Reduced variability of bursting activity during working memory

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Working memories have long been thought to be maintained by persistent spiking. However, mounting evidence from multiple-electrode recording (and single-trial analyses) shows that the underlying spiking is better characterized by intermittent bursts of activity. A counterargument suggested this intermittent activity is at odds with observations that spike-time variability reduces during task performance. However, this counterargument rests on assumptions, such as randomness in the timing of the bursts, which may not be correct. Thus, we analyzed spiking and LFPs from monkeys’ prefrontal cortex (PFC) to determine if task-related reductions in variability can co-exist with intermittent spiking. We found that it does because both spiking and associated gamma bursts were task-modulated, not random. In fact, the task-related reduction in spike variability could largely be explained by a related reduction in gamma burst variability. Our results provide further support for the intermittent activity models of working memory as well as novel mechanistic insights into how spike variability is reduced during cognitive tasks.

Working memory (WM), the holding of information “online” in a form available for processing, is central in human cognition. It has long been thought that WMs are carried by persistent spiking of neurons that maintain “attractor states”¹⁻⁴. But recent evidence suggests that WM-related activity is more dynamic⁵⁻¹¹. Advances in multiple-electrode technology have allowed sampling of larger populations of neurons as well as a closer examination of their activity in real time. This revealed that the spiking activity of many neurons does not seem persistent. Instead, it is organized in intermittent bursts, especially when activity is examined on individual trials⁵,⁶,¹²,¹³. Because burst times vary from trial to trial, averaging across trials can create the illusion of persistent spiking when the underlying activity is intermittent⁶. Further, the intermittent spiking is not just a property of individual neurons. The bursts of spiking are associated with bursts of gamma-band power⁵,⁶. This suggests that the intermittent bursting is coordinated in local networks. Thus, persistence cannot be obtained by pooling local networks of neurons. This provides support to a class of synaptic attractor models in which short-term plasticity helps maintain attractor states in between bouts of intermittent spiking¹⁴⁻¹⁷. This is consistent with observations from EEG and fMRI studies that, for extended periods of time, information held in working memory cannot be decoded from global activity. However, when the cortex is “pinged” by a task-irrelevant stimulus or by transcranial magnetic stimulation, the network “rings” back with the information held in working memory⁷⁻⁹.

On the other hand, observations that cortical spiking becomes less variable during performance of various tasks¹⁸⁻²³ have led to a counterargument to intermittent spiking. For example, a recent modelling study argued against the synaptic attractor type models (and in favor of persistent spiking alone) by suggesting that intermittent bursting should cause an increase, not a decrease, in across-trial variability of spiking (measured by Fano Factor, FF)²⁴. These arguments rest on several assumptions that may not be consistent with experimental observations. Most importantly, the model assumed that intermittent bursts of spiking occurred at random times during a WM task. However, experimental observations show that WM-related bursts of activity are not random, they wax and wane with different trial events⁵,⁶⁴⁷⁸. This raises the question of whether this task-related modulation of bursts is enough to explain the task-related reduction in spike variability that is often observed in cortex.

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To address this, we analyzed actual spiking and LFPs from the prefrontal cortex of non-human primates (NHPs) performing three working memory tasks. We previously reported intermittent bursts of spiking and associated bursts of gamma power during performance of the tasks5,6. Here, we report that variability of spiking (FF) decreased, not increased, during task performance. In fact, the largest task-related reduction in variability was found at recording sites where the spike rates and burst rates increased the most. We also gained insight into why variability was reduced. We synthesized spikes based on the gamma bursts found in the LFPs. This revealed that the reduction in spiking variability was directly related to reductions in gamma burst variability. These results provide further support for synaptic attractor type models of working memory with intermittent activity15–17,25–27.

Results

Increased bursting co-exists with reduced spike variability. We analyzed LFP power and spiking from multiple single neurons recorded from the lateral prefrontal cortex of monkeys performing three different WM tasks. From these data, we previously reported intermittent bursts of spiking and associated bursts of gamma power during performance of the tasks5,6. Here, we report that variability of spiking (FF) decreased, not increased, during task performance. In fact, the largest task-related reduction in variability was found at recording sites where the spike rates and burst rates increased the most. We also gained insight into why variability was reduced. We synthesized spikes based on the gamma bursts found in the LFPs. This revealed that the reduction in spiking variability was directly related to reductions in gamma burst variability. These results provide further support for synaptic attractor type models of working memory with intermittent activity15–17,25–27.
to the corresponding pre-stimulus period, using paired t-test for increase in number of bursts in the pre-stimulus to stimulus presentation time windows at threshold $p = 0.05$, see also “Materials and methods” section and Refs. 5,6. We referred to these sites as “gamma-modulated sites” while the sites without stimulus-induced gamma bursting were referred to as “non-modulated sites”5,6. We then demonstrated that spiking at gamma-modulated sites exclusively carried information about the stimuli held in WM.

Both gamma-modulated and non-modulated sites showed a reduction in gamma-burst variability during the task. Figure 2 shows the difference in spike rates (Fig. 2E) and gamma burst rates (Fig. 2F) between
Figure 3. Spike and burst variability in Task 2. (A) Spike rates averaged across trials and all neurons (n = 283). Gray shaded regions indicate sample presentations (S; the first sample is presented at time = 0). The two delays are marked by (D). Error bars indicate standard error of the mean in all panels. (B) Gamma burst rates averaged across trials for all recording sites. (C) FF in spike times normalized by the pre-stimulus baseline (marked by “B”; from 500 to 100 ms prior to first sample onset) average. (D) FF in burst times normalized by the pre-stimulus (from 500 to 100 ms prior to first sample onset) average. (E–H) Same as (A–D) but separately for sites that increased (blue) gamma bursting at sample onset, and sites that did not (red).

Figure 4. Burst variability in Task 3. Top row shows the gamma burst FF for short (left) and long (right) trials. Shaded areas indicate standard error of mean. Bottom row plots the corresponding gamma burst rates.
gamma-modulated and non-modulated sites. Much like the average across all sites (Fig. 2B), gamma burst rate at gamma-modulated sites increased during stimulus presentation and ramped up over the memory delay (Fig. 2F, blue lines). As expected, there was less gamma bursting and no stimulus dependent modulation at non-modulated sites (Fig. 2F, red lines), and the spike rates were lower at non-modulated sites (Fig. 2E).

Importantly, we found that the spiking FF showed a greater reduction at the gamma-modulated sites (Figs. 2G, 3G, blue lines) than non-modulated sites (Figs. 2G, 3G, red lines). Gamma bursting at both types of sites showed a reduction in FF as the trial progressed relative to the baseline but this effect was more pronounced at the gamma-modulated sites (Figs. 2H, 3H). During sample stimulus presentations, when both gamma burst and spike rates increased at the gamma-modulated sites, there were sharp reductions in the FFs at these sites (Figs. 2E,F, 3E,F). At test stimulus presentations, when the burst rates were at their peak, the reductions in spike and gamma burst FFs were most pronounced (Figs. 4, 5).

These analyses indicate that task modulation of spike activity and gamma bursts does result in task-related reduction in neural variability. The examination of gamma-modulated vs non-modulated sites and different task epochs further indicates that the higher the gamma bursting, the greater the reduction in both spike and gamma burst variability. The gamma bursts are thought to reflect coordinated spiking in local networks. This raises the question of whether the task-modulation of the gamma bursts can, in turn, explain the task-related reduction in spike variability. We address this below focusing on Task 1.

Reductions of gamma burst variability help explain the reduction in spike variability. To further examine the relationship between spiking and gamma burst variability, we used gamma burst events to construct synthetic spike trains. In particular, inside every gamma burst identified (see "Materials and methods"
section and Ref.5) at every recording site on each trial we generated an artificial Poisson spike train (see "Materials and methods" section). We systematically varied the spike rates inside bursts over a wide range and quantified the resulting synthetic spike variability across trials using FF. The analysis revealed that task-related modulations of the gamma bursts reduced the FF for the artificial spikes to a similar degree as it did for actual recorded spikes (Fig. 6). In fact, this effect was observed across a wide range of spike rates within the bursts, although it was more pronounced for the higher spiking rates. The effect gradually decreased as spike rates inside bursts were reduced to 10 sp/s (corresponding to the average of 0.54 sp/burst given the average burst duration of 54 ms in the data. Thus bursting does not necessarily have to be apparent in single neuron activity.). In addition, we added background spiking activity outside bursts (at 1 sp/s). This resulted in a weakening of the stimulus-driven reduction in spike time variability, especially for low within-burst firing rates (Fig. 6B). This suggests that most spiking has to occur during the bursts for the gamma burst variability to drive the observed drop in spike variability.

Figure 6. Variability of synthetic spike trains generated based on neurophysiological gamma burst times in Task 1. Spiking rates within the bursts varied from 10 to 1000 sp/s (black to light gray). (A) Plotted are the mean (averaged across all the recording sites) FFs normalized divisively by the pre-stimulus average, for a range of simulated spiking rates without (A) and with (B) an additional synthetic background activity at the level of 1 sp/s. Red line corresponds to the original normalized spike FF for the recorded spiking. (C) The FF for original data (red) and synthetic data (black). The synthetic data was created by using the recorded burst times and populating each trial with spikes based on firing rates measured inside and outside bursts per neuron ("Materials and methods" section). Blue marks: FFs in two time windows (normalized by the pre-stim time window; both significantly lower than pre-stimulus window using Wilcoxon rank test, p = 0.007 for sample and p < e−30 for test time window) where both burst counts and the corresponding spiking was synthesized from data. First, burst counts were produced based on the distribution of burst counts in the selected time window for each electrode. The synthetic spike trains were then produced as above (see also "Materials and methods" section).
In addition, we estimated the average firing rates inside and outside gamma bursts per neuron to generate synthetic spike trains (based on the measured gamma burst times for that neuron). This yielded FFs similar to the ones measured in the data (Fig. 6C, black vs red lines). This suggests that the timing of bursts and the corresponding modulation of firing rates could explain a large part of the reduction in spike variability. The reduction in the variability of the synthetic data was particularly pronounced at periods of increased bursting, during sample and test stimulus onsets.

Finally, we created synthetic burst times ("Materials and methods" section) in three time windows corresponding to the periods before and during the first sample stimulus as well as the first test stimulus. Based on the distribution of burst counts in each time window, we synthesized burst events in epochs corresponding to the three time windows. We then populated those synthetic trial epochs with spikes based on the estimated average firing rates inside and outside bursts for each recorded unit, similarly as in previous analysis of synthetic spike trains. There was a similar reduction in FF from pre-sample to sample stimulus onset as the original data (Fig. 6C, blue marks). Thus, taken together, the reduction in spiking variability during the task can be largely explained by the reduction of gamma burst variability reflecting coordinated population activity.

Discussion
We found that the task-related modulation of bursts of spiking and gamma power during a WM task resulted in the cross-trial reduction in the variability of neural activity. Further, we found a direct relationship between the reduction of the variability of gamma bursting and the reduction of spiking variability. They co-occurred both in time and space. The reduced spike variability was most pronounced during task events and at recording sites where the reduction in gamma burst FF was large. These were the same sites where gamma bursting and spiking were most strongly task-modulated. However, reductions in activity variability were seen both at recording sites with increased and decreased gamma bursting. Thus, the reduced burst variability was not due to increased bursting per se. This can explain why the reduction in spiking variability is largely decoupled from spike rates24.

It was the increase in the structure of bursting activity via modulation by the task that produced the reduction in variability. We investigated this by synthesizing spike trains from recorded gamma burst patterns and found that the pattern of bursts could explain reduced spike variability for a wide range of synthetic spike rates including the rates observed in the data. Thus the timing of population bursts before and after task onset can explain a large part of the reduced spiking variability long observed in empirical data19–23. It is however important to note that in addition to this, individual neurons have other sources of variability not captured by population bursts, such as stimulus preference or non-Poissonian firing patterns41.

These results provide further support for working memory models in which information is held by a combination of spiking (during bursts) and short-term synaptic plasticity (between bursts) rather than persistent spiking alone15–17,23,27. In these " synaptic attractor" models bursts of spiking induce temporary synaptic imprints that maintain a trace of the attractor state in between spiking. Further, recent evidence has shown that the short-term synaptic plasticity endows additional benefits beyond memory maintenance. For instance, it helps make recurrent networks more stable and more robust to perturbations and synaptic loss55,56.

A recent modeling study argued that intermittent bursting of activity is not compatible with task-related reductions in spiking variability43. However, this model used several problematic assumptions that were not in line with experimental observations. Most critically, the Li et al.24 model assumed intermittent bursting at random times during the task and they assumed no bursting (i.e., steady spiking) before task onset. Neurophysiological recordings paint a different picture. Task-related bursting is not random. It is highly modulated, strongly waxing and waning with different trial events5,6. Here, we demonstrated that this modulation reduces trial-to-trial spiking variability. Neurophysiological recordings, unlike the Li et al.24 model, also show bursting before task onset22. Thus, Li et al.24 made comparisons to a baseline with an artificially low level of variability not actually seen in the data. Therefore, the model assumed that the pattern of bursts could explain reduced spike variability for a wide range of synthetic spike rates including the rates observed in the data. Thus the timing of population bursts before and after task onset can explain the large part of the reduced spiking variability long observed in empirical data19–23. It is however important to note that in addition to this, individual neurons have other sources of variability not captured by population bursts, such as stimulus preference or non-Poissonian firing patterns41.

These oscillations occur at a similar time scale as the spike bursts and are related48. Here we demonstrate that this framework is consistent with task-related reduction in neuron variability that is common across cortex.
Materials and methods

Data. We analyzed data from prior studies\(^5\)\(^,\)\(^6\)\(^,\)\(^28\)\(^,\)\(^29\)\. This data consisted of three data sets with electrode recordings from PFC while monkeys performed two distinct WM tasks (Task 1, Task2, Task 3: Fig. 1). Task 1 corresponds to 3-item trials in Lundqvist et al\(^5\). We focused on 3-item trials as they allowed us to have a larger number of repeated trials. Task 2 corresponds to the recognition task in Refs.\(^28\)\(^,\)\(^29\), which is the same data analyzed in Ref.\(^6\). Task 3 corresponds to the delayed saccade task in Lundqvist et al\(^5\). Task 1 and 2 were recorded with acute electrodes in PFC, yielding both LFPs and isolatable neurons. For Task 3 we had local field-potential data from chronic Utah arrays with no isolated neurons. All the animals received postoperative antibiotics and analgesics and were always handled in accord with the National Institutes of Health guidelines, and all procedures were approved by the Massachusetts Institute of Technology Committee on Animal Care. They were trained with positive reward (juice) only and maintained in accordance with the National Institutes of Health guidelines and the policies of the Massachusetts Institute of Technology Committee for Animal Care). The study is reported in accordance with ARRIVE guidelines.

The LFPs were recorded at 30 kHz in Task 1 and Task 3, and 1 kHz in Task 2. From each task we recorded from two animals (one male Macaca mulatta and one male Macaca fascicularis in Task 1, one male and one female Macaca mulatta in Task 2, two male Macaca mulatta in Task 3). Trial conditions were always randomized between trials. Data was analyzed using automatic scripts, and researchers were therefore blind to group belongings of single trials. The data from Task 1 and Task 2 were down sampled to 1 kHz before further analysis. We kept all electrode contacts with at least one associated isolated neuron in the acute data, all electrodes from the array data. As a result, we obtained 495 units corresponding to 319 electrodes in Task 1 and 233 units on 199 electrodes in Task 2. For Task 3 we had 8 sessions with 64 electrodes in each. Only correct trials were kept for further analysis (on average, 255 + -72 trials in Task 1 and 332 + -80 trials in Task 2).

Gamma modulated sites were chosen based on activity during sample presentations. To this end, we identified sites with the average gamma burst rate over any of the sample presentations significantly exceeding the threshold of 0.05. The rest of sites were considered as non-modulated.

Burst analysis. We relied on gamma burst times extracted in the two prior studies\(^5\)\(^,\)\(^6\)\. In short, we first extracted the gamma-band power in 3 distinct bands (40–65, 55–90 and 70–100 Hz). For this purpose we adopted a multi-taper approach with frequency-dependent window lengths corresponding to 6–8 oscillatory cycles and frequency smoothing corresponding to 0.2–0.3 of the central freq, \(f_0\), i.e. \(f = f_0 ± 0.2f_0\), where \(f_0\) were sampled with the resolution of 1 Hz (this configuration implies that 2–3 tapers were used). We then extracted an estimate of the band power envelope by averaging spectral components within each band with the temporal resolution of 1 ms. Next, we thresholded the envelopes based on their mean and standard deviation during fixation in pre-stimulus conditions (using the last 300 ms prior to stimulus onset in Task 1, last 500 ms prior to stimulus onset in Task 2).

A gamma burst was defined as a time interval when the envelope exceeded the mean by two standard deviations for at least the duration of 3 oscillatory cycles (using the center of each gamma band to define the length of an oscillatory cycle). In line with our previous work\(^5\)\(^,\)\(^6\) we also used a trial-average measure of burst rates for each spectral band. It corresponds to the chance of a burst occurrence on an individual electrode at a particular time in the trial. In this work, we used the union of bursts across the three gamma bands and, accordingly, we relied on the average burst rate across those identified for the three gamma bands.

Fano factor (FF). We estimated the spiking Fano factor (FF) for each unit in a two-step process. First, we calculated the number of spikes, \(s\), within a sliding window of 150 ms and calculated its mean, \(\mu\), and variance, \(\sigma^2\), across trials to estimate \(FF = \sigma^2 / \mu\). This resulted in a raw FF time series of 1 ms resolution. Further, to facilitate comparisons we show in Figs. 2 and 3 the relative FF, which is obtained by removing the mean FF over the pre-stimulus baseline from the raw FF.

The FF for bursts was estimated using a binary time series for each site and trial with 0 corresponding to the time bin outside a burst event and 1—inside a burst event. The resulting time series was then subject to the same treatment as spike time series.

Sythetic spike train generation. We constructed synthetic spike trains based on the recorded gamma burst timings. In particular, we randomly placed spikes inside and outside burst intervals for each corresponding trial and channel to match the desirable target average firing rates inside and outside bursts. For the results shown in Fig. 6A, the average firing rates inside bursts were systematically varied from 10 to 1000 sp/s (with no spikes outside burst intervals). In another manipulation (Fig. 6B) we added background spike activity with the constant rate of 1 sp/s in each trial. Finally, following our approach for gamma burst dependent random spike generation, we produced synthetic spike trains with the target firing rates inside and outside bursts corresponding to the average firing rates in the recorded data inside and outside bursts, respectively (Fig. 6C). Importantly, these averages were estimated (and then used for spike generation) independently for each channel, i.e. burst events for each electrode were matched with spiking units measured on that electrode. In addition, in three 150 ms windows (~ 250 to 100 ms before stimulus onset, 75 to 225 ms after stimulus onset, 75 to 225 ms after test onset) we used the recorded burst count distributions to create synthetic burst events. We then populated these synthetic bursts with synthetic spikes and calculated corresponding FFs.

Data availability
The datasets used in the current study are available from the corresponding author on reasonable request.
References

1. Fuster, J. M. & Alexander, G. E. Neuron activity related to short-term memory. Science 173, 652–654 (1971).
2. Funahashi, S., Bruce, C. I. & Goldman-Rakic, P. S. Mnemonic coding of visual space in the monkey's dorsolateral prefrontal cortex. J. Neurophysiol. 51, 331–349 (1989).
3. Goldman-Rakic, P. S. Cellular basis of working memory. Neuron 14(3), 477–485 (1995).
4. Amit, D. J. & Brunel, N. Model of global spontaneous activity and local structured activity during delay periods in the cerebral cortex. Cereb. Cortex (New York) 7(3), 237–252 (1997).
5. Lundqvist, M. et al. Gamma and beta bursts underlie working memory. Neuron 90(1), 152–164 (2016).
6. Lundqvist, M., Herman, P., Warden, M. R., Brincat, S. L. & Miller, E. K. Gamma and beta bursts during working memory readout suggest roles in its volitional control. Nat. Commun. 9(1), 1–12 (2018).
7. Wolff, M. J., Jochim, J., Akyurek, E. G. & Stokes, M. G. Dynamic hidden states underlying working-memory-guided behavior. Nat. Neurosci. 20(6), 864–871 (2017).
8. Rose, N. S. et al. Reactivation of latent working memories with transcranial magnetic stimulation. Science 354(6316), 1136–1139 (2016).
9. Sprague, T. C., Ester, E. F. & Serences, J. T. Restoring latent visual working memory representations in human cortex. Neuron 91(3), 694–707 (2016).
10. Hussar, C. R. & Pasternak, T. Memory-guided sensory comparisons in the prefrontal cortex: Contribution of putative pyramidal cells and interneurons. J. Neurosci. 32(36), 2747–2761 (2012).
11. Shafi, M. et al. Variability in neuronal activity in primate cortex during working memory tasks. Neuroscience 146(3), 1082–1108 (2007).
12. Lundqvist, M., Bastos, A. M. & Miller, E. K. Preservation and changes in oscillatory dynamics across the cortical hierarchy. J. Cogn. Neurosci. 32(10), 2024–2035 (2020).
13. Bastos, A. M., Loonis, R., Kornblith, S., Lundqvist, M. & Miller, E. K. Laminar recordings in frontal cortex suggest distinct layers for maintenance and control of working memory. Proc. Natl. Acad. Sci. 115(5), 1117–1122 (2018).
14. Wang, Y. et al. Heterogeneity in the pyramidal network of the medial prefrontal cortex. Nat. Neurosci. 9(4), 534–542 (2006).
15. Lundqvist, M., Herman, P. & Lansner, A. Theta and gamma power increases and alpha/beta power decreases with memory load in an attractor network model. J. Cogn. Neurosci. 23(10), 3008–3020 (2011).
16. Mongillo, G., Barak, O. & Tsodyks, M. Synaptic theory of working memory. Science 319(5869), 1543–1546 (2008).
17. Fiebig, F. & Lansner, A. A spiking working memory model based on Hebbian short-term potentiation. J. Neurosci. 37(1), 83–96 (2017).
18. Mitchell, J. F., Sundberg, K. A. & Reynolds, J. H. Differential attention-dependent response modulation across cell classes in macaque visual area V4. Neuron 55(1), 131–141 (2007).
19. Churchland, M. M. et al. Stimulus onset quenches neural variability: A widespread cortical phenomenon. Nat. Neurosci. 13(3), 369–378 (2010).
20. Cohen, M. R. & Maunsell, J. H. Attention improves performance primarily by reducing interneuronal correlations. Nat. Neurosci. 12(12), 1594–1600 (2009).
21. Ponce-Alvarez, A., Thiele, A., Albright, T. D., Stoner, G. R. & Deco, G. Stimulus-dependent variability and noise correlations in cortical MT neurons. Proc. Natl. Acad. Sci. 110(32), 13162–13167 (2013).
22. Hussar, C. & Pasternak, T. Trial-to-trial variability of the prefrontal neurons reveals the nature of their engagement in a motion discrimination task. Proc. Natl. Acad. Sci. 107(50), 21842–21847 (2010).
23. Ito, T. et al. Task-evoked activity quenches neural correlations and variability across cortical areas. PLoS Comput. Biol. 16(8), e1007983 (2020).
24. Li, D., Constantinidis, C. & Murray, J. D. Trial-to-Trial Variability of Spiking Delay Activity in Prefrontal Cortex Constrains Burst-Coding Models of Working Memory (2021).
25. Sandberg, A., Tegnér, J. & Lansner, A. A working memory model based on fast Hebbian learning. Netw. Comput. Neural Syst. 14(4), 789 (2003).
26. Stokes, M. G. Activity-silent working memory in prefrontal cortex: A dynamic coding framework. Trends Cogn. Sci. 19(7), 394–405 (2015).
27. Kozachkov, L. et al. Robust working memory through short-term synaptic plasticity. BioRxiv 7, 237 (2022).
28. Warden, M. R. & Miller, E. K. Task-dependent changes in short-term memory in the prefrontal cortex. J. Neurosci. 30(47), 15801–15810 (2010).
29. Warden, M. R. & Miller, E. K. The representation of multiple objects in prefrontal neuronal delay activity. Cereb. Cortex 17, 141–150 (2007).
30. Wu, Y. K. & Zenke, F. Nonlinear transient amplification in recurrent neural networks with short-term plasticity. Elife 10, e67263 (2021).
31. Herrero, J. I., Gieselmann, M. A., Sanaye, M. & Thiele, A. Attention-induced variance and noise correlation reduction in macaque V1 is mediated by NMDA receptors. Neuron 78(4), 729–739 (2013).
32. Lakatos, P., Karmos, G., Mehta, A. D., Ulbert, I. & Schroeder, C. E. Entrainment of neuronal oscillations as a mechanism of attentional selection. Science 320(5872), 110–113 (2008).
33. Buschmann, T. J. & Miller, E. K. Shifting the spotlight of attention: Evidence for discrete computations in cognition. Front. Hum. Neurosci. 4(194), 1–9 (2010).
34. Schroeder, C. E., Wilson, D. A., Radman, T., Scharfman, H. & Lakatos, P. Dynamics of active sensing and perceptual selection. Curr. Opin. Neurobiol. 20(2), 172–176 (2010).
35. Engel, T. A. et al. Selective modulation of cortical state during spatial attention. Science 354(6316), 1140–1144. https://doi.org/10.1126/science.aag1420 (2016).
36. Luczak, A., McNaughton, B. L. & Harris, K. D. Packet-based communication in the cortex. Nat. Rev. Neurosci. 16(12), 745–755 (2015).
37. Heifrech, R. E. et al. Neural mechanisms of sustained attention are rhythmic. Neuron 99(4), 854–865 (2018).
38. VanBullen, R. Attention cycles. Neuron 99(4), 632–634 (2018).
39. Stringer, C. et al. Spontaneous behaviors drive multidimensional, brainwide activity. Science. https://doi.org/10.1126/science.aav793 (2019).
40. Lundqvist, M. & Wutz, A. New methods for oscillation analyses push new theories of discrete cognition. Psychophysiology 59, e13827 (2022).
41. Lisman, J. E. & Idiart, M. A. Storage of 7±2 short-term memories in oscillatory subcycles. Science 267(5203), 1512–1515 (1995).
42. Womelsdorf, T. & Fries, P. Neuronal coherence during selective attentional processing and sensory–motor integration. J. Physiol. Paris 100(4), 182–193 (2006).
43. Buffalo, E. A., Fries, P., Landman, R., Buschman, T. J. & Desimone, R. Laminar differences in gamma and alpha coherence in the ventral stream. *Proc. Natl. Acad. Sci.* **108**(27), 11262–11267 (2011).

44. Bastos, A. M., Vezoli, J. & Fries, P. Communication through coherence with inter-areal delays. *Curr. Opin. Neurobiol.* **31**, 173–180 (2015).

45. Landau, A. N., Schreyer, H. M., Van Pelt, S. & Fries, P. Distributed attention is implemented through theta-rhythmic gamma modulation. *Curr. Biol.* **25**(17), 2332–2337 (2015).

46. Cavanagh, J. F., Cohen, M. X. & Allen, J. J. Prelude to and resolution of an error: EEG phase synchrony reveals cognitive control dynamics during action monitoring. *J. Neurosci.* **29**(1), 98–105 (2009).

47. Sauseng, P. et al. Theta coupling in the human electroencephalogram during a working memory task. *Neurosci. Lett.* **354**(2), 123–126 (2004).

48. Canolty, R. T. et al. High gamma power is phase-locked to theta oscillations in human neocortex. *Science* **313**(5793), 1626–1628 (2006).

49. Maris, E. & Oostenveld, R. Nonparametric statistical testing of EEG-and MEG-data. *J. Neurosci. Methods* **164**(1), 177–190 (2007).

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**Author contributions**

M.L., P.H., and E.K.M. conceived of and wrote the manuscript. M.L. and P.H., conducted data analysis and prepared the figures. J.R., M.R.W., T.J.B. and M.L., collected the data. All authors reviewed the manuscript.

**Competing interests**

The authors declare no competing interests.

**Additional information**

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