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Transforming the Concept of Memory Reactivation

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Abstract

Reactivation refers to the phenomenon wherein patterns of neural activity expressed during perceptual experience are re-expressed at a later time—a putative neural marker of memory. Reactivation of perceptual content has been observed across many cortical areas and correlates with objective and subjective expressions of memory in humans. However, because reactivation emphasizes similarities between perceptual and memory-based representations, it obscures differences in how perceptual events and memories are represented. Here, we highlight recent evidence of systematic differences in how (and where) perceptual events and memories are represented in the brain. We argue that neural representations of memories are best thought of as spatially transformed versions of perceptual representations. We consider why spatial transformations occur and identify critical questions for future research.

Keywords

Episodic memory; reactivation; reinstatement; memory transformation; sensory cortex; frontoparietal cortex

Moving beyond memory reactivation

When remembering an event from the past, it often feels as though we are re-experiencing the content of that event, bringing to mind the sensations, emotions, or thoughts that characterized the initial experience. This subjective quality of memory is paralleled by an important neural phenomenon: that **content-sensitive patterns of neural activity** (see Glossary) evoked during the initial experience of an event are re-expressed when that event is retrieved from memory. This phenomenon is referred to as **reactivation** and has been an influential tool for studying **episodic memory** in humans. Neural measures of reactivation have been shown to be predictive of the accuracy [1–3], subjective vividness [4,5], and consequences of **memory retrieval** [6–8]. Another notable aspect of reactivation is that it

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can be measured in contexts such as rest [9–11] or sleep [12,13] when behavioral recordings are not feasible. Over the past two decades, there have been tremendous advances in both the computational sophistication and the sensitivity of the methods used to measure reactivation (see Box 1), but these methods exploit the same fundamental phenomenon of a ‘match’ between content-sensitive neural activity patterns evoked during perception and memory retrieval. While this emphasis on the similarity between perceptual and memory-based representations is appealing and is unquestionably important, here we argue that it also provides an incomplete view of how the contents of memories are expressed in the brain during memory retrieval. In particular, because reactivation, by definition, tests for similarities between perception and retrieval, it necessarily fails to capture ways in which content representations *differ* between perception and memory.

One obvious way in which memory-based and perceptual representations might differ is in terms of the amount of information they contain, with memory-based representations presumably being a weaker or degraded version of perceptual representations. Empirically, this might manifest, for example, as a reduced ability to decode the content of an event from neural activity patterns evoked during retrieval compared to patterns evoked during perception. However, a number of recent studies have demonstrated clear violations of this idea: specifically, some brain regions contain *more information* about the content of an experience when that experience is being retrieved from memory compared to when it is perceptually experienced [14–17]. The clear implication of this is that measures of reactivation—which only test whether activity patterns evoked during retrieval are similar to those evoked during perception—will systematically underestimate the amount, strength, or qualities of content representations during retrieval.

In this Opinion article, we argue that there is emerging evidence that content representations during memory retrieval are spatially-transformed versions of content representations originally expressed during perception. We use this idea of **spatial transformation** to emphasize that content information (a) is expressed in *different* brain regions during perception and memory retrieval and (b) that these differences in spatial localization are *systematic* (or predictable) as opposed to random noise or signal degradation during memory retrieval (Figure 1). While the possibility of systematic differences in content representations during perception and memory retrieval has, to date, been overshadowed by the more dominant idea of reactivation, it is important to note that cognitive theories of episodic memory have long argued that memory retrieval is more of a constructive act than a reproductive act [18]. Thus, there is a strong theoretical motivation for moving beyond measurements that only index the degree to which content representations at retrieval resemble those at perception and moving toward approaches that characterize how and why content representations at retrieval differ from those at perception. Below, we describe some of the recent evidence supporting the specific proposal of spatial transformation from perception to memory. We consider several accounts of why spatial transformations occur and highlight several avenues for future research. We argue that characterizing and understanding perception-to-memory transformations (spatial and otherwise) represents a logical, timely, and important next step for the field.

Spatial transformations within sensory cortex

While memory reactivation has been reported in many parts of the human brain, it has been most frequently measured in sensory areas, and visual cortex in particular. Early studies demonstrated that visual cortex is active both during visual perception and during retrieval of visual memories [19], demonstrating a coarse-level reactivation of the sensory modality of experience (e.g. vision vs. audition). More recently, studies have established reactivation of increasingly fine-grained information within visual cortical areas. For example, numerous studies have shown that patterns of activity in ventral temporal cortex that reflect the visual category of a stimulus (e.g., face vs. scene vs. object) are reactivated during retrieval of that stimulus [1,20–22]. Further, patterns of neural activity in early visual areas (e.g., V1) that reflect low-level stimulus properties such as spatial location, spatial frequency, edges, and orientation, are also reactivated during retrieval [23–26].

The common thread among the examples of visual cortical reactivation described above is that the same content-sensitive patterns of activity evoked during perception are re-expressed during memory retrieval. Yet, results from a number of recent studies suggest that there are subtle, but systematic differences in the spatial localization of content information within visual cortex during perception and memory retrieval. In particular, several studies have found that scene-selective responses during perception are located posterior to scene-selective responses during memory retrieval. For example, a meta-analysis of over one hundred fMRI studies found that scene perception recruits relatively posterior aspects of the parahippocampal place area (PPA) whereas retrieving scenes from memory recruits relatively anterior aspects of PPA [27]. In an unpublished study, this striking dissociation was replicated in a paradigm that directly compared perception and memory retrieval using a within-subjects design [14]. Similar posterior (perception) to anterior (memory retrieval) shifts have been observed in other scene-selective areas, including lateral occipital and medial parietal areas [14,28]. Resting state fMRI analyses have shown that anterior and posterior aspects of ventral, lateral, and medial scene-selective areas have different connectivity profiles with the rest of the brain, consistent with the idea that scene perception and scene memory are supported by segregated visual cortical networks [14,27–29]. Related work has suggested that this posterior to anterior division between visual perception and visual memory extends to stimulus classes other than scenes, such as objects, and to aspects of lateral and ventral temporal cortex outside of the scene network [30,31].

Collectively, these studies reveal a shift in the spatial localization of content-sensitive responses during perception and memory retrieval in the visual system. Specifically, information about a stimulus is preferentially expressed in different regions of visual cortex depending on whether that stimulus is currently available to sensory receptors or recalled from long-term memory. Importantly, the fact that some regions of the visual system contain *more information* about remembered stimuli than they do about perceived stimuli is fundamentally inconsistent with the idea of reactivation. Moreover, these differences in spatial localization are highly systematic (as opposed to random noise or signal degradation) and generalize across different classes of stimuli and visual regions. As such, these findings strongly suggest that content representations during perception are re-expressed in a spatially transformed manner during memory retrieval.

Spatial transformations from sensory cortex to frontoparietal cortex

Though memory reactivation has been most consistently observed in sensory areas, recent applications of pattern-based analysis methods have revealed robust, content-sensitive reactivation in frontoparietal regions [5,7,32,33]. This evidence of frontoparietal reactivation is surprising given the traditional view that frontoparietal regions are involved in controlling or evaluating memories (i.e., memory *processes*) without actively representing retrieved content [34]. More surprisingly, however, several studies have directly contrasted the strength of content representations during perception and memory retrieval and have found that, among some frontoparietal regions, content representations are not only present during memory retrieval, but are *stronger* during retrieval than during perception [15–17].

In one study, lateral parietal activity patterns contained more information about the features of visual objects (i.e. the patterns were more correlated within feature than across feature) during memory retrieval than during perception [17]. This contrasted sharply with visual cortical areas, where feature representations were stronger during perception than during memory. Similarly, another study found that representations of individual stimuli were stronger in lateral frontal and lateral parietal regions during memory retrieval than during perception, whereas ventral temporal cortex exhibited the opposite pattern [16]. Strikingly, the fine-grained representational structure among stimuli (i.e., the relative distance between stimuli in representational space) was preserved from perception to memory retrieval despite the change in spatial localization from ventral temporal cortex to frontoparietal cortex. This evidence of preserved information despite changes in spatial localization is strongly consistent with the idea that memory retrieval involves a spatial transformation of content information expressed during perception.

In both of the studies discussed above, content representations were stronger in frontoparietal regions during memory retrieval than during perception—a finding that is fundamentally inconsistent with the idea of reactivation. Specifically, ‘perfect’ reactivation should correspond to memory representations that are as strong as, but not stronger than, perceptual representations. Instead—and consistent with posterior/anterior dissociations in visual cortical areas described in the prior section—these findings indicate that some brain regions preferentially represent stimuli or events that are retrieved from memory, and that other brain regions preferentially represent stimuli or events that are perceptually available. That said, it is important to note that even in studies that have observed a shift of content representations from visual cortical areas during perception to frontoparietal regions during retrieval, there is still some degree of reactivation both in ventral temporal cortex and in frontoparietal cortex [15–17]. Thus, the presence of transformation does not require the absence of reactivation. Rather, reactivation and transformation likely *co-occur*. By definition, however, these phenomena explain unique variance in retrieval-related representations. Understanding the factors that determine the relative degree of reactivation vs. transformation is a critical open question (see Outstanding Questions).

Why is there spatial transformation?

Why might stimulus representations be spatially transformed from perception to memory? While current evidence does not allow for a definitive answer to this question, below we discuss three potential explanations (which are not mutually exclusive) that may help guide future studies.

Distinct neuroanatomical origins

One reason why perception- and memory-based representations may be preferentially expressed in different brain regions is because the signals that drive these representations have distinct neuroanatomical origins. While perceptual representations originate from sensory receptors such as those in the retina, retrieved memory representations originate from a pattern completion process commonly thought to be triggered by the hippocampus [35,36]. Although computational perspectives on memory have emphasized the relevance of feedback connections from the hippocampus to sensory cortex in enabling sensory reactivation [37], the connectivity profile of the hippocampus suggests that memory retrieval involves more than sensory reactivation. Most notably, the hippocampus is intimately connected to the **default mode network** [38,39] and this coupling is particularly strong during memory retrieval [40,41]. The default mode network is so-named because regions in this network tend to show higher activation during internally-oriented cognition (including memory retrieval) than to perceptual stimuli. However, several recent studies have found that this bias toward internally-oriented cognition also manifests in stronger content representations in the default mode network during memory retrieval than during perception [15–17].

Connectivity with the hippocampus may also explain the anterior bias for memory-based representations in the visual system, as the same anterior regions that exhibit a memory bias also exhibit relatively stronger connectivity to the hippocampus [14,27]. It is also of note that the hippocampus, itself, has been shown to more strongly represent retrieved content than perceived content [16,42]. Thus, just as the hippocampus plays a critical role in driving reactivation in sensory cortex, it is also likely to play a key role in mediating transformation by driving unique cortical responses during memory retrieval that were weak or not present during perception. From this perspective, the degree to which a given cortical area displays a preference for perceptual vs. memory-based content may be explained by the degree to which that cortical area is more strongly driven by signals from sensory receptors vs. signals from the hippocampus. This idea can be readily tested in targeted studies that combine connectivity measures with measures of content-sensitivity. This account also makes interesting and testable predictions for memory tasks that vary in their dependence on the hippocampus. For example, recognition memory tasks, which only partially depend on hippocampal processing [43], should involve less transformation under this account. Of course, this account leaves open the question of *why* there are partially distinct neuroanatomical pathways for perception and memory retrieval and whether having segregated content representations is adaptive. Although speculative, it may be the case that segregated representations of perceived and remembered content serve the simple purpose of avoiding confusion between the current environmental state and past environmental states.

Distinct task demands

Another potential account of spatial transformation—particularly from visual regions to frontoparietal cortex—is that memory retrieval imposes unique (and perhaps greater) task demands compared to perception. This account is motivated by the known role of frontoparietal regions in implementing attention and control processes [44,45] but specifically requires that frontoparietal regions meet task demands by *actively representing retrieved content* [5,32,46]. For example, frontoparietal regions have been shown to flexibly prioritize retrieved content that is relevant to current memory demands [17,47,48] and to actively insulate retrieved content from sensory distraction [49,50]. From this perspective, whether frontoparietal cortex displays a preference for perception vs memory-based content can be explained by the degree to which content representations must be brought in line with task demands during perception vs memory.

In our view, a “task demands” account does not fully explain existing evidence of transformation. First, this account more readily explains transformations from sensory to frontoparietal cortex than transformations within sensory cortex. Second, this account would predict that the specific frontoparietal regions exhibiting a bias toward memory representations would be the same regions most involved in control or attention. In fact, a bias toward retrieved content is particularly pronounced in regions that belong to the default mode network [16,17]—a network that is associated with internally-oriented cognition and not top-down attentional control. Finally, task relevance does not appear to be a requirement for frontoparietal regions to exhibit a bias toward memory-based representations. In one study [17], task relevance did influence the strength of memory representations in dorsal regions of parietal cortex (the intraparietal sulcus), but in ventral parietal regions (including components of the default mode network), the strength of memory-based representations was insensitive to task demands, with equivalent representation of mnemonic information that was relevant vs. irrelevant to the current task. Strikingly, regions of the rodent parietal cortex have also been shown to contain a bias toward past experience over current perceptual experience, even when past experiences are entirely irrelevant to current behavioral decisions [51]. Taken together, while some frontoparietal regions may be particularly involved in representing retrieved content in a way that aligns them with task demands, this framework, we would argue, is unlikely to fully account for spatial transformations from perception to memory. That said, the “task demands” account is also testable in that task demands can be independently manipulated during perception and memory retrieval in order to determine whether—or for which brain regions—content representations become stronger as the demand for attention or control increases.

Shift toward conceptual representations

Another account of spatial transformation is that the *nature of content representations* changes from perception to memory. The most obvious version of this account is that, compared to a perceptual representation of an event, a memory-based representation of the same event will reflect relatively more conceptual information. For example, whereas perceiving an apple typically requires at least some low-level processing of stimulus features (contrast, color, spatial frequency, etc.), retrieving the same apple from memory may produce a representation that omits some of these low-level features (i.e., that is less

perceptually specific) and is instead biased toward higher-level, conceptual properties derived from general knowledge (“apples are a healthy snack”). This perspective is motivated by evidence that some of the same frontoparietal [52] and anterior ventral temporal [53,54] regions that demonstrate biases toward memory-based representations are also involved in conceptual processing. From this perspective, the degree to which a given cortical area displays a preference for memory-based content should be explained by the degree to which that cortical area codes for conceptual features. Notably, the idea that memories involve a shift toward conceptual representations makes contact with the idea of consolidation—a point we consider in Box 2. Additionally, it is interesting to note that conceptual representations necessarily involve integrating across distinct experiences (e.g., prior encounters with apples), raising the question of whether this integration over *time* is a relevant quality. Indeed, regions of the default mode network that exhibit biases toward retrieved memories and that have been implicated in conceptual processing have also been shown to have wider temporal receptive windows than sensory regions [55–57].

While an account that emphasizes a shift toward conceptual representations is appealing in many respects, it remains to be seen whether this account can explain recent evidence suggesting spatial transformation in the rodent brain [51] or emerging evidence of spatial transformation in working memory tasks that rely on simple visual stimuli (see Box 3). This account also makes a notable testable prediction: if spatial transformations reflect differences in the perceptual vs. conceptual content of representations, then the degree of transformation should be modulated by the kind of representation accessed during perception and memory retrieval. For example, if perception and retrieval both require conceptual representations, the resulting content representations should be more similar (i.e., relatively less transformation and relatively more reactivation).

Other forms of transformation

Though this opinion article focuses on spatial transformation, **transformation** is likely to occur along other dimensions as well. Below, we briefly mention other forms of transformation that do not fit our definition of spatial transformation. Although detailed consideration of these other forms of transformation is beyond the scope of the current article, these other lines of evidence reinforce the critical conceptual point that memory representations differ from perceptual representations in systematic ways—differences that strict measures of reactivation will not capture. Ultimately, theoretical accounts of how memories are expressed in the brain during retrieval should integrate all of these lines of work coherently.

First, recent evidence suggests that even when the same neurons or voxels are active during perception and memory retrieval, their tuning for stimulus features may change (**tuning transformation**). For example, the same visual cortical voxels are tuned to more foveal eccentricities and lower spatial frequencies during memory retrieval than perception [58]. These voxels also pool over larger extents of visual space during memory retrieval than perception [58,59]. Similarly, at least some neurons in monkey lateral prefrontal cortex exhibit different motion direction tuning during perception and during working memory maintenance [60]. Other work suggests that electrophysiological responses in sensorimotor

and premotor cortex may switch from reflecting sensory tuning during speech perception to reflecting motor tuning during speech rehearsal [61].

Second, a large body of evidence from animal studies—and more recently from human studies—indicates that neural activity undergoes several kinds of **temporal transformations** from perception to memory retrieval. In rodents, it is well established that **replay** of hippocampal place cell sequences involves *temporal compression*, in that sequences of cell firing occur much faster offline (during rest or sleep) than online (during exploration) [62]. Behaviorally, humans also express memory compression [63,64], and initial evidence for human hippocampal replay and compression has recently been reported using non-invasive measurements of population activity [65–68]. Additionally, memories are influenced by *event segmentation*, a process that chunks continuous perceptual experience into discrete events [69]. While some segmentation processes influence online perception as well as memory, other segmentations processes occur after perception and may specifically influence how temporal sequences are remembered [70,71]. The extent to which these processes result in activity patterns during retrieval that diverge from earlier perceptual activity is an interesting question for future research. Finally, memory retrieval may involve a *reversal* of the temporal activity flow present during perception, such that higher-level brain regions engaged during relatively late stages of perceptual processing are invoked during initial stages of memory retrieval [72–74]. These findings and other observations about the temporal structure of reactivation have been recently reviewed in detail [75].

Concluding remarks

It has long been appreciated that perception and memory retrieval require distinct cognitive processes [76,77] and engage distinct neural structures [78–80]. Yet, measures of memory reactivation fundamentally focus on the similarity of content representations across perception and memory retrieval. While reactivation has been a crucially important phenomenon in the study of memory, we argue that there is a pressing need to better understand the differences between neural representations of perceived and remembered events. Rather than viewing these differences as the product of noise or imprecision in memory retrieval, we argue that there are systematic spatial transformations from perception to memory that are better understood as reformatting of information, either within brain regions or across them. Many open questions remain concerning the nature, causes, and consequences of transformations from perception to memory. These questions represent exciting areas for future research that will hopefully lead to a more comprehensive understanding of how the brain recreates experiences from the past. To conclude, we highlight some specific goals that could guide future research.

One specific target for future studies is to mathematically model the transformation from perceptual to memory-based neural representations. In the studies reviewed here, we have focused on predictable shifts in brain regions that represent perceptual and memory-based content. However, no studies to date (to our knowledge) have specified *transfer functions* that predict the spatial activity pattern that an individual memory will evoke based on the corresponding activity pattern evoked during perception. Techniques for predicting neural activity patterns across different brain regions have recently been described [81,82] and

these techniques can be readily applied to predict perception-to-memory transformations as well. A critical advantage of approaches which yield concrete (mathematical) predictions of spatial activity patterns is that they allow for precisely quantifying the amount of variance in memory-based activity patterns that is explained by perceptual activity patterns in different brain regions (transformation) versus the re-expression of the same perceptual activity patterns (reactivation). Model-based approaches to identifying transfer functions would represent a significant advance over current approaches (reviewed in this article). Model-based approaches could also be extended to quantify the degree to which variables such as task demands and stimulus properties influence the relative degree of transformation versus reactivation (see Outstanding Questions). Finally, comparing transfer functions across individuals will also allow for deeper understanding of the extent to which transformations are idiosyncratic versus shared across individuals [33].

A second important target for future studies is to incorporate finer-grained measures of content representations when considering spatial transformations. The studies reviewed here relied on category-level representations [15,28], exemplar-level representations [16,33] and, to a lesser extent, feature-level representations [17]. The fact that evidence for spatial transformation can be observed at many levels of representational specificity is important, but feature-level representations hold unique appeal in that they allow events to be decomposed such that potentially subtle changes in content representations can be measured. For example, memory-based representations may differ from perceptual representations in terms of dimensionality (number of features) or the weighting of specific features (e.g., perceptual vs. conceptual features). Thus, feature-based approaches [17,25] may provide key insight into spatial transformations. Indeed, a complete understanding of how perceptual experiences are transformed into memories will require understanding not only how and when spatial patterns of neural activity differ between perception and memory but also understanding subtle differences in the information contained within those patterns of neural activity.

Finally, while there is some evidence from rodents consistent with our proposal of spatial transformation [51], the ideas described here are largely motivated by human neuroimaging studies. Given that memory reactivation has been extensively studied in both animal models and humans, it will be valuable to test the predictions of spatial transformation more systematically across species. In fact, the ideas we propose here are particularly well suited to cross-species comparisons (so long as multi-site recordings are feasible in the animal model) given that there are well-developed analytic approaches for measuring content representations that transcend recording methodologies and species (e.g. [83]).

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Glossary

Content-sensitive patterns of neural activity

Patterns of neural activity that reflect ‘what’ is being currently experienced or remembered. Content sensitivity may reflect sensitivity to broad visual categories (e.g., faces vs. scenes vs. objects), specific exemplars (e.g., apple vs. butterfly), or specific feature values of an exemplar (e.g., color: red vs. yellow). Content sensitivity can be contrasted with process-level responses that are invariant to the content of an experience.

Default mode network

A network of regions spanning medial and lateral temporal cortex, medial and lateral parietal cortex, and medial prefrontal cortex. These regions show higher activation during internally-oriented tasks than externally-oriented ones and may play a key role in representing the content of retrieved memories.

Episodic memory

A form of declarative long-term memory that allows for recollection of unique previous experiences.

Memory retrieval

Accessing a previously stored event from memory

Reactivation

Phenomenon whereby content-sensitive patterns of neural activity evoked during perception are re-expressed when that experience is remembered. Reactivation has been measured using multiple methodologies (e.g., fMRI, EEG) and analytic approaches (e.g., univariate activation, pattern similarity, decoding models).

Replay

Reactivation of temporally-structured sequences of neural activity (most typically, single-cell firing).

Spatial transformation

A form of transformation in which the spatial localization of content-sensitive neural activity patterns changes from perception to memory retrieval. Whereas activity patterns in some brain regions preferentially carry information about perceptual experience, activity patterns in other brain regions preferentially carry information about these same experiences when they are retrieved from memory.

Temporal transformation

A broad class of transformations in which the temporal organization of neural activity changes from perception to memory retrieval.

Transformation

In the context of this article, the term transformation is used to refer to a systematic and predictable change in neural activity from perception to memory retrieval that cannot be attributed to memory noise or error.

Tuning transformation

A form of transformation in which the preferred feature values of an individual neuron, voxel, or population of neurons/voxels changes from perception to memory retrieval.

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Box 1: Methods and approaches for detecting reactivation

Non-invasive neuroimaging methods

Memory reactivation has been extensively studied using non-invasive imaging techniques (e.g., fMRI, EEG, MEG), which can simultaneously record population neural activity from large parts of the brain. Below, we detail several popular analytic approaches for measuring reactivation in neuroimaging data. All of these approaches test for similarity in profiles of neural activation during perceptual experience and memory retrieval. Notably, these methods can be applied at varying levels of specificity: stimulus category (e.g., faces vs. scenes vs. objects), specific exemplar (e.g., apple vs. butterfly), or stimulus feature value (e.g., color: red vs. yellow).

Univariate Activation

Brain regions are identified for which the mean level of neural activity varies according to the stimulus content. Reactivation is demonstrated by brain regions exhibiting similar content sensitivity (changes in mean activity level) during perception and memory retrieval [19].

Pattern Similarity

Neural activity patterns (activity across voxels or sensors) are measured during perception from a specific brain region. Patterns are then measured within the same brain region during memory retrieval. Reactivation is demonstrated by correlation between a perception pattern and a corresponding retrieval pattern that exceeds the correlation between non-corresponding patterns [2,84].

Decoding Models

Supervised models are trained to learn a mapping between stimuli or stimulus classes and neural activity measured during perception. These models are then tested on neural activity measured during memory retrieval. Reactivation is demonstrated by classifier transfer from perception to retrieval: i.e., the perception-trained model accurately labeling stimuli retrieved from memory [1,22].

Encoding Models

Models that specify mathematical relationships between physical stimulus properties (e.g. pixel values) and neural activity are fit to neural data measured during perception. Reactivation is demonstrated if model parameters generalize from perception to retrieval: i.e., if the perception parameters can be used to accurately predict brain activity during memory retrieval [85]. Predicted brain activity can also be used to decode remembered stimuli (encoding-decoding) [24,25] or encoding models can be inverted to reconstruct remembered stimuli (inverted encoding model) [46].

Invasive physiological and optogenetic methods

While the current article focuses on findings from non-invasive neuroimaging methods (which benefit from large-scale spatial coverage of the brain), all of the forementioned methods can be applied to LFP or single-cell electrophysiological activity measured

invasively in humans (when treated for neurological diseases requiring electrode implants) or in animal experiments [86]. Other approaches, such as large-scale or multiple-region electrophysiological recordings, are more prevalent in animal models, and some approaches are restricted to animal experiments. For example, large single-unit datasets collected from rodents have been essential in characterizing the phenomenon of replay, or the temporally-structured reactivation of single-cell firing sequences [62]. Only recently has replay of single cell firing been shown in humans [87]. Optogenetic techniques, applicable in animal studies solely, have revealed that *experimentally-induced* reactivation in rodents is sufficient to evoke memory behavior [88].

Box 2: Transformation versus consolidation

The idea that memories undergo transformation is not new to the field. For instance, numerous prior empirical and theoretical papers have argued that memories are transformed from hippocampally-dependent traces to cortically-dependent traces via a consolidation process that operates over long timescales [89,90]. This clearly makes contact with the idea of spatial transformation proposed here. It also makes contact with some of the explanations we offer for transformation—namely, consolidation is thought to enable the extraction of conceptual knowledge from individual episodes [90,91]. However, our proposal is also distinct from the standard view of consolidation. In particular, we argue for an *immediate* transformation wherein there are systematic differences in how perceptual and memory-based content is represented. In contrast, consolidation can be thought of as more gradual, memory-to-memory transformations. That said, the distinction between these ideas may be blurred in some cases. For example, several recent papers have raised the possibility that repeated retrieval accelerates consolidation processes that normally occur over longer timescales [92–94]. Further, a recent paper demonstrated that just one session of retrieval practice (as opposed to restudying information) is sufficient to shift the spatial localization of content representations during subsequent memory retrieval [95]. Thus, it is possible that repeated retrieval may also be a factor that influences the degree of perception-to-memory representations.

Box 3: Parallels to working memory

Many neuroimaging studies have demonstrated that visual cortex contains representations of stimuli held in working memory [96,97] and that the quality of these representations relates to behavioral performance in working memory tasks [98,99]. Often referred to as “sensory recruitment” rather than reactivation, these findings convincingly demonstrate that visual cortex encodes stimuli during a delay period in a similar format to perception. However, recent work has raised the possibility that, in addition to sensory recruitment, sensory representations may also be transformed during working memory maintenance. In particular, one recent study [50] showed that while early visual areas code perceived and maintained orientations in a shared sensory-like format, parietal cortex codes perceived and maintained orientations in distinct formats. While speculative, it is possible that despite the dissociable mechanisms underlying working memory and long-term memory [100,101], common principles of spatial transformation apply across these domains. Indeed, the debate over the functional relevance of working memory representations in sensory vs. frontoparietal regions [102,103] in many ways parallels questions and findings related to the functional significance of long-term memory reactivation in sensory vs. frontoparietal regions [7,16,17,47,104,105]. Ultimately, understanding the nature of transformation in working memory and long-term memory and the extent to which they are similar will require a coordinated effort between working memory and long-term memory researchers as well as direct experimental comparisons across tasks.

Outstanding Questions

1. To what extent do changes in information content account for spatial transformation from perception to retrieval? Are certain stimulus features that are present during perception systematically lost or distorted in memory? Do memory representations *gain* information that is absent or weakly present during perception through integration with other memories or existing knowledge structures (schemas)?
2. What determines the relative degree of neural reactivation vs. transformation across brain regions observed during memory retrieval? For example, is greater transformation observed when memory tasks promote conceptual processing at retrieval? Conversely, is relatively greater reactivation in sensory areas observed when memory tasks promote perceptual processing? Does the relative degree of reactivation vs. transformation depend on whether memory tasks involve recall vs. recognition judgments? Do reactivation and transformation trade-off or are they independent?
3. Does the degree of transformation across brain regions depend on the temporal lag between perception and memory retrieval? Transformation potentially occurs in working memory paradigms with delays on the order of seconds, yet there is also considerable work documenting consolidation-related transformations at timescales of hours to years. What are the similarities and differences between transformations that occur across these vastly different timescales?
4. What is the relationship between transformation within sensory areas and transformation from sensory to frontoparietal regions? These two forms of transformation have been studied separately to date, and it is thus unclear whether they are related, and if so, how. Notably, the frontoparietal and sensory regions that exhibit biases toward memory-based representations are functionally connected with the hippocampus. To what extent can connectivity with the hippocampus explain both sets of findings?

Highlights

A foundational finding in the field of memory is that content-sensitive patterns of neural activity expressed during perceptual experiences are re-expressed when experiences are remembered—a phenomenon termed reactivation. However, reactivation obscures key differences in how perceptual events and memories are represented in the brain.

Recent findings suggest systematic, spatial transformations of content-sensitive neural activity patterns from perception to memory retrieval. These transformations occur within sensory cortex and from sensory cortex to frontoparietal cortex.

We consider why spatial transformations occur and identify critical questions to be addressed in future research. Understanding the ways in which memory representations differ from perceptual representations will critically inform theoretical accounts of memory and will help clarify how the brain recreates the past.

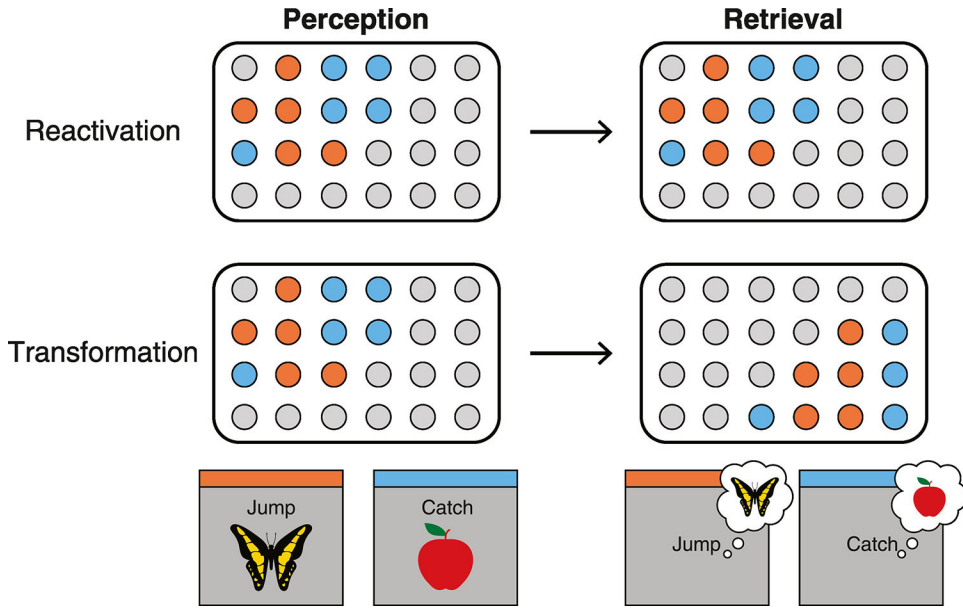


Figure 1. Schematic comparison of reactivation and spatial transformation.

During perception (left column), content-sensitive representations of perceptual events are encoded in cortex. We schematize this by highlighting the units that code for each stimulus (a butterfly and an apple) in different colors. The distinction between reactivation and transformation concerns how these content-sensitive representations at perception relate to content-sensitive representations during memory retrieval (right column). Reactivation (top row) proposes that the representation of an event measured during retrieval will be *similar* to the representation of the same event measured during perception. We schematize reactivation by highlighting the same neural units active during perception and memory retrieval. Though we illustrate it this way, the idea of reactivation does not argue that there will be a perfect match between representations measured during perception and retrieval; noise or forgetting may degrade this similarity. However, theories of reactivation do not propose or define any systematic differences between perceptual and memory-based representations, and typical measures of reactivation do not characterize potential differences. In contrast, the idea of spatial transformation (bottom row) proposes a *systematic change* in the neural localization of content representations from perception to memory retrieval. We schematize spatial transformation by highlighting different active units during perception and memory retrieval. Critically, spatial transformation cannot be attributed to noise or degradation because the changes are systematic—illustrated here by the rightward shift to a distinct cortical location. Changes in neural localization could occur within cortical regions (e.g., posterior-to-anterior shifts in sensory cortex) or across them (e.g., from sensory cortex to frontoparietal cortex).