

Review

The Distributed Nature of Working Memory

Thomas B. Christophel, 1,2,3,12,* P. Christiaan Klink, 4,5,6,12 Bernhard Spitzer,⁷ Pieter R. Roelfsema,^{5,6,8,12} and John-Dylan Haynes^{1,2,3,9,10,11,12}

Studies in humans and non-human primates have provided evidence for storage of working memory contents in multiple regions ranging from sensory to parietal and prefrontal cortex. We discuss potential explanations for these distributed representations: (i) features in sensory regions versus prefrontal cortex differ in the level of abstractness and generalizability; and (ii) features in prefrontal cortex reflect representations that are transformed for guidance of upcoming behavioral actions. We propose that the propensity to produce persistent activity is a general feature of cortical networks. Future studies may have to shift focus from asking where working memory can be observed in the brain to how a range of specialized brain areas together transform sensory information into a delayed behavioral response.

Storage of Working Memory Contents

Cognition critically depends on the ability to memorize information and change it adaptively. This ability is typically ascribed to working memory, which has often been conceptualized as comprising two basic functions: (i) short-term storage of information and (ii) executive processes that retain no information themselves but control what is retained [1,2]. Here, we argue that instead of being discrete, these functions could be considered extreme functional specializations of a distributed working memory network combining storage and control. Different brain regions might contribute to working memory in accordance with the general nature of their representations. Thus, we consider working memory to rely on the interplay between brain regions that retain low-level or more abstract sensory information, and the transformation of this information for guidance of memory-guided behavioral plans.

Traditionally, working memory research in neuroscience has focused on the question of where contents are stored across delay periods. However, this is subject to ongoing debates [3-8]. Current views emphasize the representation of memorized contents in the prefrontal [3,8–10], parietal [7], or sensory [5,11–13] cortices. Some views more clearly consider the representation of memorized contents at multiple levels of the cortical hierarchy [2,4,14-17]. Localizing working memory storage has proven difficult. The first neural models of working memory were primarily shaped by lesion studies in humans and non-human primates [18-20]. For example, a seminal study [18] performed lesions in prefrontal cortex (PFC) of monkeys and found deficits in their ability to maintain task-relevant information across brief delays, while performance remained intact for tasks that did not require memorization. This and similar work [21-25] led to the belief that working memory is primarily a function of prefrontal cortex.

Later, electrophysiological [26] and functional magnetic resonance imaging (fMRI) studies (reviewed in [27]) revealed persistent neural activity in various brain regions during memory delays, including sensory cortices [28]. However, if a brain region X (say PFC) exhibits delay

Trends

There is no evidence for a single site of working memory storage. Rather, persistent neuronal activity that is informative about a currently memorized stimulus can be found in sensory, parietal, and prefrontal brain regions.

Working memory entails a gradient of abstraction from sensory areas reflecting low-level sensory features to prefrontal regions encoding more abstract, semantic, and responserelated aspects of stimuli.

We hypothesize that all regions of neocortex have the capability to briefly retain their specialized representations in the service of upcoming task demands. Persistent activity in most. if not all, cortical regions can exert control over future behavior.

The contributions of individual brain regions to working memory are best understood as different representational stages with various levels of transformation and abstraction.

¹Bernstein Center for Computational Neuroscience, Charité Universitätsmedizin, Berlin, Germany ²Berlin Center for Advanced Neuroimaging, Charité Universitätsmedizin, Berlin, Germany ³Clinic for Neurology, Charité Universitätsmedizin, Berlin, Germany ⁴Department of Neuromodulation & Behaviour, Netherlands Institute for Neuroscience, Amsterdam, the Netherlands ⁵Department of Vision & Cognition, Netherlands Institute for Neuroscience,



Amsterdam, the Netherlands

Medical Center, University of

⁶Department of Psychiatry, Academic



period activity during working memory delays, and if lesions to X decrease memory performance, it does not necessarily mean that X encodes the working memory content during that delay. Instead, region X might just have an influence on storage taking place in other regions. Any neural signal that encodes working memory content should exhibit at least the following two properties. First, it should contain information about what is memorized, meaning that different memory contents lead to different patterns of activity (stimulus selectivity). Second, stimulusselectivity activity should be present over extended delays in the absence of the stimulus (persistent activity or delay-period activity). The current review focuses on observations of such persistent stimulus-selective activity (see Glossary) in single-cell and local field potential recordings in non-human primates and in human neuroimaging [electroencephalography (EEG) and fMRI]. Please note that delay activity in both sensory and prefrontal regions occasionally fails to extend throughout the maintenance period [29], or can be absent altogether [30,31]. For this reason, working memory mechanisms that do not rely on persistent activity (activity-silent working memory [6,32-34]), that postulate encoding in dynamic firing trajectories [35,36] or that involve gating of spiking activity by local field potentials [37] have also been postulated (see Outstanding Questions). Importantly, persistent activity does not imply that a single selective neuron is firing throughout the entire delay period. Instead, it means that the activity of a neural population encodes stimulus-specific information at any point during the delay. Furthermore, persistent stimulus-selective activity has been proposed to require recurrent excitation [38,39], changes in synaptic facilitation or combinations of both [32,34,40].

Electrophysiological Recordings in Non-Human Primates

First reports of persistent working memory signals date back to the 1970s and describe studies on non-human primates performing delayed response tasks. Most of these early studies focused on the PFC because they were inspired by previous findings that lesions in the PFC cause deficits in delayed response tasks [18]. In one study [41], monkeys were presented with two lamps, one in the left and one in the right visual hemifield. At the beginning of a trial, one of the lamps was briefly turned on. After a delay of a few seconds, the monkeys were trained to press a key on the left or right, corresponding to the memorized side of the light. Activity in a subset of cells in the dorsolateral prefrontal cortex (dIPFC) was modulated during the delay period. Some cells discriminated between left and right stimuli, thus potentially providing evidence for contentspecific delay-period activity in the dIPFC. However, in this study the position of the lights was correlated with the location of the response keys so that the selective delay-period activity might as well have represented motor preparation rather than stimulus memory. The identification of which feature is being represented is a general problem with memory-related representations that is elaborated upon below.

Subsequent studies revealed cells in the PFC that encode memorized positions during the delay period, even when controlling for response preparation [42]. Content-specific delay-period activity in the PFC has been shown for many sensory features (Figure 1, left): objects and natural images [43-45], color [46], and visual motion [29]; also for more abstract features such as numerosity [47] and for other modalities such as the frequency of tactile vibrations [48]. Detailed analyses of visual motion signals in the PFC during working memory has furthermore revealed that they exhibit a tuning profile that is comparable to those found in sensory regions [29,49].

While working memory deficits after lesions in the PFC led most early electrophysiological studies to focus on the PFC, further recordings in non-human primates have also revealed persistent stimulus-selective activity for spatial location in V1 [50,51], for motion in MT [52] and MST [49], for shapes and real-life stimuli in V4 and the temporal cortex [53-55], and for color in the inferior temporal cortex [56,57] (Figure 1, left). Persistent stimulus-selective activity has also been observed in the sensory cortices of other modalities, such as for pure tones in the auditory cortex [58], for haptic texture in the primary somatosensory cortex [59] and, to some extent, for

Netherlands ⁷Department of Experimental Psychology, University of Oxford, Oxford, UK ⁸Department of Integrative Neurophysiology, Centre for Neurogenomics and Cognitive Research, VU University, Amsterdam, The Netherlands ⁹Berlin School of Mind and Brain. Humboldt Universität, Berlin, Germany ¹⁰Cluster of Excellence NeuroCure,

Amsterdam, Amsterdam, The

Charité Universitätsmedizin, Berlin, Germany ¹¹Department of Psychology, Humboldt Universität zu Berlin, Berlin,

Germany ¹²These authors contributed equally to

this work.

*Correspondence: tbchristophel@gmail.com (T.B. Christophel,).



vibration in the secondary somatosensory cortex [60]. An overview of evidence for persistent stimulus-selective activity in primate electrophysiology is shown in Figure 1 (left; see Table S1 in the supplemental information online).

Apart from the PFC and sensory cortices, persistent stimulus-selective activity has furthermore been reported in the primate frontal eye fields (FEFs) [61], parietal cortex [62], premotor cortex [60,63] and the medial temporal lobe [64]. Activity preceding delayed responses has also been reported in subcortical areas such as the mediodorsal nucleus of the thalamus (see [65]), superior colliculi [66], basal ganglia [67], and spinal cord [68] (not included in Figure 1).

Multivariate Decoding in Humans

In the human brain, it can be more challenging to identify content-specific delay-period activity due to the limited spatial resolution of noninvasive neuroimaging techniques. Early neuroimaging studies did not distinguish between content-selective and non-selective delay-period signals [27]. Others reported category-specific differences, for example, between spatial locations and objects [69] or houses and faces [70]. However, with the development of multivariate decoding (also known as multivoxel pattern analysis; MVPA [71-74]), content-specific activity also became accessible to human neuroimaging studies. This was further advanced by the development of inverse encoding models [75-78]. Such studies have identified delay-period information about sensory features in a range of sensory cortical regions (Figure 1, right). Low-level visual features such as orientation, color, motion, or complex patterns can be decoded from early visual areas [79–83]. Auditory stimuli can be read out from the primary auditory cortex [84,85]. Delay-period information about complex visual pattern stimuli has also been found in parietal areas [81,86]. More recent human imaging studies eventually found content-specific delay activity also in frontal areas [77,85,87–92], which was not observed in early MVPA studies [81,82]. Retinotopically organized area FEF, for instance, carries information about memorized spatial position in several tasks [87]. Also, complex shapes [92] and oriented gratings [77] can be decoded from human FEF (often also referred to as superior precentral sulcus). Furthermore, delay-period representations of natural objects [88], Chinese script [93], auditory pure tones [85], and to a limited extend, oriented gratings [77] can be decoded from signals in the lateral PFC. Additional evidence for content-specific delay activity in human PFC comes from EEG studies [89-91] that showed tactile stimulus frequency representations in prefrontal electrode locations.

Taken together, findings in non-human primates and humans reveal that the content of working memory is widely distributed and can be found across sensory, parietal, temporal, and prefrontal cortices (see Figure 1 and Table S1 in the supplemental information online, for an overview; see also [77]).

Why Are Working Memory Representations Distributed?

The presence of working memory signals in both low-level and high-level cortices suggests some kind of distributed representation (Box 1). There are several accounts for this finding (Figure 2). For example, it could in theory reflect a full duplication of sensory information in the PFC (Figure 2A; [8]). While the redundancy of such duplicate representations might, at first sight, stand in contradiction to evolutionary frugality [4,5], it could be useful in establishing the robustness of working memory contents against distraction (see below and [43,94-97]). However, there also appear to be substantial differences in the functional roles of persistent stimulus-selective activity between low-level and high-level brain regions. In the following, we discuss two of these differences (see also Box 2): (i) persistent stimulus-selective activity reflects different levels of abstraction with early sensory regions encoding simple features of stimuli in sensory representations and prefrontal regions encoding memories using more abstract representations or a verbal format (Figure 2B); and (ii) persistent stimulus-selective activity reflects different functional roles of these representations across multiple cortical regions,

Glossary

Abstract representation: high-level representation of memorized stimuli that is less detailed with regards to specific physical stimulus features, but that allows for generalization across a large number of stimuli that share the same semantic property or that belong to the same category.

Activity-silent working memory: neural mechanisms of working memory that are not directly detectable as changes in spike rate (or BOLD activity), for example: temporary changes in synaptic efficacy.

Distributed representations: neural representations of a memorized content (i.e., stimulus information) that can be found in multiple neural populations (see Box 4).

Dynamic firing trajectories: encoding of working memory contents in a dynamic temporal pattern of spiking activity.

Persistent stimulus-selective activity: any neural activity that takes place after a sensory stimulus is no longer present and that holds information about that stimulus. Sensory representation: low-level representation of memorized stimuli that carries detailed information about the originally presented stimulus. Turing machine: hypothetical machine, invented by mathematician Alan Turing as a thought experiment

on computational operations (see



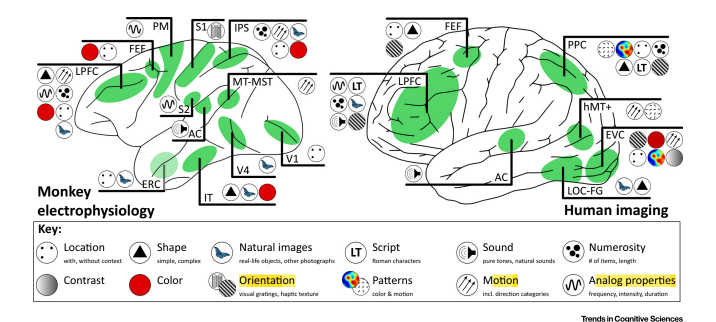


Figure 1. Overview of Content-Specific Activity during Working Memory Delays in the Macaque (Left) and Human (Right) Brain. Icons indicate persistent stimulus-selective activity for each stimulus type indicated by the icon (see legend) at the respective locations. Brain areas are identified by abbreviations (a full list of individual studies is reported in Table S1 in the supplemental information online). Both left- and right-sided effects are shown on the left hemisphere. Data from delayperiods during which subjects can prepare a specific motor output instead of memorizing a stimulus (e.g., delayed response tasks, e.g., [137]) and tasks that involve any form of explicit mental transformation [192,193] or explicitly learned association [132] are excluded. AC, auditory cortex; ERC, enthorinal cortex; EVC, early visual cortex; FEF, frontal eye fields; FG, fusiform gyrus; hMT+, human analog to MT/MST; IPS, intraparietal sulcus; IT, inferior temporal cortex; LOC, lateral occipital complex; IPFC, lateral prefrontal cortex; PM, premotor cortex; PPC, posterior parietal cortex.

ranging from representing the incoming low-level sensory features to planning a memorydependent behavioral response (Figure 2C).

Different Levels of Abstraction

Imagine a task that involves memorizing one out of two visually presented animals (say a yellow lion and a grey elephant, see Figure 2). It is possible to memorize the distinction between these animals at different levels of abstraction, for example, using low-level visual features such as outlines or colors, or abstract categories like type-of-animal. Importantly, the memory representations can go beyond the features presented in the samples. In our example, it would be possible to memorize images of animals based on their typical sounds, their verbal labels, or even the emotions they evoke. In early work many authors believed working memory to be of a purely verbal nature [98-102], but we now know that working memories can also have basic sensory qualities [103-106]. Working memory signals in sensory and prefrontal cortices could reflect such different levels of abstraction. This specialization between sensory and prefrontal regions for different levels of abstraction is a general feature of cortical processing [2,107]. In the field of working memory, it is supported by several lines of evidence.

An early route to distinguishing sensory from abstract memory representations has been to require participants to memorize nonsense patterns or shapes that are hard to verbalize. Behavioral studies using this strategy have shown that working memory can rely exclusively on visual features [103]. Content-specific delay-period activity for such nonsense stimuli has been found in ventral occipitotemporal cortex, dorsal occipitoparietal areas, and the frontal eye fields, but not in PFC [53,81,86,92]. Interpreting these findings, however, relies on interpreting null results, which can be difficult for imaging data in particular.



Box 1. Distributed Representations

The term distributed representation is ambiguous and requires clarification. Several distinct interpretations can be identified

- (1) Local pattern information: it has been repeatedly shown that the orientation of a grating stimulus during a working memory delay can be decoded from patterns of brain activity in primary visual cortex [79]. In this case a distributed representation means that the orientation information is disseminated across a local population of units within a single
- (2) Separable information in multiple brain regions: orientation information can also be decoded from signals in the IPFC, independent of the decodable information in the primary visual cortex [77]. Distributed here means that there are multiple local response patterns in parallel; each of which can be independently used to decode orientation. In this review, this is what is meant by the term distributed representation.
- (3) Inseparable information across multiple brain regions: finally, information might be encoded in global patterns of brain activity that is not encoded at the level of individual regions. In this case, a single area would not allow for decoding of such information, but considering the signals across multiple regions together would allow the extraction of information.Distinguishing between Cases 2 and 3 experimentally requires an assessment of whether the different regions contain sufficient information on their own to decode a stimulus feature, or whether only the combination of signals across areas allows decoding. Thus, an important criterion is whether the information in the multiple regions is redundant. The distinction between Case 2 and 3 is probably gradual rather than categorical. Memory codes could, for example, be redundant or synergistic so that the information present in two regions is less or more than the sum, respectively.

If multiple, redundant, working memory representations indeed exist, it raises the interesting question which of them will be used to solve any given task. Lesion studies are one way to answer this question, as partially evidenced from the fact that many working memory tasks can be performed even after PFC lesions [20,156]. Another approach could be to investigate the choice probabilities [157] associated with activity in particular areas to analyze which signals are maximally predictive of behavioral choices in a memory task.

A different approach to dissociate levels of abstraction is to use stimuli with both low-level and abstract features and to vary the degree to which the task emphasizes sensory detail. One recent neuroimaging study [88] capitalized on this strategy using real-life visual objects. In one (visual) condition, participants were shown an object fragment and asked whether this fragment belonged to a memorized item or not. In another (nonvisual) condition, they had to report whether a new object (e.g., a watch) belonged to the same category as the memorized item (e. g., a clock). Sensory areas exhibited delay-period information about the stimulus during the visual task but not during the nonvisual task. Prefrontal areas showed the opposite pattern. Similarly, the PFC was found to represent color information when individuals were asked to discern clearly categorizable colors, but not when discriminating subtle color hues [31,46]. This distinction between memory representations for fine-grained color hues and color category is further supported by recent behavioral experiments and modeling work [108].

Another way of testing whether a brain region encodes abstract information is to assess to which degree neuronal memory representations generalize across different stimuli. In one series of studies [109,110], for instance, macaques were trained to memorize morphed images of animals whose low-level visual features could vary either within category or across a category boundary between cats and dogs. Prefrontal neurons generalized more across different exemplars of the same category than neurons in the inferior temporal cortex. Likewise, converging evidence from human [91] and non-human primate electrophysiology [63] indicates that neurons in frontal areas that were initially demonstrated to represent memorized tactile vibration frequencies [48], can in fact represent parametric memories from a range of sensory modalities (visual, auditory, and tactile). Frontal areas thus appear to encode these memories in an abstract, amodal magnitude format [111], while parametric delay activity in modality-specific sensory cortices is mostly absent [89,91,112]. Prefrontal areas have also been shown to host memory representations of abstract number information [47,113] in formats that may generalize across sensory modalities [114,115].

Taken together, these findings are compatible with a division of labor, in which sensory regions encode low-level details and prefrontal regions encode abstract, categorical information that



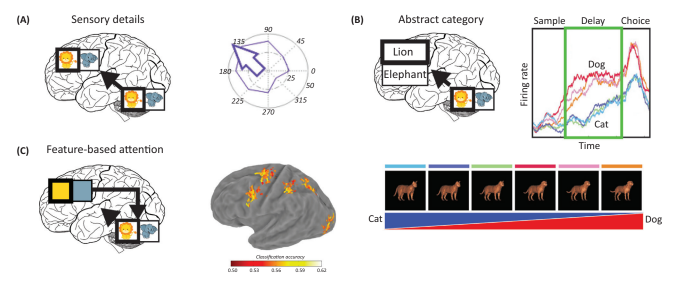


Figure 2. Potential Explanations for Persistent Stimulus-Selective Activity in Prefrontal Cortex during Working Memory Delays, Shown Here for the Example of Memorizing a Lion (versus an Elephant). (A) The prefrontal cortex maintains sensory details with similar resolution to sensory brain regions [29,77]. The right shows a plot of responses of a cell in monkey lateral prefrontal cortex to different directions of motion, which exhibits similar properties to responses in MT [194]. (B) Representation of abstract category information. The right panel shows responses to six morphs gradually varying between pure cat and pure dog images. This cell largely encodes the perceived category with the three morphs that are perceptually classified as dogs yielding a similar and high categorical response [109,195]. (C) Prefrontal cortex prospectively encodes feature-based attention during the delay that prepares for an upcoming search for the target. This can serve as a potential bias signal for feature-selective representations in early visual cortex [196].

generalizes across modalities [5,116,117]. A computational advantage of storing low-level features in sensory areas is that it circumvents the necessity to duplicate low-level feature spaces for persistent storage in higher brain regions (Figure 2A, [5,116]). Persistent stimulusselective activity in the PFC can then be used to build flexible and task-dependent representations, such as arbitrary categorical labels (Figure 2B; [116]). However, some studies have suggested that also low-level features can be represented in the PFC [46,77], which appears to be at odds with a distinction based purely on the level of abstraction. It has been proposed that such representations of low-level sensory features in the PFC might reflect memorization using a nonsensory neural code [31] (see also Box 2). For example, orientation information can be memorized using symbolic, numerical cues and this does not necessarily involve persistent stimulus-selective in sensory cortices (see Supplementary Material in [79]). Alternatively, as we outline next, the prevalence of sensory information in the PFC might reflect the transformation of sensory information in a form that serves to prepare for an upcoming behavioral response.

Transformation from Sensory Input to a Behavioral Response

A central function of working memory is to bridge the gap between a stimulus and a response that is to be executed while the stimulus is no longer present. The PFC seems to encode the tobe-memorized information in a format that can directly guide behavior ([5,14,15,118,119]). Persistent activity thus not only reflects a retrospective memory of the stimulus but also a prospective action plan. Such entangled tuning for different cognitive variables has been observed repeatedly in the PFC [120] and has been proposed as a general model of PFC function [121].

The transformation of sensory signals into an intermediate format that bridges between stimuli and delayed target responses can be easily appreciated. Consider a task that involves memorizing a picture of a lion and selecting this animal from one of two alternative targets after a delay. The incoming target stimulus to which a memorized sample must be matched will drive the



Box 2. A Hierarchy of Spatial Scale

Spatial maps can be found in visual, parietal, and frontal cortices alike [158-160]. This reflects the predominance of space as an organizational principle across many brain areas. The encoding of memorized positions has also been observed across many stages of the cortical hierarchy including in the IPFC [31,61,62,76,87]. As for nonspatial features, there is a gradient of representation across the hierarchy: (i) higher-level areas exhibit larger spatial receptive fields [161-163]; (ii) spatial mnemonic representations in parietal and frontal cortices are closely tied to the preparation of actions and the control of attention shifts [164-166]; and (iii) spatial representations in higher-level regions generalize across many different sensory modalities [167-169] and involve nonretinal frames of reference [135,170]. In contrast, lower-level maps are recruited when tasks demand memory of fine spatial detail in retinal coordinates [50].

Mnemonic maps of visual space could also be involved in the representation of other feature information. For example, real-life objects are not spatially homogeneous and are known to elicit differential distributions of attention and saliency across space, even for objects of similar size [171]. The frontal eye fields and parietal cortex encode the spatial distribution of saliency within natural images [172,173]. Thus, information that appears to be specific to the nonspatial features of an object might also depend on spatial memory of particularly salient subregions within the shape of the object.

Stimulus-selective information in the PFC has been shown for low-level nonspatial features such as grating orientation [77] and motion direction [29,49]. Recent behavioral and electrophysiological work suggests that spatial strategies might play a role in memorizing such features [174]. Consistent with this possibility, orientation-selective responses in V1 correlate with the topography of spatial selectivity, demonstrating joint selectivity for orientation and space [175]. In line with this, selectivity for orientation, motion, and space during working memory largely overlap in parietal and frontal cortices (see Figure 1). Hence, feature-selective representations during working memory could arise at multiple representational scales of space that are distributed across the cortical hierarchy.

sensory cortex in a similar way as the sample. Thus, their traces would likely interfere if they were maintained in the same low-level area (see [43,94-97]). A solution would be to transform the representation and store it in a manner that allows the comparison to subsequent stimuli. For example, the relevant features of the lion could be maintained in the PFC as a feature-selective attentional search template [122,123]. This would have the additional advantage of avoiding interference between sample and target stimuli, avoiding that they overwrite the representation of the sample [124]. If this is the case, the nature of stimulus representations in PFC should be strongly task dependent [123]. Finding a yellow lion among yellow giraffes, for instance, benefits from an attentional focus on shape rather than color, whereas finding a yellow lion among grey elephants will benefit more from an attentional focus on color rather than shape ([125], Figure 2C).

The transformation of a sensory item into a behavioral response can be observed at multiple hierarchical levels of the brain, ranging from sensory to prefrontal cortices [122,123,126-128], and culminates in concrete preparatory motor activity in the premotor and primary motor cortices [129,130]. Furthermore, as this process unfolds during the delay-period, task-related preparation signals are expected to gradually increase in strength and be strongest immediately before the onset of the test stimulus. This is indeed what is found in the PFC; at least in a subset of cells [6,43] (see also [95,131]), which suggests that these forms of persistent activity can also play a preparatory role, going beyond the memorization of sensory information. Such a response-oriented, prospective nature of representations in the lateral prefrontal cortex (IPFC) is further supported by delayed paired association tasks [132], where animals are required to learn an association between a sample and test stimulus, separated by a brief delay. Under these conditions, PFC neurons encode the transformation from a memory representation of the sample stimulus to a representation of the expected paired associate in preparation of the test stimulus presentation.

A Working Memory Gradient

We conceptualize working memory as a hierarchical process that links detailed sensory representations to specific behavioral responses via intermediate task-relevant representations and action plans in a network of brain areas that each represent the working memory in a format



that matches their functional specialization (see also [15]). Thus, we propose that every cortical region can produce persistent stimulus-selective activity if the features that are coded in that region need to be memorized across a delay to prepare for an upcoming response. Such a universal cortical capability to produce persistent activity explains a simple observation that may not have received the attention it deserves, and which we here put forward as a central hypothesis: all features that are represented in the neocortex can be briefly memorized. Violations of this rule would provide important insights into the mechanisms underlying working memory. As far as we know, however, this proposal holds for low-level as well as high-level sensory features, which can be stored as persistent activity in lower and higher areas of the cerebral cortex, respectively. It also holds for transformed representations that are used to guide upcoming behavior.

The two potential explanations for the distributed nature of working memory outlined above are not mutually exclusive. In fact, they both roughly map on a similar posterior-to-frontal axis of functional brain organization. At the posterior end, the sensory cortices represent incoming sensory information in a relatively pure and detailed form. At the frontal end of the gradient, the frontal cortex represents information that is abstracted and transformed in support of upcoming behavior. Some of the frontal representations are abstract and categorical [109], however, other representations more directly support the execution of actions [15,133], and code for remembered action features such as the location of objects in eye coordinates for eye movements [134], in arm coordinates for hand movements [129], or even in object coordinates if this is required for the task [135].

Areas in between the sensory and prefrontal cortices might maintain intermediate transformations in many different formats. The gradients mentioned can be observed at multiple stages and reflect the increasing levels of processing and abstraction across subsequent stages of stimulus processing. One major visual stream, for example, originates from the primary visual cortex, and then proceeds via the extrastriate and inferior temporal cortices to the IPFC. It is well established that the tuning of single cells in these regions exhibits increasing degrees of spatial invariance, stimulus generalization, and abstraction. Whereas cells in V1 are selective to the detailed lowlevel features of stimuli, cells in inferior temporal cortex exhibit translational invariance and they can generalize across different low-level features [136]. Consistently, inferior temporal cells exhibit persistent stimulus-selective activity for objects during working memory delays [54]. Our review of the literature (see Figure 1) is generally consistent with the notion of a gradient with increasing levels of abstraction of memory signals across different levels of processing.

Which nodes maintain working memory representations depends on which format is the most suitable to perform the task at hand. When, for instance, detailed sensory information is required to perform a task, it makes sense to store stimulus information in areas suitable to represent such details, like the early sensory cortex. In contrast, when a task promotes the use of categorical information, it may be stored in a more abstract or verbal format by neurons higher in the processing hierarchy. Finally, if the appropriate response is known as soon as a sample stimulus is shown, working memory can also be encoded in the form of a pure response plan [137].

The notion of a distributed working memory network that postulates the ability of retention as a general principle in the neocortex has a long history [133,138]. Already early on, behavioral studies have revealed the effects of varying levels of processing in memory [139] and have distinguished between stores for different stimulus modalities [1]. More recent accounts of distributed working memory have emphasized different timescales of retention in different brain areas [140,141], the role of attentional processes in the retention of memories [4,11,142], and the utility of synaptic plasticity and dynamic firing trajectories for working memory [2,6,32,35,143].



Our hypothesis is ignorant about the precise mechanism for persistent activity: it might be generated within the cortex itself or rely on recurrent loops through subcortical structures, such as the thalamus or basal ganglia [144-147]. Furthermore, the working memory gradient outlined above does not require a separate homunculus control mechanism [145,146,148], but can instead be produced by the underlying neural circuits themselves, similar to the idea of a **Turing** machine ([149], Box 3). Indeed, recent computational studies have reported biologically plausible learning rules that can induce persistent activity for working memory, specifically for those features that need to be kept online during memory delays [150,151].

The ability of the brain to simultaneously encode working memory representations in multiple regions and at different levels of abstraction and response preparation necessarily involves redundancy. One advantage of redundancy is that it strongly increases the robustness of working memory representations. One study investigated simultaneous representations of a motor plan in the left and right premotor cortex. It was found that memory signals could be restored from the contralateral (i.e., redundant) side when one side was optogenetically silenced [152]. More generally, simultaneous and partially redundant representations might reflect the effects of representational loops that have been postulated for visual (attention-based rehearsal, [11]) and verbal (phonological loop, [1]) working memory.

Given the abundance of neurons capable of persistent activity, it is remarkable that the capacity to retain information in working memory is severely limited [153,154]. Hence, there must exist restrictions on the number of coactive memories per node and/or the number of coactive nodes. While individual stores might already have capacity restrictions [46,155], we suggest that properties of the overall working memory network might contribute to limitations of capacity as well (see Box 4). Specifically, if the maintenance of every individual memory item relies on the

Box 3. Distributed Control of the Working Memory Network

Here, we propose that signals in all cortical regions can exhibit a short-term buffering of information, depending on the nature of the task. Accordingly, working memory contents can range from low-level to abstract features and from sensory to motor codes. This raises the question how the distributed storage across multiple regions is coordinated. Theoretically, it might be conceivable that one or a few regions specifically control encoding, storage and retention, similar to a central executive [1]. However, it is also conceivable that control itself is distributed [176,177]. On the one hand, prefrontal regions can exert top-down attentional control and influence which low-level features are selected to be encoded in working memory. On the other hand, a low-level sensory cue (or its working memory) encoded in the sensory cortex can define a task and thus influence activity in the PFC. Thus, control cannot only go top-down but also bottom-up, and some sense of control thus seems to be present across the entire hierarchy.

The proposal that every brain region can exert an influence over processing in (most) other regions when it is required by the task is conceptually related to models that only involve states and transitions between states. Such models, for instance Markov processes [178] or Turing machines [179], do not require a centralized control system. A Turing machine is a hypothetical apparatus developed by Alan Turing to address the nature of computational operations. Turing machines consist of a finite-state machine and an infinite tape. Actions are selected based on the internal state of the machine and the symbol of the tape currently 'seen' by the machine. Selected actions then cause the machine to change its internal state by writing a new symbol to the tape, and/or shifting the tape to reveal the next symbol. The computational versatility of Turing machines arises from this continuous cycle of actions and the updating of internal states and 'tape memory'. Importantly, the tape can specify simple input but also entire programs using a similar format. Clearly, there are many differences between Turing machines and brains (e.g., Turing machines are serial and lack the distinction between long-term and working memory). Yet, there are also interesting analogies with working memory processes in the brain [149]. For instance, working memories are internal neural states, which may encode previous sensory inputs or even complex task rules. These memories in turn influence the selected actions. Motor actions influence incoming sensory information (e.g., arm movements influence proprioception and eve movements change visual input). Likewise, internal actions, like attention shifts or the recall of associations, modulate active mnemonic codes in the frontal cortex and lower brain areas. Thus, selected actions can impact the pattern of sensory input, the activated mnemonic codes, or both (just as actions of Turing machines change the internal state, the tape, or both). Consequentially, new actions can be selected at the next time step, causing new changes in sensory input and working memory contents. This succession of action selection and updating of sensory and mnemonic codes in working memory resembles the workings of a Turing machine.



Box 4. The Limited Capacity of Working Memory

The wide distribution of persistent stimulus-selective responses in the primate brain yields the question whether capacity limitations in working memory reflect properties of individual storage regions, or a property of the distributed network. Behavioral models assume either a slot-based [180,181] or resource-based [182,183] limitation of capacity. At a neural level, several factors might limit memory capacity.

Competition for representation: multiple items may compete for representation within the same representational maps (e. g., of visual space in V1) or in any other limited 'cortical real estate' [155], such that individual memorized items interfere with each other during the delay period or already during encoding [46]. In line with this, individual visual working memory capacity is correlated with the volumetric size of V1 [184], and persistent stimulus-selective activity per memory item decreases with increasing load [12]. Capacity increases with the sparseness of neural representations that leads to less interference between representations [40]. It can be further improved by synaptic facilitation [40]. In contrast, capacity might be a global rather than local property of the working memory network. For example, memorization of low-level visual features reduces the ability to retain low-level auditory information [185].

Distributed storage: the above-mentioned interference among memory representations could decrease if different individual stimuli of the same type (e.g., multiple oriented gratings) were represented in different brain areas. Memorized stimuli that are currently relevant for an ongoing task and that are in the focus of attention [186] might, for instance, be retained by detailed representations in sensory cortex, while secondary memory items that are stored for later use are retained in a coarser, more abstract form in anterior regions [187]. This would explain why persistent stimulus-selective activity in sensory regions is weaker or absent for items that are stored outside the focus of attention, while it is reinstated as soon as they regain their relevance [30,76,188].

Task-relevant detail: the level of detail required for a working memory task has implications for capacity. The posterior-tofrontal gradient of increasing abstraction proposed here (see 'Different Levels of Abstraction') can be interpreted as a form of complexity reduction [189] or 'chunking' [153]. For example, 'd', 'o', and 'g' are more easily retained as a word ('dog') than as individual letters [190]. Similarly, and in accordance with the flexible allocation of memory resources [182,183,191], higher cortical areas can retain larger chunks of low-level features that would be difficult to store individually in lower level regions. However, such chunking involves a step of abstraction that might discard low-level visual features

activity within a widespread network, the representations of multiple memoranda will more readily interfere with each other. Consequently, the selection of the nodes that will retain stimulus- or response-selective information across a delay interval should strongly depend on task requirements [80,88] to avoid interference (e.g., from distractors, [43,94-97]) and thereby boost memory capacity (see Outstanding Questions).

Concluding Remarks

There is abundant evidence for widely distributed stimulus-related information in sensory, parietal, and prefrontal cortices during working memory delays. In the absence of evidence implicating any of these regions as an exclusive and localized store of memory contents, we suggest that working memory is better characterized as a distributed network that gradually transforms sensory information towards an appropriate behavioral response, across a temporal delay. Persistent stimulus-specific activity might be observed anywhere in such a network and at any stage of transformation. Localization will thus strongly depend on the precise requirements and context of the task. This notion suggests that perhaps the field of working memory should shift its focus from asking where in the brain working memories are stored to unraveling how a range of highly specialized brain areas together transform a sensory stimulus into an appropriate response and how this process is sustained as a working memory across delays.

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Outstanding Questions

What is the mechanism of retention? Two main neurocomputational mechanisms of retention have been proposed: (i) activity-based retention, such as spiking activity, reverberatory activity, feedforward-connected subgroups of neurons (synfire chains), oscillatory dynamics, or high-dimensional dynamic trajectories; and (ii) activity-silent retention, where stimulus-selective representations are maintained as a pattern of synaptic weights. These two mechanisms are not mutually exclusive and might jointly contribute to retention.

How does the current relevance of a memorized item affect its retention? Behavioral studies suggest a distinction between attended and unattended (accessory) items held in memory, but persistent stimulus-specific activity has only been reported for attended items. Unattended representations might be retained (i) as activity-silent synaptic representations, (ii) using a different neural code than attended representations (e.g., on different levels of abstraction; see Box 4), or (iii) as weaker activity patterns then attended ones.

How can we distinguish between true mnemonic activity of a feature and encoding of other stimulus-related factors? Different sensory stimuli are known to be associated with differences in valence, motor affordance, and ease of recognition. All these features are known to affect signals across the neocortex. Thus, signals that appear to be stimulus selective might instead reflect encoding of these other features.

How is the retention of multiple items coordinated and how do these memories interact? Storing stimulus information in memory in multiple formats and areas in parallel can increase working memory robustness. However, to date it is unclear: (i) how multiple memory representations interact, (ii) which mechanism selects which memory will guide behavior, (iii) how parallel representations can compensate for perturbations and interference (e.g., by distracting stimuli), and (iv) how the retention of a particular working memory content is terminated.



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