Low-Resolution Neurocognitive Aging and Cognition: An Embodied Perspective

Jordan Mille**, Simona M. Brambati‡, Marie Izaute§ and Guillaume T. Vallet¶

1 CNRS, LAPSCO (UMR CNRS 6024), Université Clermont Auvergne, Clermont-Ferrand, France, 2 CRIUGM, Université de Montréal, Montréal, QC, Canada

Consistent with embodied cognition, a growing evidence in young adults show that sensorimotor processing is at the core of cognition. Considering that this approach predicts direct interaction between sensorimotor processing and cognition, embodied cognition may thus be particularly relevant to study aging, since this population is characterized by concomitant changes in sensorimotor and cognitive processing. The present perspective aims at showing the value and interest to explore normal aging throughout embodiment by focusing on the neurophysiological and cognitive changes occurring in aging. To this end, we report some of the neurophysiological substrates underpinning the perceptual and memory interactions in older adults, from the low and high perceptual processing to the conjunction in the medial temporal lobe. We then explore how these changes could explain more broadly the cognitive changes associated with aging in terms of losses and gains.

Keywords: embodiment, aging, neurocognition, perceptual processing, conjunction, distinctiveness

INTRODUCTION

Embodied cognition defines the body and the interaction with the world as shaping cognition and not just as simple inputs/outputs (Wilson, 2002). As a consequence, perceptual and motor systems should play a crucial role in cognitive functioning. Growing evidence, especially in young adults, has shown that sensorimotor components are at the core of language (Pulvermüller et al., 2005), attention (Bradley, 2007), memory (Versace et al., 2014) or action (Hommel, 2009). However, few studies are conducted in normal aging (Vallet, 2015). This is particularly surprising given that aging is marked by perceptual (and motor) decline in the one hand and by cognitive decline in the other hand. This perspective aims at proposing an embodied account of age-related cognitive decline focusing on perceptual and memory interactions in older adults. To this end, the neurophysiological substrates at the origin of the interactions between perceptual and memory should be understood. The low and high levels of sensory neurophysiological changes in older adults will be firstly described as their impact on the emergence of memory representations. Then, with the support of the hierarchical representational model (Murray and Bussey, 1999; Saksida and Bussey, 2010), we will examine how changes in the perceptual-mnemonic conjunctive processes, occurring in the medial temporal lobe (MTL), alter the emergence of representation. Finally, we will discuss how these changes could explain more broadly the cognitive changes associated with...
aging in terms of losses and gains based on the Activation-Integration model (Versace et al., 2009, 2014).

**SENSORY-PERCEPTUAL DECLINE AND LOW-RESOLUTION REPRESENTATION IN AGING**

Biological aging affects the whole body including, at a low-sensory level (sensory organs), many sensory modalities. Recently, the five Aristotelian senses (hearing, vision, taste, touch, and smell) was simultaneously assessed in older adults aged 57–85 years (Correia et al., 2016). The results showed that 74% of the participants had a deficit in identifying taste, 70% in touch, 22% in smell, 20% in corrected vision and 18% in corrected hearing. This study also provides for the first time an estimate of the proportion in which the sensory modalities are jointly altered. Two thirds of the participants had a deficit of two or more modalities, 27% had a deficit of only one of these modalities, while only 6% showed no impairment. At a higher level, aging worsens the transmission of sensory information from these organs to the brain (Ullhake et al., 2002) and higher perceptual thresholds are also found (e.g., Fozard and Gordon-Salant, 2001). At a cortical level, the occipital sensory cortex is less affected in aging with regard to structural integrity (e.g., Peters, 2006), but long-term peripheral sensory alteration may promote atrophy of the perceptual areas of the brain (Baltes and Lindenberger, 1997; Boucard et al., 2009; Golub, 2017). Functionally, the dopaminergic modulation deficit in aging, regulating the neurons sensitivity to related signals, reduces the functional specialization of neuronal activation. This is also true in neuronal circuits that are still relatively intact (Li et al., 2001), especially for visual stimuli in posterior regions (Park et al., 2004). The impoverished perceptual signal results in weakened unsensory and strengthen multisensory information processing (de Dieuleveult et al., 2017). As older adults exhibited reduced activity in occipital regions coupled with increased frontal activity, a functional compensation occurs (Davis et al., 2008).

Embodied cognition predicts that these perceptual changes should directly impact cognitive functioning. In this approach, all forms of knowledge (e.g., semantic, episodic) remains grounded in its sensorimotor components (Glenberg et al., 2013). The cognitive representations are not retrieved in memory, but instead emerge from the simulation of these components associated with the individual’s previous experiences based on the constraints of the present situation (Figure 1). Simulation refers here to the automatic and mandatory re-enactment of the brain activities of the perceptual, motor and emotional states produced by past experiences in the modal and heteromodal areas (Barsalou, 2008). Thus, the neurophysiological degradation occurring in aging in perceptual processing should deteriorate the simulation mechanism at the core of representation emergence (Vallet, 2015).

A less efficient simulation in older adults should lead to the emergence of “low-resolution” impoverished representations, that is, representations with lower details. In other words, the signal-to-noise ratio of the sensorimotor simulation should be lower, mainly due to a decrease in central perceptual processing. Whereas the perceptual discriminability in memory is underlyed by occipital regions regardless of age (Bowman et al., 2019), older adults exhibits reduced representation fidelity in these regions (Zheng et al., 2018). Impoverished sensory input may decrease the activation of specific sensory components of perceptually present information, but it should generally not alter the ability to simulate perceptually absent (and therefore mnemonic) information. Coherently, it has been shown that older adults did not suffer from a retrieval deficit, but instead exhibit less precise mnemonic representations of the items (i.e., less accurate responses on the color and orientation) and of the context (e.g., location) of the information to be learned (Korkki et al., 2020). This study also showed that the reduction in accuracy is not fully explained by a deficit in low-level sensory functioning alone (visual acuity), which may rather occur from neural dedifferentiation in the parahippocampus (Koen et al., 2019). The lower-resolution hypothesis could also account for associative (e.g., source memory) deficits in aging. Indeed, older adults have the greatest deficit in access associations requiring a high level of specificity (e.g., the old man was in this park), whereas they performed as well as younger adults to recognize general associations (e.g., the old man was in a park) and fuzzy associations (e.g., the old man was out somewhere) (Greene and Naveh-Benjamin, 2020). As all forms of knowledge (e.g., semantic, episodic, autobiographical) emerges from simulation, a consequence of this hypothesis is that memory deficits in older people should not be limited to newly learned knowledge. Coherently, older adults also recall less specific perceptual or spatiotemporal details in autobiographical memory tasks (very long-term memory see, Frankenberg et al., 2021). A better understanding of the changes requires to study the neurophysiological mechanisms underlying memory in the MTL (the hippocampus and surrounding cortex) and the effect of age on them.

**LOW-RESOLUTION REPRESENTATIONS INDUCE MORE INTERFERENCES IN AGING MEMORY**

In the MTL, the general coding principle is based on the functional theory of the hippocampus (e.g., Marr, 1971; Rolls, 2013). According to this theory, the emergence of specific memories (i.e., episodic memory) relies on the pattern separation (PS) and the pattern completion (PC) mechanisms. PS is defined as the ability to reduce interference from similar percepts by processing non-overlapping representations, whereas the PC allows recalling a whole and specific memory from an incomplete input signal by complementing (activating) the missing components. The input signals projected from the sensory cortex and then, the entorhinal cortex processed it into a non-overlapping pattern in the dentate gyrus via the mossy fiber (PS). This non-overlapping pattern is then projected as distinct representations into the CA3 field of the hippocampus, after from which the representation could be retrieved from CA3 by diffusing activation to the cortex (PC, Rolls, 2016; Pishdadian et al., 2020).
Modern approaches of neurocognition state that PS occurs beyond the hippocampus, all along the ventral stream (Kent et al., 2016; Cowell et al., 2019; Ekstrom and Yonelinas, 2020). These modern approaches emphasize the content of information to be processed focusing on the visual modality for the moment. All visual representations are, for instance, processed hierarchically from the simplest processing within the visual cortex to the most complex processing in the hippocampus (Murray and Bussey, 1999; Saksida and Bussey, 2010). This representational-hierarchical model states that non-overlapping representations (PS) could be obtained by the conjunction of perceptual features along the ventral stream regions to the hippocampus (see Kent et al., 2016). Basic sensory interference between individual stimulus characteristics (e.g., lines and colors) can be resolved in the sensory cortex. When more perceptually complex features are introduced (e.g., objects), the interference is resolved at higher processing level (e.g., the perirhinal cortex -PrC). Finally, combinatorial codes (e.g., conjunctive representations of object scenes in a spatial context) are resolved in the hippocampus.

The representations are assumed to be widely distributed into a system where characteristics of objects and scenes are progressively combined as the processing progress hierarchically. Moreover, and consistently with the embodiment, memory representation would emerge from the activation of perceptual units, and thus, the distinction between memory and perception are no more relevant (see also, Graham et al., 2010). Thus, conjunctions would not only reduce memory interference, but would also eliminate the ambiguity of visually similar stimuli in supposedly non-memory tasks. Accordingly, the PrC is involved in processing of complex perceptual objects (Buckley et al., 2001; Bussey et al., 2002; Barense et al., 2007) and lesions to the PrC induce false recognition due to interference from similar memories (Burke et al., 2010; McTighe et al., 2010). Similarly, growing evidence indicates that the hippocampus underlies the processing of conjunctions of complex spatial information in memory (Girardeau et al., 2009) and in presumed non-mnemonic tasks (see, Lee et al., 2012).

Applied in aging, the functionally weakened sensory signals in the sensory cortex would increase perceptual/mnemonic interference as conjunctions would be less efficient along the ventral stream. Indeed, and on the contrary to the occipital cortex, the MTL is structurally affected by aging, both in the PrC (Ryan et al., 2012; Fidalgo et al., 2016) and the hippocampus (Fraser et al., 2015). As a result, conjunction processing is expected to be impoverished in older adults. This should increase interference mainly between similar memories (Surprenant et al., 2006; Ekstrom and Yonelinas, 2020). Actually, when the memories are sufficiently distinct (efficient PS or dissimilar memories), then the PC mechanism can easily complete a specific trace, but when the memory traces overlap, then the system will enter into an unstable attractor that may lead to the emergence of altered/confused information (inefficient PC) (Ekstrom and Yonelinas, 2020; Zotow et al., 2020). Furthermore, the contribution of conjunctive representations to reduce interference could be more important as the delay increases. Delay-dependent impairments after MTL damage result from deficient conjunction representations to resolve the ambiguity of simpler representations in lower-level regions that are more likely to be encountered during delay (Yonelinas, 2013). Therefore, the
processing of similar information in aging could be impacted at low levels of similarity in long-term memory, while deficits would be evident at shorter time frames (e.g., short-term memory, perception) only at higher levels of similarity.

On a behavioral standpoint, less efficient PS in aging is mainly studied in memory using a perceptual lure discrimination index. In a study phase, the participant learns images. In a subsequent recognition task, these images (targets) are presented along with new images that are visually distinct (foils) or that are visually similar (lures) to the targets. The results show a linear decrease in perceptual lure discrimination as perceptual similarity increases in older adults (see Leal and Yassa, 2018). Older adults thus indicated more often having already seen a new image, especially when it is perceptually similar to a target (declining perceptual lure discrimination index). Similarly, false recognition was more likely for items in categories that are visually more similar than those that are more distinct (Boutet et al., 2019). It should be highlighted that the perceptual lure discrimination index is associated with more global cognitive functioning in older adults (Pishdadian et al., 2020). As such, the alteration of the conjunction processing in the MTL could have wider consequences than the specific memories.

DISTINCTIVENESS OF MEMORY TRACES ON OTHER COGNITIVE DOMAINS

The consequence of reduced sensory processing in aging should not be limited to memory according to embodiment. According to the embodied and situated memory models Act-In (Activation-Integration, Versace et al., 2014), representations (e.g., semantic, episodic) emerge from the same sensorimotor components of the different memory traces. All experiences of the individual are supposed to be accumulated as memory traces. These traces are distributed across modal and heteromodal neuronal systems coding the multiple sensorimotor components of the experiences.

The different components of a given memory trace are bound together (Opitz, 2010), following the conjunction processing and therefore the PS. The binding allows the PC, described in Act-In as an intra-trace activation. A specific memory emerges when the activation does not propagate to similar traces (called inter-trace activation). Reversely, the activation of multiple similar traces (inter-trace activation) should produce categorial (non-specific/semantic) knowledge. A strong inter-trace diffusion achieves categorization by eliminating specific details and context of events (see Versace et al., 2009, 2014), mechanism underlined by the CA1 subregion of the hippocampus and neocortical upstream (Kumaran and McClelland, 2012). The intra-trace and inter-trace activations are mutually repulsive so strengthen or weaken one kind of activation should directly weaken or strengthen the other. Given that intra-trace activation is facilitated by distinctiveness, better simulations (higher resolution representations) should allow more activation of contextual details limiting the activation of common/similar components of other traces (inter-trace activation) (e.g., Ekstrom and Yonelinas, 2020). Furthermore, distinctiveness is also a function of the number of experiences accumulated, then more traces should increase the likelihood of their overlap. This, in turn, should increase the inter-trace activation. Consequently, the decline of perceptive and conjunctive processing associated with more traces (more events experienced by older adults) should induce less distinct traces in older adults (Vallet, 2015), then it should in return bias the dynamics of the simulation in favor of the inter-trace activation (less distinct processing).

This balance between specific and non-specific knowledge could be illustrated by the fact that older adults produce fewer internal (specific) details associated with more external (categorial) details when they recall an event compared to young adults (e.g., Levine et al., 2002). The effect of aging on the visual cortex and the hippocampus reduces specific details (neural dedifferentiation), while aging enhances categorial representations in the anterior temporal lobe (neural hyperdifferentiation) (Deng et al., 2021). Therefore, older adults exhibit relatively well-preserved performance in semantic tasks (see, Salthouse, 2010, for a review). However, a more qualitative analysis shows that they produce more categorial (e.g., cat) and less unique (e.g., botfly) semantic knowledge (Murphy and Naveh-Benjamin, 2020; Frankenberg et al., 2021), but they also less benefit from distinctiveness (see Smith, 2006) due to a more generic (categorial) and less distinct processing (Koutstaal and Schacter, 1997; Smith, 2006).

Aging might be characterized by an imbalance toward generic (non-specific) processing constrained by the weight of prior knowledge at the expense of specific processing. Compared to young adults, older people produced indeed less specific details and more generic details in a basic image description task and in future imagination task (Gaesser et al., 2011; Schacter et al., 2013). Yet, an integrated view of neurocognitive functioning suggests that deficits in modal (e.g., less specialization in occipital processing) and heteromodal (e.g., structural impairment of MTL) regions should alter the processing done within the connected regions as the prefrontal cortex (Davis et al., 2008). The neuromodulation deficit induces noise in the neural processing, including the prefrontal cortex, and leads to less specific and more general processing (Li et al., 2001). As the processes are less specific, and the similarity of the previous processes favors the automation of processes (see Logan, 1988), new processes are less likely to emerge. The same over-repeated processes will more likely emerge, making more rigid and less flexible other processing. This rigidity is found for their executive functions such as to change categorization rules in the Wisconsin tasks (e.g., Daigneault et al., 1992; Ashendorf and McCaffrey, 2008). Interestingly, the largest executive switching costs were found under conditions of ambiguous sensory stimuli and overlap between sets of responses (Mayr, 2001). This is consistent with the hypothesis that perceptual deficits
reduce trace distinctiveness and lead to increased inter-trace activation, requiring more inhibition in prefrontal cortex (Li et al., 2001). Similar link between sensory functioning and inhibition is observed in young adults with degraded vision (e.g., cataract vision simulation, as found in pathological visual aging) in the Stroop task (see Monge and Madden, 2016 for a review). A more evidence comes from the Perceptual Lure Discrimination Index in the Mnemonic Similarity Task that is related to inhibition (Foster and Giovanelli, 2020). Finally, it is noteworthy that the MoCA (global measure of cognitive aging) were associated with lure discrimination performances in older adults (Pishdadian et al., 2020).

CONCLUSION

Normal aging is characterized by sensory-perceptual (and motor) decline, on the one hand, and cognitive decline on the other. Further studies, using a longitudinal design, are required to fully explore the progressive sensory (motor) and cognitive changes occurring throughout life. Embodied cognition provides a theoretical framework explaining the possible these links given that any representation at the source of cognitive functioning remains grounded in these sensorimotor components. As such, memory and perception (and action) are functionally equivalent. This perspective investigated the neurophysiological mechanisms underlying these links. The sensory decline (organ level) should have a minimal impact, mainly on overlapping stimuli by impoverishment of the related signal. Higher perceptual decline should affect the simulation mechanism leading to the emergence of a less specific and detailed representation. Functional changes in the primary perceptual areas may reduce the benefit of distinct perceptual information, while structural changes in the MTL may reinforce of overlapping perceptual and memory information. Since embodied representation should be at the core of cognition, such changes should have wider cognitive consequences than memory. Thus, aging could be characterized by less specific and more rigid processing.

This perspective highlights the interest to study aging in an embodied cognition approach, which could represent an alternative to other theories of cognitive aging due to how the sensory (motor)-cognitive interactions are defined. The focus of the present article on perceptual-memory interactions also suggest that early sensory improvement and environmental enrichment could improve cognitive aging (Leon and Woo, 2018). Similar effects should also be found with motor and action interactions. We hope that this brief overview of the contribution of embodied cognition to characterize neurocognitive aging will encourage further investigation of cognitive functions in aging from an embodied perspective.

DATA AVAILABILITY STATEMENT

The original contributions presented in the study are included in the article/supplementary material, further inquiries can be directed to the corresponding author(s).

AUTHOR CONTRIBUTIONS

JM: conceptualization, writing – original draft, and project administration. GV: conceptualization, writing – review and editing, supervision, funding acquisition, and resources. SB: writing – review and editing. MI: writing – review and editing, and supervision. All authors contributed to the article and approved the submitted version.

FUNDING

JM and GV are supported by a grant from the Auvergne-Rhône-Alpes region for the project Vieillissement, Maladie Chronique et Stimulation Cognitive (ViMaCC). The ViMaCC project was co-financed by the European Union within the framework of the Fonds européen de développement régional (FEDER). MI was supported by the FACTOLAB framework, sponsored by Michelin Tyres Manufacturer and by the French government research programs “Investissements d’Avenir” through the IDEX-ISITE initiative 16-IDEX-0001 (CAP 20-25) and the IMobS3 Laboratory of Excellence (ANR-10-LABX-16-01).

REFERENCES

Ashendorf, L., and McCaffrey, R. J. (2008). Exploring age-related decline on the Wisconsin card sorting test. Clin. Neuropsychol. 22, 262–272.

Baltes, P. B., and Lindenberger, U. (1997). Emergence of a powerful connection between sensory and cognitive functions across the adult life span: a new window to the study of cognitive aging? Psychol. Aging 12, 12–21. doi: 10.1037/0882-7974.12.1.12

Barense, M. D., Gaffan, D., and Graham, K. S. (2007). The human medial temporal lobe processes online representations of complex objects. Neuropsychologia 45, 2963–2974. doi: 10.1016/j.neuropsychologia.2007.05.023

Barsalou, L. W. (2008). Grounded cognition. Annu. Rev. Psychol. 59, 617–645. doi: 10.1146/annurev.psych.59.103006.093659

Boucard, C. C., Hernowo, A. T., Maguire, R. P., Jansonius, N. M., Roerdink, J. B. T. M., Hooymans, J. M., et al. (2009). Changes in cortical grey matter density associated with long-standing retinal visual field defects. Brain 132, 1898–1906. doi: 10.1093/brain/aws119

Boutet, I., Dawod, K., Chiasson, F., Brown, O., and Collin, C. (2019). Perceptual similarity can drive age-related elevation of false recognition. Front. Psychol. 10:743. doi: 10.3389/fpsyg.2019.00743

Bowler, S. N., Wallace, J. L., Nematollahi, S., Uprety, A. R., and Barnes, C. A. (2010). Pattern separation deficits may contribute to age-associated recognition impairments. Behav. Neurosci. 124, 559–573. doi: 10.1037/a0020993

Buckley, M. J., Booth, M. C., Rolls, E. T., and Gaffan, D. (2001). Selective perceptual impairments after perirhinal cortex ablation. J. Neurosci. 21, 9824–9836. doi: 10.1523/JNEUROSCI.21-24.2001

Burke, S. N., Wallace, J. L., Nematollahi, S., Uprety, A. R., and Barnes, C. A. (2010). Pattern separation deficits may contribute to age-associated recognition impairments. Behav. Neurosci. 124, 559–573. doi: 10.1037/a0020993

Mille et al.
Mille et al. Embodied Perspective on Neurocognitive Aging

Bussey, T. J., Sakaida, L. M., and Murray, E. A. (2002). Perirhinal cortex resolves feature ambiguity in complex visual discriminations. *Eur. J. Neurosci.* 15, 365–374. doi: 10.1046/j.1460-9552.2001.01081.x

Correia, C., Lopez, K. J., Wroblewski, K. E., Huising-Scheetz, M., Kern, D. W., Chen, R. C., et al. (2016). Global sensory impairment in older adults in the United States. *J. Am. Geriatr. Soc.* 64, 306–313. doi: 10.1111/jgs.13955

Cowell, R. A., Barense, M. D., and Sall, P. S. (2019). A roadmap for understanding memory: decomposing cognitive processes into operations and representations. *Eur. J. Neurosci.* 6: ENEURO.0122-19.2019. doi: 10.1111/euro.12219

Daigneault, S., Braun, C. M., and Whitaker, H. A. (1992). Early effects of

Davis, S. W., Dennis, N. A., Daselaar, S. M., Fleck, M. S., and Cabeza, R. (2008).

Deng, L., Davis, S. W., Monge, Z. A., Wing, E. A., Geib, B. R., Raghunandan,

Hommel, B. (2009). Action control according to TEC (theory of event coding).

Girardeau, G., Benchenane, K., Wiener, S. I., Buzsáki, G., and Zugaro, M. B. (2009).

Fozard, J. L., and Gordon-Salant, S. (2001). “Changes in vision and hearing with

Foster, C. M., and Giovanello, K. S. (2020). Domain general processes moderate

Ekstrom, A. D., and Yonelinas, A. P. (2020). Precision, binding, and

Fraser, M. A., Shaw, M. E., and Cherbuin, N. (2015). A systematic review and

Glenberg, A. M., Witt, J. K., and Metcalfe, J. (2013). From the revolution to

Graham, K. S., Barense, M. D., and Lee, A. C. H. (2010). Neuropsychologia

Glenberg, A. M., Witt, J. K., and Metcalfe, J. (2013). From the revolution to

Haider, G., Benchenane, K., Wiener, S. I., Buzsáki, G., and Zugaro, M. B. (2009).

Hommel, B. (2009). Action control according to TEC (theory of event coding).

Koutstaal, W., and Schacter, D. L. (1997). Gist-based false recognition of pictures in older and younger adults. *J. Mem. Lang.* 39, 555–583. doi: 10.1006/jmla.1997.2529

Kumaran, D., and McClelland, J. L. (2012). Generalization through the recurrent interaction of episodic memories: a model of the hippocampal system. *Psychol. Rev.* 119, 573–616. doi: 10.1037/a0028681

Lee, S. L., and Yassa, M. A. (2018). Integrating new findings and examining clinical applications of pattern separation. *Nat. Neurosci.* 21, 163–173. doi: 10.1038/s41593-017-0065-5

Lee, A. C., Yeung, L. K., and Barense, M. D. (2012). The hippocampus and visual perception. *Front. Hum. Neurosci.* 6: 91. doi: 10.3389/fnhum.2012.00091

Leon, M., and Woo, C. (2018). Environmental enrichment and successful aging. *Front. Behav. Neurosci.* 12:155. doi: 10.3389/fnbeh.2018.00155

Levine, B., Svoboda, E., Hay, J. F., Winocur, G., and Moscovitch, M. (2002). Aging and autobiographical memory: dissociating episodic from semantic retrieval. *Psychol. Aging* 17, 677–689. doi: 10.1037/0882-7974.17.4.677

Li, S. C., Lindenberger, U., and Säktström, S. (2001). Aging cognition: from neuromodulation to representation. *Trends Cogn. Sci.* 5, 479–486. doi: 10.1016/S1364-6613(00)01769-1

Logan, G. D. (1988). Toward an instance theory of automatization. *Psychol. Rev.* 95, 492–527. doi: 10.1037/0033-295X.95.4.492

Marr, D. (1971). Simple memory: a theory for archicortex. *Philos. Trans. R. Soc. B* 262, 23–81. doi: 10.1098/rstb.1971.0078

Mayr, U. (2001). Age differences in the selection of mental sets: the role of inhibition, stimulus ambiguity, and response-set overlap. *Psychol. Aging* 16, 96–109.

McTighe, S. M., Cowell, R. A., Winters, B. D., Bussey, T. J., and Sakaida, L. M. (2010). Paradoxical false memory for objects after brain damage. *Science* 330, 1408–1410. doi: 10.1126/science.1194780

Monge, Z. A., and Madden, D. J. (2016). Linking cognitive and visual perceptual decline in healthy aging: the information degradation hypothesis. *Neurosci. Biobehav. Rev.* 69, 166–173. doi: 10.1016/j.neubiorev.2016.07.031

Murphy, D. H., and Castel, A. D. (2020). Age-related similarities and differences in the components of semantic fluency: analyzing the originality and organization of retrieval from long-term memory. *Aging Neuropsychol. Cogn.* 1–14. doi: 10.1080/13825585.2020.1817844

Murray, E. A., and Bussey, T. J. (1999). Perceptual–mnemonic functions of the perirhinal cortex. *Trends Cogn. Sci.* 3, 142–151. doi: 10.1016/S1364-6613(99)01303-0

Optiz, B. (2010). Neural binding mechanisms in learning and memory. *Neurosci. Biobehav. Rev.* 34, 1036–1046. doi: 10.1016/j.neubiorev.2009.11.001

Park, D. C., Polk, T. A., Park, R., Minear, M., Savage, A., and Smith, M. R. (2004). Aging reduces neural specialization in ventral visual cortex. *Proc. Natl. Acad. Sci. U.S.A.* 101, 13091–13095. doi: 10.1073/pnas.0405148101

Peters, R. (2006). Ageing and the brain. *Postgrad. Med. J.* 82, 84–88. doi: 10.1136/pgmj.2005.036665

Pishdadian, S., Houng, N. V., Baker, S., Moscovitch, M., and Rosenbaum, R. S. (2020). Not only memory: investigating the sensitivity and specificity of the Mnesonic Similarity Task in older adults. *Neuropsychologia* 149,107670. doi: 10.1016/j.neuropsychologia.2020.107670

Pulvermüller, F., Hauk, O., Nikulin, V. V., and Ilmoniemi, R. J. (2005). Functional links between motor and language systems. *Eur. J. Neurosci.* 21, 793–797. doi: 10.1111/j.1460-9586.2005.03900.x

Rolls, E. T. (2013). The mechanisms for pattern completion and pattern separation in the hippocampus. *Front. Syst. Neurosci.* 7:74. doi: 10.3389/fnsys.2013.00074

Rolls, E. T. (2016). Pattern separation, completion, and categorisation in the hippocampus and neocortex. *Neuropsychol. Learn. Mem.* 12, 4–28. doi: 10.1037/lnm0000038

Ryan, L., Cardoza, J. A., Barense, M. D., Kawa, K. H., Wallentin-Flores, J., Arnold, W. T., et al. (2012). Age-related impairment in a complex object discrimination
task that engages perirhinal cortex. *Hippocampus* 22, 1978–1989. doi: 10.1002/hipo.22069

Saksida, L. M., and Bussey, T. J. (2010). The representational–hierarchical view of amnesia: translation from animal to human. *Neuropsychologia* 48, 2370–2384. doi: 10.1016/j.neuropsychologia.2010.02.026

Salthouse, T. A. (2010). Selective review of cognitive aging. *J. Int. Neuropsychol. Soc.* 16, 754–760. doi: 10.1017/S1355617710000706

Schacter, D. L., Gaesser, B., and Addis, D. R. (2013). Remembering the past and imagining the future in the elderly. *Gerontology* 59, 143–151. doi: 10.1159/000342198

Smith, R. E. (2006). "Adult age differences in episodic memory: item-specific, relational, and distinctive processing," in *Distinctiveness and Memory*, eds R. R. Hunt and J. B. Worthen (Oxford: Oxford University Press), 259–287. doi: 10.1093/acprof:oso/9780195169669.003.0012

Surprenant, A. M., Neath, I., and Brown, G. D. A. (2006). Modeling age-related differences in immediate memory using SIMPLE. *J. Mem. Lang.* 55, 572–586. doi: 10.1016/j.jml.2006.08.001

Ulfhake, B., Bergman, E., and Fundin, B. T. (2002). Impairment of peripheral sensory innervation in senescence. *Auton. Neurosci.* 96, 43–49. doi: 10.1016/S1566-0702(01)00368-X

Vallet, G. T. (2015). Embodied cognition of aging. *Front. Psychol.* 6:463. doi: 10.3389/fpsyg.2015.00463

Versace, R., Labeye, É, Badard, G., and Rose, M. (2009). The contents of long-term memory and the emergence of knowledge. *Eur. J. Cogn. Psychol.* 21, 280–306. doi: 10.1080/09541440801951844

Versace, R., Vallet, G. T., Riou, B., Lesourd, M., Labeye, É, and Brunel, L. (2014). Act-In: an integrated view of memory mechanisms. *J. Cogn. Psychol.* 26, 280–306. doi: 10.1080/20445911.2014.892113

Wilson, M. (2002). Six views of embodied cognition. *Psychon. Bull. Rev.* 9, 625–636.

Yonelinas, A. P. (2013). The hippocampus supports high-resolution binding in the service of perception, working memory and long-term memory. *Behav. Brain Res.* 254, 34–44. doi: 10.1016/j.bbr.2013.05.030

Zheng, L., Gao, Z., Xiao, X., Ye, Z., Chen, C., and Xue, G. (2018). Reduced fidelity of neural representation underlies episodic memory decline in normal aging. *Cereb. Cortex* 28, 2283–2296. doi: 10.1093/cercor/bhx130

Zotow, E., Bisby, J. A., and Burgess, N. (2020). Behavioral evidence for pattern separation in human episodic memory. *Learn. Mem.* 27, 301–309. doi: 10.1101/LM.051821.120

Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

Publisher's Note: All claims expressed in this article are solely those of the authors and do not necessarily represent those of their affiliated organizations, or those of the publisher, the editors and the reviewers. Any product that may be evaluated in this article, or claim that may be made by its manufacturer, is not guaranteed or endorsed by the publisher.

Copyright © 2021 Mille, Brambati, Izaute and Vallet. This is an open-access article distributed under the terms of the Creative Commons Attribution License (CC BY). The use, distribution or reproduction in other forums is permitted, provided the original author(s) and the copyright owner(s) are credited and that the original publication in this journal is cited, in accordance with accepted academic practice. No use, distribution or reproduction is permitted which does not comply with these terms.