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Electrophysiological evidence for a sensory recruitment model of somatosensory working memory

Sensory recruitment in tactile working memory

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Abstract

Sensory recruitment models of working memory assume that information storage is mediated by the same cortical areas that are responsible for the perceptual processing of sensory signals. To test this assumption, we measured somatosensory event-related brain potentials (ERPs) during a tactile delayed match-to-sample task. Participants memorized a tactile sample set at one task-relevant hand to compare it with a subsequent test set on the same hand. During the retention period, a sustained negativity (tactile contralateral delay activity, tCDA) was elicited over primary somatosensory cortex contralateral to the relevant hand. The amplitude of this component increased with memory load and was sensitive to individual limitations in memory capacity, suggesting that the tCDA reflects the maintenance of tactile information in somatosensory working memory. The tCDA was preceded by a transient negativity (N2cc component) with a similar contralateral scalp distribution, which is likely to reflect selection of task-relevant tactile stimuli at the encoding stage. The temporal sequence of N2cc and tCDA components mirrors previous observations from ERP studies of working memory in vision. The finding that the sustained somatosensory delay period activity varies as a function of memory load supports a sensory recruitment model for spatial working memory in touch.

Introduction

Working memory (WM) is responsible for the active maintenance of information that is no longer perceptually present. Visual and tactile working memory are both assumed to be based on distributed neural networks that include prefrontal
cortex (PFC) and modality-specific perceptual areas. The activation of PFC during the maintenance of visual and tactile stimuli in working memory is well established (Curtis and D'Esposito 2003; Curtis, Rao, D'Esposito 2004; Fuster and Alexander 1971; Kostopoulos, Albanese, Petrides 2007; Romo and Salinas 2003; Postle 2005). Additionally, modality-specific visual (Harrison and Tong 2009; Supèr, Spekreijse, Lamme 2001) or somatosensory areas (e.g., Kaas et al. 2013; Zhou and Fuster 1996) show persistent activation during the retention of visual or tactile stimuli. Although the exact role of this delay-period activity in visual areas during working memory maintenance and their link to selective visual attention are still debated (e.g., van Dijk et al. 2010; Lewis-Peacock et al. 2012; Postle et al. 2013), its existence has led to the “sensory recruitment” model of working memory (D'Esposito 2007; Harrison and Tong 2009; Pasternak and Greenlee 2005; Postle 2006). This model postulates that perceptual brain regions which are responsible for the sensory processing of visual or tactile stimuli are also involved in working memory storage. The sustained activation of perceptual areas might be particularly important when working memory tasks require the maintenance of detailed sensory information (e.g., Lee, Kravitz, Baker 2013; see also Sreenivasan, Curtis, D'Esposito 2014).

Support for the sensory recruitment model comes from ERP studies of visual working memory (e.g., Vogel, McCollough, Machizawa 2005; Vogel and Machizawa 2004). In these studies, bilateral sample displays were followed after a retention interval by test displays, and participants had to match sample and test objects on one side of these displays. A sustained negativity at posterior electrodes contralateral to the side of the memorized objects (contralateral delay activity, CDA) started 300 ms after sample onset and persisted throughout the retention interval. The fact that this CDA component is sensitive to manipulations of visual working memory load and to individual differences in working memory capacity strongly suggests that the CDA
directly reflects the maintenance of visual information in working memory. The contralateral nature and posterior scalp topography of the CDA is consistent with neural generators in extrastriate visual areas (McCollough, Machizawa, Vogel 2007), in line with the sensory recruitment model. The CDA is typically preceded by an N2pc component that emerges around 200 ms post-stimulus, has a similar posterior scalp topography (e.g. McCollough, Machizawa, Vogel 2007), and reflects the attentional selection and encoding of task-relevant objects in ventral visual cortex (Eimer 1996; Luck and Hillyard 1994).

While ERP markers of visual working memory are well established, corresponding electrophysiological correlates of tactile working memory have not yet been described. Here, we demonstrate the existence of two somatosensory ERP components that are elicited during the encoding and maintenance of tactile stimuli in working memory, and both show modality-specific topographies over primary somatosensory cortex. We employed a task that was modelled on the delayed match-to-sample task used in earlier studies of visual working memory (e.g., Vogel, McCollough, Machizawa 2005; Vogel and Machizawa 2004). On each trial, a set of tactile sample stimuli was followed after a 2000 ms retention period by tactile test stimuli. Sample and test stimuli were delivered simultaneously to both hands, but the memory task had to be performed for one of these hands only. Participants had to encode and maintain tactile sample stimuli on the currently task-relevant hand, and to match them to subsequent test stimuli on the same relevant hand. On low-load trials, a single tactile stimulus had to be maintained and matched. On high-load trials, two tactile pulses had to be memorized.

Results revealed the existence of two somatosensory ERP components that have not yet been described in the literature on tactile attention and working memory. During the retention interval, a sustained tactile contralateral delay activity (tCDA)
emerged with a modality-specific scalp distribution over somatosensory areas. This tCDA component was sensitive to memory load and to individual differences in tactile working memory capacity. It was preceded by a central contralateral negativity (N2cc component) with a similar modality-specific topography that was also modulated by working memory load. Analogous to the visual N2pc and CDA, these N2cc and tCDA components reflect the spatially selective encoding and maintenance of task-relevant information in tactile working memory.

Materials and Methods

Participants

Eighteen neurologically unimpaired paid adult participants were tested. The study was conducted in accordance with the Declaration of Helsinki, and was approved by the Psychology Ethics Committee, Birkbeck College. All participants gave informed written consent prior to testing. Two participants were excluded from analysis because their tactile WM capacity measured by Cowan’s K (Cowan 2001) was below 1. Sixteen participants remained in the sample (mean age 32 years, range 25-44 years, 3 male, 13 right-handed).

Stimuli and task design

Participants were seated in a dimly lit recording chamber, viewing a monitor showing a central white fixation cross against a black background. Both hands were covered from sight and were placed on a table at a distance of approximately 40 cm. Eight mechanical tactile stimulators were attached to the distal phalanges of the index, middle, ring and small fingers of the left and right hand. Stimulators were driven by an eight-channel sound card (M-Audio, Delta 1010LT) and custom-built amplifiers, controlled by Matlab (MathWorks, Natick, MA). Continuous white noise
was delivered via headphones to mask sounds produced by the tactile stimulators. All tactile stimuli were 100 Hz sinusoids (duration: 200 ms; intensity: 0.37 N).

Figure 1 illustrates the experimental procedure. Each trial started with a set of tactile sample stimuli that were delivered simultaneously to the left and right hand. After a 2000 ms retention period, a set of tactile test stimuli was presented simultaneously to both hands. Prior to the start of each block, instructions displayed on the monitor informed participants whether the left or right hand was relevant in the upcoming block. Participants had to decide whether sample and test stimulus locations on this hand were identical (match trials) or different (mismatch trials). The task-relevant hand was swapped after each experimental block. Two load conditions were randomized within each block. In the low-load condition, one sample pulse was presented with equal probability to one of the four fingers of the left hand and the right hand. On match trials, the test pulse was delivered to the same finger of the relevant hand as the sample pulse. On mismatch trials, one of the three other fingers on that hand was stimulated at test. In the high-load condition, two sample pulses were presented to two randomly selected fingers of the left hand and the right hand, respectively. On match trials, test pulses were delivered to the same two fingers of the relevant hand. On mismatch trials, at least one of the two test pulses was presented to a different finger of that hand. Because one of the two sample locations could be repeated at test on mismatch trials, participants had to retain the location of both sample stimuli on the relevant hand to perform the task in the high-load condition. Match and mismatch trials were equiprobable. On the currently task-irrelevant hand, sample and test stimuli were also presented at matching or mismatching locations, and this was independent of whether there was a match or mismatch on the relevant hand.
Participants signalled a match or mismatch between sample and test on the relevant hand with a vocal response ("a" for match and "e" for mismatch) that was recorded with a headset microphone between 200 ms and 1700 ms after test stimulus offset. A question mark replaced the fixation cross on the monitor during this period. The interval between the offset of this question mark and the onset of the sample pulses on the next trial varied between 800 and 1100 ms. The experiment included ten blocks of 48 trials, with twelve trials per block for each of the four combinations of high versus low load trials and match versus mismatch trials. Instructions emphasized accuracy over speed, and the need to avoid head and arm movements and to maintain central gaze fixation. Feedback on hit and correct rejection rates was provided after each block. Two training blocks were run prior to the first experimental block.

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Insert Figure 1 about here

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**Processing of EEG data**

EEG data, sampled at 500 Hz using a BrainVision amplifier, were DC-recorded from 64 Ag/AgCl active electrodes at standard locations of the extended 10-20 system. Two electrodes at the outer canthi of the eyes monitored lateral eye movements (horizontal electrooculogram, HEOG) and electrodes sites TP9/10 were used as mastoid references. Continuous EEG data was referenced to the left mastoid during recording, and was offline re-referenced to the arithmetic mean of both mastoids and submitted to a 40Hz low-pass finite impulse response filter (Blackman
window, filter order 666). EEG epochs for the 2000 ms interval following sample stimulus onset were computed relative to a 200 ms pre-stimulus baseline. Blind source separation of EEG data was performed with the Independent Component Analysis (ICA) algorithm implemented in the EEGLab toolbox (Delorme and Makeig 2004; Delorme, Sejnowski, Makeig 2007). Independent components related to artifacts at anterior scalp regions (in particular, eye movements and blinks), were identified by visual inspection and subtracted from the EEG data. To obtain reliable ICA decompositions, a copy of the data was segmented into eight 250 ms frames covering the 2000 ms retention period. These frames were corrected using whole-epoch baselines to achieve data stationarity (cf., Groppe, Makeig, Kutas 2009) without high-pass filtering, which would have removed slow brain potentials. The copy was discarded after ICA decompositions had been applied to the original data set. Epochs with lateral eye movements that escaped ICA artifact correction were identified and removed with a differential step function on the bipolarized HEOG (step width 100 ms, threshold 24 µV). The resulting HEOG waveforms contained no systematic eye gaze deflections towards the task-relevant hand (Figure 2, bottom panel). After artifact rejection and elimination of trials with incorrect responses, 90.2% of all epochs were retained for statistical analyses (low load: 93.4%; high load: 87.1%).

Results

Behavioral performance

Participants responded correctly on 97.1% of all low-load trials and 90.4% of all high-load trials. Sensitivity indices (d') were analysed in a two-way repeated measures ANOVA with the factors memory load (low, high) and relevant hand (left, right). Performance was reduced with high load relative to low load ($F(1,15) = 71.728$,
and did not differ between blocks where the left or right hand was relevant ($F(1,15) = 1.081, p > 0.3$). A significant memory load x relevant hand interaction ($F(1,15) = 6.222, p = 0.025$) was due to the fact that the performance decrement with high as compared to low memory load was larger when the memory task was performed with the left hand (8.5%) relative to blocks where the right hand was relevant (4.9%).

Mean vocal reaction times (RTs) in trials with correct responses were faster in the low-load relative to the high-load condition (799 ms versus 817 ms; main effect of memory load: $F(1,15) = 8.801, p = 0.010$). RTs did not differ between blocks where the left or right hand was task-relevant ($F(1, 15) = 1.846, p > 0.1$). The memory load x relevant hand interaction was significant ($F(1,15) = 5.25, p = 0.037$), as the RT costs for the low-load versus high-load condition were larger when the memory task was performed with the right hand (26 ms) relative to blocks where the left hand was relevant (10 ms). In other words, there was an asymmetric speed-accuracy tradeoff between the two hands for task performance in the high-load versus low-load condition.

Electrophysiological data

Figure 2 shows ERP waveforms averaged across lateral central electrodes (FC3/4, FC5/6, C3/4, C5/6, CP3/4, CP5/6) contralateral and ipsilateral to the task-relevant hand for the 2000 ms interval between the bilateral sample stimulus and the subsequent test stimulus. Results are shown separately for the low-load and high-load conditions. Following the early sensory-evoked ERP components to the sample stimulus, ERP waveforms were characterized by a gradually developing sustained negativity that reached its maximal amplitude immediately before the onset of the test.
stimuli. This sustained negativity that was present at contralateral as well as ipsilateral electrodes reflects the Contingent Negative Variation (CNV; see Birbaumer et al. 1990) that is elicited in anticipation of expected task-relevant events such as the test stimulus set used in this study. More importantly, sample stimuli triggered a transient enhanced negativity contralateral to the task-relevant hand. This N2cc component emerged around 180 ms after sample stimulus onset, and its amplitude was larger in the high-load as compared to the low-load condition. The N2cc was followed by a sustained contralateral negativity (tCDA) that remained present throughout the retention period. This tCDA component was larger when two tactile stimuli had to be memorized relative to the low load condition. The topographical maps in Figure 2 illustrate the scalp distribution of N2cc and tCDA components in the low-load and high-load conditions. Data shown in these maps were collapsed across blocks where the left or right hand was task-relevant by flipping ERPs at contralateral electrodes in blocks with a left-hand memory task over the midline. Both N2cc and tCDA components were maximal over somatosensory areas in the postcentral gyrus and adjacent parietal regions (see also Figure 4 below).

Difference waveforms were computed by subtracting ERPs ipsilateral to the currently task-relevant hand from contralateral ERPs. Statistical tests were conducted on mean amplitudes of these difference waves for a time window centered on the N2cc component (180-260 ms post-stimulus), and a second window centered on the
tCDA (300-2000 ms). Difference values that statistically differ from zero mark the presence of reliable lateraled components in the ERP waveforms. The N2cc was present in both the low-load ($t(15) = -5.593, p < 10^{-4}$) and high-load condition ($t(15) = -7.037, p < 10^{-5}$). N2cc amplitudes were significantly larger with high relative to low memory load ($t(15) = 4.235, p < 10^{-3}$). The tCDA component was present with low load ($t(15) = -2.951, p = 0.010$) as well as with high memory load ($t(15) = -6.126, p < 10^{-4}$). Similar to the N2cc, tCDA amplitudes were significantly larger in the high-load relative to the low-load condition ($t(15) = 3.801, p = 0.002$).

An additional analysis of mean amplitudes in the tCDA time window obtained for the unsubtracted ERP waveforms revealed a main effect of contralaterality (electrodes contralateral versus ipsilateral to the task-relevant hand; $F(1,15) = 38.006, p < 10^{-4}$) that interacted with load ($F(1,15) = 14.448, p = 0.002$), due to the fact that the tCDA was larger in the high-load condition. There was also a main effect of load ($F(1,15) = 14.862, p=0.002$), with larger CNV components with high memory load. This load-dependent modulation of CNV amplitudes was reliable at contralateral as well as ipsilateral electrodes $t(15) = -4.500$ and $-2.481, p < 0.001$ and 0.026, respectively).

Tactile working memory capacity was calculated for each individual participant on the basis of their performance in the high-load condition, using the formula $K = (\text{hits} + \text{correct rejections} -1) \times 2$, where 2 denotes memory set size in this condition (Cowan 2001). As illustrated in Figure 3, individual memory capacity was reliably correlated with the difference of tCDA amplitudes between the high-load and low-load conditions ($r = -0.640, p = 0.008$). Participants with higher tactile working memory capacity showed a more pronounced increase of the tCDA component on trials with high versus low memory load than participants with lower capacity. No correlation
was found between individual K values and the difference of N2cc amplitudes between high- and low-load conditions ($p > 0.7$).

To obtain additional evidence for a link between tCDA amplitudes and behavioral performance at the level of individual trials in the high-load condition, we computed tCDA components in the high-load condition separately for trials with vocal RTs above and below the median RT (with RT median splits conducted individually for each participant and trial condition). Trials with fast responses were more accurate than slow response trials (Cowan’s K: fast = 1.786, slow = 1.453; $t(15) = 6.362$, $p < 10^{-4}$). Critically, tCDA amplitudes were larger for fast as compared to slow response trials (-0.749 µV versus -0.594 µV), and this amplitude difference was significant ($t(15) = -2.564$, $p = 0.022$).

An additional current source density (CSD) analysis was conducted to further illustrate the modality-specific scalp topographies of the N2cc and tCDA components, and to demonstrate that the selection of lateral central electrodes for the analysis of these components was appropriate. ERP data were collapsed across the low- and high-load conditions, after conversion of scalp potentials to surface Laplacians ($\lambda = 10^{-5}$, iterations = 50, $m = 4$; cf. Tenke and Kayser 2012). This transformation minimizes effects of volume conduction from remote sources, and leads to a reference-independent representation of EEG/ERP data. CSD topographies provide a conservative estimate of the neural generator patterns that
contribute to scalp-recorded ERPs (Nunez and Westdorp 1994; Tenke and Kayser 2012). Robust lateralized effects were found over somatosensory brain regions (Figure 4), as demonstrated by significant differences of contra- minus ipsilateral activity recorded at central electrodes in the time window of N2cc ($t(15) = -6.476, p < 10^{-4}$) and tCDA ($t(15) = -4.066, p = 0.001$). Apart from an almost significant contralateral positivity at anterior regions during the N2cc time window ($t(15) = 2.107, p = 0.052$), no statistically reliable lateralization was evident over posterior (electrodes P3/4, P5/6, PO3/4, PO7/8) and anterior (electrodes AF3/4, AF7/8, F3/4, F5/6) scalp regions (all ps > 0.2; see Figure 4).

Discussion

We employed a tactile memory task that was modelled on the delayed match-to-sample task used in previous research on visual working memory (e.g., Vogel and Machizawa 2004) to identify ERP correlates of the selection and maintenance of task-relevant tactile stimuli. When participants memorized the spatial locations of one or two tactile sample pulses on the left or the right hand, an enhanced negativity with a centroparietal focus emerged contralateral to the hand where the memorized tactile sample was delivered. This tCDA component was sensitive to tactile working memory load, as it was larger on trials where participants had to remember two tactile stimulus locations than when only a single tactile location had to be memorized.
The load-dependent increase of tCDA amplitudes was more pronounced for participants with higher tactile working memory capacity than for individuals whose capacity (measured by Cowan’s K) was closer to 1, mirroring previous findings for the visual CDA component (Vogel and Machizawa 2004). Furthermore, the tCDA component was reliably larger on trials with fast vocal responses in the high-load condition, which were also more accurate than slow responses. This demonstrates that the tCDA component is linked to behavioral performance on individual trials. These observations strongly suggest that the tCDA is an electrophysiological correlate of the maintenance of somatosensory information in tactile working memory.

Analogous to the visual CDA, which has a modality-specific topography over posterior visual cortex (McCollough, Machizawa, Vogel 2007), the tactile CDA component emerged at contralateral central electrodes. The scalp topography of the tCDA in a CSD-transformed map (Figure 4) also suggests neural generators that are located within the somatosensory system. We conclude that the tCDA component reflects the spatially selective activation of modality-specific brain regions contralateral to the task-relevant hand during the retention of tactile stimuli in working memory. These results provide new support for the sensory recruitment model, which assumes that brain regions involved in the perceptual processing of sensory stimuli are also active during the maintenance of these stimuli in working memory. It should be noted that topographical distributions of CSD-transformed scalp maps only allow relatively coarse approximations of the neural origins of components such as the tCDA, and that the exact anatomical basis of this component needs to be determined in future work.

Previous research has used transcranial magnetic stimulation (TMS; Harris et al. 2002) and EEG source reconstruction techniques in studies with human
participants (Spitzer and Blankenburg 2011), as well as single-cell recordings in
monkeys (Romo and Salinas 2003) to show that the activity of neurons in primary
(SI) and secondary (SII) somatosensory cortex is modulated in tactile working
memory tasks. For example, a suppression of alpha activity indicative of attentional
processing was found over contralateral SI during the retention period of a
vibrotactile frequency discrimination task (Spitzer and Blankenburg 2011).
Asymmetric alpha band oscillations have also been suggested as the physiological
basis of the visual CDA component (van Dijk et al. 2010). Indirect evidence for a
recruitment of somatosensory brain areas comes from a tactile EEG study that used
task-irrelevant probe stimuli presented during the retention period to examine how
working memory influences somatosensory encoding (Katus, Andersen, Müller
2012). The retention of locations in working memory was mirrored by spatially
selective modulation of early ERP components to tactile probe stimuli with putative
origins in somatosensory areas such as SII (Frot and Mauguière 1999). These lines
of evidence point towards close links between the maintenance of tactile information
in working memory and the spatially specific activation of early somatosensory areas.
The critical new finding of the present study is the discovery of the tCDA component
that reflects the maintenance of tactile information in a sustained and load-dependent
manner. Because the tCDA is computed by comparing ERPs at electrodes
contralateral and ipsilateral to the location of memorized tactile events, it only reflects
the difference in the absolute activation of contralateral versus ipsilateral
somatosensory areas, and should therefore not be interpreted as evidence that
tactile working memory storage is exclusively contralateral. In fact, there is
electrophysiological evidence that ipsilateral somatosensory cortex may also be
involved in the maintenance of tactile pattern information (Li Hegner et al. 2007).
The tCDA component was preceded by an earlier contralateral negativity, (N2cc component) which emerged around 180 ms after sample stimulus onset. Similar to the tCDA, the N2cc showed a centroparietal scalp topography (see Figures 2 and 4), and was larger in the high-load as compared to the low-load condition. This new N2cc component is likely to represent the somatosensory equivalent of the well-known visual N2pc component. The N2pc is triggered at contralateral posterior electrodes at a similar post-stimulus latency during the attentional selection of targets among distractors in visual displays (Eimer 1996; Luck and Hillyard 1994), and precedes the CDA in visual working memory studies that employ a similar delayed match-to-sample task as the one used in the present study (e.g., Anderson, Vogel, Awh 2011; McCollough, Machizawa, Vogel 2007). The load-dependent increase of the tactile N2cc component observed in the present study mirrors previous findings for the visual N2pc, which increases in size with the number of attended objects in visual displays (e.g., Drew and Vogel 2008; Mazza and Caramazza 2011).

The absence of N2cc components in previous ERP studies of tactile spatial attention is due to the fact that instead of employing bilateral stimuli, tactile events were delivered to a single location on the left or right hand. In these studies, modality-specific components of the somatosensory event-related potential, such as the P100 or N140, were found to be larger for tactile stimuli at currently attended as compared to unattended positions (e.g., Forster and Eimer 2005), demonstrating that spatial attention enhances the sensory processing of tactile events. Analogous to the visual N2pc, which is elicited when target and distractor objects appear in both visual hemifields, measurement of the N2cc component requires that relevant and irrelevant tactile events are presented simultaneously to both hands, or to other homologous locations on the left and right side of the body. Note that the modality-specific somatosensory N2cc component found here is distinct from another ERP component
with the same label that has been observed in stimulus-response compatibility experiments, and is linked to visuospatially guided response selection (Praamstra and Oostenveld 2003). The question whether the effects of memory load on the N2cc and tCDA components reflect load-sensitive modulations of two distinct processing stages (i.e., the attentional selection and the subsequent storage of task-relevant tactile information in working memory), or of a single memory maintenance stage that temporally overlaps with the N2cc component needs to be investigated in future studies where the demands on attentional target selection and working memory load are independently manipulated. In addition to the N2cc and tCDA components, a sustained bilateral CNV component that was observed in the interval between sample and test stimuli was also modulated by memory load. This modulation may primarily reflect differences in the preparation for the match/mismatch decision in response to the test stimulus, which is more demanding in the high-load condition. However, the presence of load effects at ipsilateral electrodes could in principle as well reflect contributions of ipsilateral somatosensory cortex to working memory maintenance (Li Hegner et al. 2007; see also van Ede, Lange, Maris 2013).

When considered together with the results of previous ERP investigations of visual working memory (Anderson, Vogel, Awh 2011; McCollough, Machizawa, Vogel 2007; Vogel, McCollough, Machizawa 2005; Vogel and Machizawa 2004), the current findings reveal striking similarities between the mechanisms that underlie the spatial selection and selective maintenance of sensory stimuli in vision and touch. During both visual and tactile working memory tasks, two contralateral ERP components are elicited successively, with a highly similar time course in both modalities. N2pc and N2cc components that emerge around 180 ms after sample display onset reflect spatial selection during encoding of task-relevant visual or tactile information. The subsequent CDA and tCDA components are linked to the sustained maintenance of
stored information during the retention period. The fact that the load-sensitive tCDA component observed in this study showed a topography over lateral central somatosensory areas (see Figure 4), while the visual CDA component is elicited over lateral posterior visual cortex (McCollough, Machizawa, Vogel 2007) strongly suggests that the maintenance of visual or tactile information in working memory involves the activation of distinct modality-specific regions, in line with the sensory recruitment model of working memory (D’Esposito 2007; Pasternak and Greenlee 2005; Postle 2006; Sreenivasan, Curtis, D’Esposito 2014). In both vision and touch, neural networks that mediate the perceptual processing of sensory signals contribute to the storage and maintenance of information in working memory.

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References


**Figure Legends**

**Figure 1.** Illustration of the experimental setup. Participants memorized a tactile sample set at one task-relevant hand to compare it with a test set on the same hand after a 2 second retention period. Memory load was varied between trials (low load: one pulse, high load: two pulses per hand). The relevant hand (left, right) was varied between blocks. The example shown here illustrates a high-load trial where the locations of tactile sample and test stimuli (symbolized by white dots) are identical at the left hand (match), but not at the right hand (mismatch).

**Figure 2.** Grand mean ERPs elicited in the 2000 ms interval following sample stimulus onset in the low-load and high-load conditions. ERPs were averaged across
lateral central electrode clusters contralateral (blue lines) and ipsilateral (red lines) to
the hand where the memory task was performed. Difference maps show the
topographical distribution of lateralized effects in the N2cc (bottom) and tCDA (top)
time windows. These maps represent the amplitude difference of contralateral minus
ipsilateral recordings, collapsed across blocks where the memory task was
performed with the left or right hand. Enhanced contralateral negativities are shown
in blue. The two bottom panels show difference waveforms for the low-load and high-
load condition, obtained by subtracting electrodes ipsilateral to the task-relevant hand
from contralateral electrodes, and HEOG difference waveforms, calculated by
subtracting HEOG electrodes ipsilateral to the task-relevant hand from contralateral
electrodes after artifact rejection. In these HEOG difference waves, any eye
movements towards the task-relevant hand would be reflected by negative
(downward) HEOG deflections.

Figure 3. Correlation of individual participant’s tactile working memory capacity $K$ (x-
axis) and the increase of tCDA amplitudes in the high-load relative to the low-load
condition measured for each participant (y-axis). $K$ was calculated on the basis of
individual performance in the high-load condition.

Figure 4. Grand mean current source density (CSD) topographical maps, showing
the scalp distribution of lateralized effects in the N2cc and tCDA time windows. These
maps represent the amplitude difference of contralateral minus ipsilateral recordings,
collapsed across blocks where the memory task was performed with the left or right
hand, and averaged across the low- and high-load conditions. Six electrodes at
lateral central scalp regions (black dots) were averaged for each recording cluster
(contra- and ipsilateral to the task-relevant hand). The presence of lateralized effects was also tested for different sets of electrodes over anterior (white triangles) and posterior (white crosses) scalp areas. Reliable lateralized effects were present only for the central electrode cluster.
FIGURE 1
FIGURE 2
Correlation

$r = -0.640$, $p = 0.008$

Figure 3

Memory capacity ($K$)

cDCA change from low to high load

$K = 1$

$K = 2$
FIGURE 4

N2cc
(180-260 ms)

- µV/m²
- +0.05
- 0
- -0.15

△ Anterior
p > 0.05

● Central
p < 0.001

× Posterior
p > 0.3

tCDA
(300-2000 ms)

△ Anterior
p > 0.9

● Central
p = 0.001

× Posterior
p > 0.2