space spanned by the 20 PCs we extracted from the data (Methods 1.4.3), dimensionality 2028 matched to the persistent and amplifying subspaces (i.e. 5 orthogonal dimensions), and computed the same 2029 subspace overlap measures for the data in the given time step with respect to these random subspaces (Fig. 5f and Extended Data Fig. 10f-g; gray line and shading). For testing the significance of differences between overlaps 2031 (amplifying vs. persistent at a given time step, or amplifying or persistent between two different time steps), our 2032 test statistic was this difference (i.e. a paired test), and our null distribution was constructed by measuring it for 2033 = 200 pairs of random subspace overlaps at the appropriate time step(s). Once again, in all these cases we n computed the two-tailed p-value of the test statistic as computed on the real data (Fig. 5f and Extended Data 2035 Fig. 10f-g, green and red lines) with respect to the corresponding reference distribution. 2036

Note that we did not compute p-values across multiple splits of the data because this led to p-value inflation as we increased the number of splits. Instead, we repeated all relevant analyses on 10 different random 1:1 train:test splits to see if our results were robust to the choice of data split. Indeed, we obtained qualitatively and quantitatively (in terms of p-values for quality of fits, and overlaps) similar results for all these splits.

Permutation tests do not assume that the data follows any pre-defined distribution. No statistical methods were used to predetermine experimental sample sizes. Sample sizes for permutation tests (n above) were chosen so as to be able to determine p-values to a precision of 0.02 (quality of fits) or 0.01 (subspace overlaps).

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Extended data figures



Extended Data Fig. 1 | **Dynamics of network models of working memory. a**, Neural network dynamics in a bump attractor network⁵ performing the task shown in Fig. 1a. Left: trajectory in neural state space in a single cue condition during the cue period (pale purple line, ending in pale purple circle) and delay period (dark purple line). Purple arrow heads indicate direction of travel along the trajectory, black cross shows attractor state, gray arrow shows overlap between cue input and late delay activity. Center: time course of relative (i.e. mean-centered) firing rates of one neuron for two cue conditions (purple vs. blue, see also inset). Yellow lines indicate cue onset and offset times. Right: cross-temporal decoding of neural activity produced by the network across all 6 cue conditions. Pink rectangles indicate generalized decoding between the cue/early delay period and the late delay period and cyan rectangles indicate generalized decoding between time points in the late delay period. The gray tick on the color bar indicates chance-level decoding. **b**, Same as **a** but for a discrete attractors model^{5,28,30}. **c**, Same as **a** but for a linear integrator model with transient dynamics that are orthogonal to the attractor subspace⁶. **d**, Same as **a** but for a network optimized to perform the task shown in Fig. 1a using gradient descent (cf. Fig. 6d, right).



Extended Data Fig. 2 | Attractor network dynamics with or without constraints on the initial condition of the dynamics. a, Illustration of an attractor network with symmetric connections. b-d, Analysis of neural responses in symmetric attractor networks (such as shown in a). b, Sub-threshold activity for all 6 cue conditions (color trajectories) with initial conditions optimized within the full state space (Methods 1.3.1). Open circles show the optimized initial conditions and crosses show stable fixed points. We show neural activity projected onto the top two principal components of the persistent subspace. c, Analysis of neural responses when initial conditions are constrained to lie within the 5-dimensional persistent subspace. Top left: distribution of Pearson correlations between initial and final mean-centered neural firing rates across all 6 cue conditions and 10 networks (same as Fig. 2c, but for persistent subspace-constrained inputs, corresponding to green line in Fig. 2f). Top right: sub-threshold activity for 2 cue conditions in an example network (color trajectories; same as Fig. 2d, but for persistent subspace-constrained inputs, corresponding to green line in Fig. 2f). Open circles (with arrows pointing to them from the origin) show the optimized initial conditions, black crosses show stable fixed points, dashed gray line is the identity line. Horizontal axis (persistent PC1) shows neural activity projected on to the 1st principal component (PC1) of network activities at the end of the delay period (across the 2 conditions shown), vertical axis (initial PC1 (orthogonalized)) shows projection to PC1 of initial neural activities orthogonalized to persistent PC1. Bottom: same as b, but for persistent subspace-constrained inputs, corresponding to green line in Fig. 2f. d, Same as c, but for persistent nullspace-constrained inputs. Note that the distribution of Pearson correlations of neural firing rates (top left) is distinct from a delta function at 0 because we constrained the initial conditions in the space of sub-threshold activities (rather than firing rates). In the bottom panel, which shows sub-threshold activity, we see that indeed all the colored circles overlap at the origin, indicating orthogonality of the initial conditions to the persistent subspace. e, Illustration of an attractor network without a symmetric connectivity constraint. f-h, Same as b-d but for attractor networks without a symmetric connection constraint (i.e. panels f, g, and h, respectively correspond to the networks shown by the black, green, and red lines in Fig. 2). Note initial conditions being near the origin in f mean that they are strongly orthogonal to the persistent subspace (as in d, but without constraining them explicitly to be in the persistent nullspace).



Extended Data Fig. 3 | Dynamics of optimized ring attractor networks. a, Neural activity for 6 cue conditions (color trajectories) with optimized initial conditions in a ring attractor network with unconstrained connectivity (see Methods 1.3.1 and 1.3.4). Open circles show the optimized initial conditions and black crosses show fixed points. We show neural activity projected onto the top two principal components of the persistent subspace. Thus, all circles being near the origin means that initial conditions are strongly orthogonal to this subspace (cf. Extended Data Fig. 2f). b, Tuning curves for 6 neurons (purple curves) whose preferred angles (coloured crosses) correspond to the 6 cue conditions shown in a. c, Distribution of Pearson correlations between initial and final mean-centered neural firing rates across the 6 cue conditions and 10 networks (cf. Fig. 2i). d, Cross-temporal decoding of neural firing rate activity (cf. Fig. 2k). Note that only the first second of the delay period is shown on both axes because the dynamics of these networks, using a tanh nonlinearity, are faster than those shown in other figures (e.g. Fig. 2), using a ReLu nonlinearity (but the same time constant; Methods 1.2, and Table 1). e, Overlap (mean±1 s.d. across 10 networks) of the 2 locally most persistent (green), most amplifying (red), or random directions (black), obtained using a local linearization around the origin, with the 'persistent subspace' and 'persistent nullspace' of the original non-linear dynamics, obtained without linearization, and the subspace spanned by the 'optimal' initial conditions of the original nonlinear dynamics (cf. Extended Data Fig. 7a, bottom; see Methods 1.4.2 and 1.7.3). We used 2-dimensions from the local linearization because we found empirically that the ring attractor lay in a 2-dimensional subspace (see also a). f, Percent variance explained (mean±1 s.d. across 10 networks) by the subspace spanned by either the 2 most persistent (green) or 2 most amplifying (red) modes as a function of time for a 20-dimensional linear dynamical system fitted to the neural activities of the nonlinear ring network (Methods 1.4.3; cf. Extended Data Fig. 7d, bottom far right). We used 2-dimensions because we found empirically that the ring attractor lay in a 2-dimensional subspace (see also a and e). Gray lines show chance level overlap defined as the expected overlap with a randomly chosen 2-dimensional subspace.



Extended Data Fig. 4 | Analysing the total energy produced by different initial conditions in linear networks. a, The norm of neural activity integrated over time (i.e. a measure of total energy used by the network) for each of 1000 random initial conditions (10 initial conditions for each of 100, 100-neuron networks) relative to the energy produced by the most amplifying initial condition, plotted as a function of their overlap with the persistent mode for symmetric (top) and unconstrained (bottom) linear integrator networks. A positive value on the yaxis means that the total energy produced by the given random initial condition is greater than that produced by the most amplifying initial condition. Initial conditions are scaled so that they all produce the same level of persistent activity (i.e. the same level of performance) after 2 s of simulation. b, Same as a, but initial conditions are plotted as a function of their overlap with the most amplifying mode. Note that overlap with the most amplifying mode (but not in general with the most persistent mode) is strongly predictive of total energy (with an inverse relationship between the two). c, Overlap (mean±1 s.d. across the 100 networks from a and b) of optimal initial conditions (Eq. S39), producing an overlap of 1 with the persistent mode after a given delay length (x-axis) while using the minimal total energy over time (Eq. S38), with either persistent (green), most amplifying (red), or random (black) directions, for symmetric (top) and unconstrained (bottom) networks. In unconstrained networks, for very short delay lengths, initial conditions must align exactly with the persistent mode, by necessity (green lines at 0 s). For longer delay lengths, initial conditions make greater use of the most amplifying direction (red lines). d, Total energy over time (mean across the 100 networks from a and b; we do not show error bars for visual clarity) for dynamics starting from initial conditions that produce an overlap of 1 with the persistent mode after a given delay length (x-axis) and are aligned with the optimal initial condition (blue; i.e. the one using the least energy, cf. panel c), the most persistent (green), most amplifying (red), or a random direction (black), in symmetric (top) and unconstrained (bottom) networks. In unconstrained networks, for very short delay lengths, initialising along the most persistent mode achieves near-optimal energy-efficiency (green is close to blue), but for longer delay lengths, initialising along the most amplifying mode becomes more energy efficient (red is closer to blue). (Note that for symmetric networks (top), we have offset the curves for the most amplifying, persistent, and optimal directions because these 3 directions are the same and therefore produce the same total energy.)



Extended Data Fig. 5 | Analysis of linear networks of different sizes. a, Distributions of absolute overlap with the persistent mode for persistent (pale green), most amplifying (pale red), or random initial conditions (gray) across 100 randomly sampled linear symmetric (top) and unconstrained networks (bottom) consisting of either 10 (solid), 100 (dashed), or 1000 (dotted) neurons (cf. Fig. 3d). The persistent initial conditions produced delta functions at 1 (arrows). Results for persistent and most amplifying initial conditions are identical in symmetric networks (top). b, Time course of mean (across the 100 networks from a) absolute overlap with the persistent mode when starting network dynamics from persistent (green), most amplifying (red), or random initial conditions (black) in symmetric (top) and unconstrained networks (bottom) consisting of either 10 (solid), 100 (dashed), or 1000 (dotted) neurons (cf. Fig. 3e). Results for persistent and most amplifying initial conditions are identical in symmetric networks (top). c, Mean (across 100 networks) overlap of initial conditions that were optimized so as to generate persistent activity in 100-neuron noisy symmetric (top) and unconstrained (bottom) networks with 100 orthogonal modes ordered by their persistence (green) or amplification (red) (i.e. corresponding to the rank ordered eigenvectors of the weight matrix, green, or of the observability Gramian of the dynamics, red; Methods 1.7.1). In symmetric networks (top), the optimized initial conditions overlap only with the most amplifying mode and no other mode (note that the most persistent mode is identical to the most amplifying mode in this case). In unconstrained networks (bottom), optimized initial conditions overlap strongly with the most amplifying mode and only weakly with other modes. (The non-zero overlap with the most persistent mode is simply due to the fact that there is a non-zero overlap between the most persistent and amplifying mode in random networks, and it is at the level that would be expected based on this overlap.) d, Time course of mean (across the 100 networks from c) absolute overlap with the persistent mode for 100-neuron symmetric (top) and unconstrained networks (bottom) when the network is started from optimized initial conditions (blue), and for comparison for the most amplifying (red dashed) initial conditions (cf. Fig. 3e). Note the close agreement between the two indicating that the most amplifying mode is indeed optimal in these networks. Horizontal black bar on x-axis shows the time period in which we applied the cost function to optimize the initial conditions (Methods 1.7.3).



Extended Data Fig. 6 | Analysis of canonical nonlinear attractor systems. a, State space of a canonical nonlinear system with two attractors and a symmetric (top) and non-symmetric Jacobian (bottom, see also Methods 1.6, Supplementary Math Note S2; cf. Fig. 3b). Pale blue arrows show flow field dynamics (direction and magnitude of movement in the state space as a function of the momentary state). Black crosses indicate asymptotically stable fixed points (i.e. attractor states), dashed black line shows the separatrix (the manifold separating the basins of attraction of the two attractors). Thin green and red lines indicate the locally most persistent and amplifying modes around the origin, respectively (lines are offset slightly in the top panel to aid visualisation). Pale green, red, and gray arrows with open circles at the end indicate most persistent, amplifying, and random initial conditions, respectively. Blue ellipses show the fixed initial condition norm around the origin to highlight the different axis scales. Dark green, red, and black arrows show neural dynamics starting from the corresponding initial condition. b, Time course of dynamics of the system along the persistent mode (i.e. the projection onto the green line in a) when started from the persistent (green), most amplifying (red), or random (black) initial conditions for the symmetric (top) and the unconstrained system (bottom). c, Late overlap with the locally persistent mode as a function of initial overlap with the locally most amplifying mode in the canonical nonlinear systems shown in panels a-b (solid gray line) and, for comparison, in the linear networks of Fig. 3a-c (dashed gray line) for symmetric (top) and unconstrained systems (bottom). Late overlap is measured as the mean overlap of activity along the persistent mode (panel **b**, from t = 0.8 to t = 2 for the canonical nonlinear system; Fig. 3c, from t = 0.8 s to t = 2 s for the linear networks). Open circles and squares indicate the random (gray), persistent (pale green), and most amplifying (pale red) initial conditions used respectively in panels a and b for the canonical nonlinear system, and in Fig. 3b-c for the linear networks.



Extended Data Fig. 7 | Linear analyses of the nonlinear attractor networks of Fig. 2. a, Overlap (mean±1 s.d. across 10 networks) of the 5 locally most persistent (green), most amplifying (red), or random directions (black) of the symmetric (top) and unconstrained (bottom) networks from Fig. 2, obtained using a local linearization around the origin, with the 'persistent subspace' and 'persistent nullspace' of the original non-linear dynamics, obtained without linearization (as used in Fig. 2f and I, red and green), and the 5-dimensional subspace spanned by the 6 'optimal' initial conditions of the original nonlinear dynamics (used in Fig. 2b-e, h-k, and f and I, black). For comparison, we also show the overlap (mean±1 s.d. across 100 networks) of the single most persistent (pale green), most amplifying (pale red), and random (gray) direction with the optimal initial condition of the linear networks from Extended Data Fig. 5c,d ('optimal (lin. model)'). b, Time course of the overlap (mean±1 s.d. across 10 networks, s.d. not shown in bottom for visual clarity) of the linearized dynamics of symmetric (top) and unconstrained networks (bottom) with the subspace spanned by their most persistent modes when started from initial conditions that were optimized for the decoding accuracy of the nonlinear dynamics while constrained to be within the locally most persistent (green), most amplifying (red), or a random subspace (black). The linear dynamics, the persistent subspace wrt. which overlap is measured, and the subspaces within which initial conditions were constrained while being optimized, were all based on a local linearization of the nonlinear dynamics around the origin. Compare with Fig. 3e for the analogous plots for linear networks. For reference, blue line shows overlap of the same linearized dynamics when started from the initial conditions directly optimized for the decoding accuracy of the nonlinear dynamics without subspace constraints (used in Fig. 2b-e, h-k, and f and I, black). For consistency with Fig. 3b-e (where initial conditions were constrained to have unit norm), we scaled activity by the norm of the initial condition (which was constrained to be 3 here; Methods 1.4.2). c, Performance (mean±1 s.d. across 10 networks) of a delay-trained decoder (black bar indicates decoder training time period; Methods 1.7.4) on neural activity in stochastic nonlinear symmetric (top) and unconstrained networks (bottom) over time. Colors indicate initial conditions as in b. (Blue line shows same data as black line in Fig. 2f and I). Gray dotted line shows chance level decoding. Green, red, and blue lines are vertically offset slightly in the top panel to aid visualization. Compare with Fig. 4a (noise matched) for the analogous plots for linear networks (though with non-instantaneous inputs). d, Percent variance explained (mean±1 s.d. across 10 networks) by the subspace spanned by either the 25% (i.e. 5) most persistent (green) or 25% (i.e. 5) most amplifying (red) modes as a function of time for 20-dimensional linear dynamical systems fitted to the neural responses generated by the symmetric (top) and unconstrained (bottom) nonlinear networks when started from the same (optimized) initial conditions analyzed in b-c: constrained to be within the locally most persistent (far left), most amplifying (center left), or a random subspace (center right), as determined by the local linearization of the dynamics, or without subspace constraints (far right). Gray lines show chance level overlap defined as the expected overlap with a randomly chosen subspace occupying 25% of the full space (i.e. 5 dimensions). Compare with Fig. 4c for the analogous plots for linear networks (though with non-instantaneous inputs, and performancematched levels of noise, see also Supplementary Math Note S3) and with Fig. 5f and Extended Data Fig. 10f,g for analogous plots of linear dynamical systems fitted to experimental data.



Extended Data Fig. 8 | **Analysis of two variants of an integrator model and feedforward model. a**, Cross-temporal decoding of model neural activity (cf. Fig. 2e,k, Fig. 4b, and Fig. 5c) for a linear integrator model⁶ (see also Methods 1.5). Yellow lines indicate cue onset, offset, and go times. **b**, Same as **a** for the same model but for inputs aligned with purely random directions (as opposed to inputs aligned with both persistent and random directions as in the original formation of Ref.⁶). **c**, Same as **a** but for a linear feedforward network model^{21,26}. **d**, Percent variance explained by the subspace spanned by either the 25% most persistent (green) or 25% most amplifying (red) modes as a function of time for the linear integrator model from **a** (cf. Fig. 4c,b, Fig. 5f, and Fig. 6e). Yellow lines indicate cue onset, offset, and go times. Gray dotted line shows chance level overlap with a subspace spanned by 25 random orthogonal directions. **e**, Same as **d** for the same model but for inputs aligned with or a linear feedforward network model^{21,26}.



Extended Data Fig. 9 | **Recording locations for the two monkeys.** Left: recording locations in monkey K (T1-weighted image). In order to image the interior of the chamber, we filled the chamber with cut cottons soaked in iodine. In the upper picture, the yellow arrow indicates the principal sulcus. In the bottom picture, locations of the 11 by 15 grid holes were superimposed over the MR picture. Right: recording locations in monkey T (T2-weighted image). The bottom picture shows the location for the grid of the 32 semi-chronic electrodes. Yellow dots indicate electrode penetrations and recording sites, red dots indicate non-visited sites.



Extended Data Fig. 10 | **Supplemental analysis of experimental data and comparison to models. a**, Cross-temporal decoding analysis for monkey K (cf. Fig. 5c for the same analysis for monkey T and for explanation of plotting scheme and annotations). **b**, Subspace overlap between different task epochs, measured as the percent variance explained (PVE) by projecting neural activity from one task epoch (tested) through the top 10 PCs of another task epoch (fitted). Diagonal elements show the PVE within each task epoch. We show results for monkey K (left) and monkey T (right). **c**, Time course of overlap with delay epoch subspace, measured as the percent variance explained by the top 2 PCs obtained from delay period activity (black bar shows time period of activity from which these PCs were obtained) on held-out test data taken in different time bins. This metric is called the alignment index⁶⁴ and is very similar to that used in Ref. 6 (Methods 1.7.3). We show mean (over 10 different data splits) results for both monkeys. Yellow ticks on horizontal axis indicate cue onset, cue offset, and go times. (*Caption continued on next page.*)

Extended Data Fig. 10 | Supplemental analysis of experimental data and comparison to models (cont'd). d, Schematic of 3 different hypothetical scenarios for the relationship between cue and late delay activities (panels), illustrated in neural dynamics for 2 neurons and 2 cue conditions. Coloured curves show neural trajectories, black squares indicate cue onset, open circles indicate cue offset, and filled circles show late delay activity. Left vs. right: populations encoding the cue during cue and late delay periods are overlapping vs. non-overlapping, respectively. Top vs. bottom: cue and delay activities are non-orthogonal vs. orthogonal, respectively. (Note that we are not showing dynamics for non-overlapping, non-orthogonal dynamics because no overlap necessarily implies orthogonality.) e, Relationship between cue and late delay activities in various different models and our experimental recordings (x-axis). Top: population overlap measured as the mean difference between cue and delay epoch classifier weights (left for each model and data) and, as a control, when randomly shuffling classifier weights across neurons (right for each model and data) (Methods 1.7.6). Box plots show medians (black lines), guartiles (boxes), and 1.5 times the inter-guartile range (whiskers). Dotted gray line shows chance level overlap. Bottom: orthogonality measured as 1 minus the mean overlap between cue and delay epochs (given by the corresponding elements of the subspace overlap matrices shown in panel b and Extended Data Fig. 11c, center right). The discrete attractors, bump attractor, and integrator models show high overlap but low orthogonality. The simple feed-forward network shows high orthogonality but low overlap (note that recurrent networks with embedded feed-forward connectivity²¹ may show high overlap). The just-in-time network shows high overlap and orthogonality, similar to the experimental data in both monkeys. f-g, Same analysis as in Fig. 5f, but either after randomly shuffling data across time (but consistently across conditions and neurons, and applied to the same time period as in the main analysis; f, see also Methods 1.4.3), or applied to the late delay time period (without acrosstime shuffling) in which we do not expect information loading dynamics (g). h, Decoding of stimulus information within the subspace spanned by either the 25% most persistent modes (green), or the 25% most amplifying modes (red) in the original linear systems shown in Fig. 5f. Comparisons use two-sided permutation tests (*, p < 0.05; **, p < 0.01; n.s., not significant; see Methods 1.8) **i**, Top inset: original data analysis of overlaps repeated from Fig. 5f to indicate the comparisons (coloured numbers) we show in the table below (numbered columns). Bottom: table showing p-values (in each cell for experimental data, top: monkey K, bottom: monkey T) from two-sided permutation tests for each comparison of the main analysis (row 4, repeated from the main text associated with Fig. 5f) and the control analyses shown in panels f and g of this figure (rows 5-6). Top 3 rows show predictions for the sign of each comparison under different information loading strategies in unconstrained linear networks (Fig. 4c, bottom): using inputs aligned with random directions (1st row), persistent directions (2nd row), or the most amplifying directions (3rd row). In the column headings, pers., amp., and ch. respectively refer to overlap with most persistent, most amplifying and random subspaces (chance), to refers to the beginning of the analysis time window, i.e. cue onset (rows 1-5) or 1 s before the timing of the go cue (row 6), and $t_1 = t_0 + 1$ s refers to the end of the analysis time window. The colored numbers above each column correspond to the comparisons shown in the inset above the table. Gray indicates no significant difference between data points, red and blue indicate a significant difference for both monkeys where the first data point is respectively greater or smaller than the second data point, and pale red indicates a significant difference for one of the two monkeys (see Methods 1.8).



Extended Data Fig. 11 | Cue-delay and just-in-time trained networks. a–b, Same as Fig. 6c green and red, and Fig. 6d left and right, but with a regularisation strength of $\alpha_{nonlin}^{(2)} = 0.0005$ used during training (Methods 1.3.2). **c**, Subspace overlap between different task epochs, measured as the percent variance explained (PVE) by projecting neural activity from one task epoch (tested) through the top 4 PCs of another task epoch (fitted; cf. Extended Data Fig. 12d, Extended Data Fig. 13d, and Extended Data Fig. 10b). Diagonal elements show the PVE within each task epoch. We show results for cue-delay (left two panels) and just-in-time trained networks (right two panels) trained with either a regularisation strength of $\alpha_{nonlin}^{(2)} = 0.0005$ (left panel for each model, as in Fig. 6) or $\alpha_{nonlin}^{(2)} = 0.0005$ (right panel for each model, as in panels **a–b**). **d**, Neural activity plotted in the top two PCs of delay-epoch activity for all 6 initial conditions for cue-delay and just-in-time trained networks for each of the network-regularization combinations shown in **c** (cf. Extended Data Fig. 2b–d and f–h.) Purple traces show state-space trajectories, squares indicate cue onset, open circles indicate cue offset, and crosses indicate asymptotically stable fixed points, colours indicate cue condition as in Fig. 2d.



Extended Data Fig. 12 | **After-go-time trained networks. a**, Cost function for after-go-time training on the fixed delay task (Methods 1.3.3). Cue onset, cue offset, and go cue times are indicated by the yellow vertical lines. The boxcar shows the interval over which stable decoding performance was required (i.e. the cost was only applied after the go cue). **b–c**, Same as Fig. 6c orange and Fig. 6d center, but with a regularisation strength of $\alpha_{nolin}^{(2)} = 0.0005$ used during training and when either a random (**b** orange, **c** left) or a fixed delay task is used (**b** blue, **c** right, Methods 1.7.4). **d**, Subspace overlap between different task epochs, measured as the percent variance explained (PVE) by projecting neural activity from one task epoch (tested; cf. Extended Data Fig. 11c, Extended Data Fig. 13d, and Extended Data Fig. 10b) through the top 4 PCs of another task epoch (fitted) for the networks shown in **b–c**. Diagonal elements show the PVE within each task epoch. **e**, Neural activity plotted in the top two PCs of delay-epoch activity for all 6 initial conditions for random delay (left) and fixed delay (right) trained networks (cf. Extended Data Fig. 2b–d and f–h; and Extended Data Fig. 11d.) Purple traces show state-space trajectories, squares indicate cue onset, open circles indicate cue offset, and crosses indicate asymptotically stable fixed points (if there are any), colours indicate cue conditions as in Fig. 2d.



Extended Data Fig. 13 | **Full-delay trained networks. a**, Cost function for full-delay training on the random delay task (Methods 1.3.3). Yellow ticks indicate cue onset and offset times, the yellow bar indicates range of go times in the variable delay task. Boxcars show intervals over which stable decoding performance was required in three example trials with different delays (Methods 1.3.3). **b–c**, Same as Fig. 6c–d, but when training with the full-delay cost with a regularisation strength of $\alpha_{nonlin}^{(2)} = 0.0005$ (**b** solid, **c** left) or $\alpha_{nonlin}^{(2)} = 0.0005$ (**b** dashed, **c** right, Methods 1.7.4). **d**, Subspace overlap between different task epochs, measured as the percent variance explained (PVE) by projecting neural activity from one task epoch (tested; cf. Extended Data Fig. 11c, Extended Data Fig. 12d, and Extended Data Fig. 10b) through the top 4 PCs of another task epoch (fitted) for the networks shown in **b–c**. Diagonal elements show the PVE within each task epoch.