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#### Monitoring multiple deadlines relies on spatial processing in posterior parietal cortex

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#### Abstract

Proactively coordinating one's actions is an important aspect of multitasking performance due to overlapping task sequences. In this study, we used functional magnetic resonance imaging (fMRI) to investigate neural mechanisms underlying monitoring of multiple overlapping task sequences. We tested the hypothesis that temporal control demands in multiple-task monitoring are offloaded onto spatial processes by representing patterns of temporal deadlines in spatial terms. Results showed that increased demands on time monitoring (i.e., responding to concurrent deadlines of 1–4 component tasks) increasingly activated regions in the left inferior parietal lobe (IPL) and the precuneus. Moreover, independent measures of spatial abilities correlated with multiple-task performance beyond the contribution of working memory. Together, these findings suggest that monitoring and coordination processing. We suggest that the precuneus is involved in tracking of multiple task timelines, whereas the IPL constructs spatial representations of the temporal relations of these overlapping timelines. These findings are consistent with the spatial offloading

hypothesis and add new insights into the neurocognitive mechanisms underlying the coordination of multiple tasks.

to Review Only

#### Introduction

Modern life imposes increasing demands on our ability to keep track of multiple overlapping tasks that include various timelines and deadlines. For example, we may pay attention to TV programs or cell phone notifications, while taking care of the household or work. During multitasking, several different tasks, or instances of the same task, are combined and executed in concert with overlapping timelines (Burgess, Veitch, de Lacy Costello, & Shallice, 2000; Oswald, Hambrick, & Jones, 2007; Strobach, Wendt, & Janczyk, 2018). Previous behavioural (Hambrick, Oswald, Darowski, Rench, & Brou, 2010; Konig, Buhner, & Murling, 2005; Logie, Cocchini, Delia Sala, & Baddeley, 2004; Meyer & Kieras, 1997) and neuroimaging (Collette et al., 2005; Deprez et al., 2013; Hsu, Zanto, Anguera, Lin, & Gazzaley, 2015; Kübler & Schubert, 2017; Medeiros-Ward, Watson, & Strayer, 2015; Nijboer, Borst, van Rijn, & Taatgen, 2014; Stelzel, Kraft, Brandt, & Schubert, 2008; Strobach, Antonenko, Schindler, Flöel, & Schubert, 2016; Szameitat, Lepsien, Cramon, Sterr, & Schubert, 2006; Tschernegg et al., 2017; Verghese, Garner, Mattingley, & Dux, 2016) studies indicate that executive functions play a critical role in multitasking. However, most studies typically concern isolated, experimentally tractable, aspects of dual- and multitasking, using structured or bottom-up controlled paradigms (see also Logie, Trawley, & Law, 2011).

In contrast, everyday multitasking usually involves planned (top-down controlled) task management (e.g., preparing breakfast or carrying out multiple errands, Craik and Bialystok 2006; Logie et al. 2011), with high demands on time keeping (i.e. keeping track of overlapping task timelines and deadlines). Executive control demands are typically high in proactive tasks, but these accentuated control requirements can be alleviated by relying on prior knowledge and experience (Loukopoulos, Dismukes, & Barshi, 2009; Wickens, 2008), as well as cognitive offloading

strategies (Risko & Gilbert, 2016). Specifically, studies in different domains of cognitive sciences have shown that humans can change the task requirements (and affordances) by relying both on internal and external representations of the task (Wilson & Golonka, 2013; Zhang & Norman, 1994; for an overview, see also Risko & Gilbert, 2016).

Reflecting this *offloading* perspective, we have previously proposed a mechanism for alleviating control demands in multitasking performance. In detail, when task-specific knowledge is not available or insufficient, temporal control demands during multitasking are offloaded onto spatial abilities by representing patterns of temporal deadlines in spatial terms (Mäntylä, 2013; Todorov, Del Missier, Konke, & Mäntylä, 2015). This idea resonates with similar views of "timeto-space-mappings" in other domains of cognitive sciences which suggest that temporal and spatial processing are closely related (e.g., Bonato, Zorzi, & Umiltà, 2012; Dehaene & Brannon, 2011; Gijssels, Bottini, Rueschemeyer, & Casasanto, 2013; Núñez & Cooperrider, 2013). This spatial offloading hypothesis is supported by several studies showing that spatial ability (i.e. mental rotation) and executive functioning (i.e. working memory updating) are independent predictors of multitasking performance, and that spatial ability predicts multitasking over and beyond executive functioning (Logie et al., 2011; Mäntylä, 2013; Mäntylä, Coni, Kubik, Todorov, & Del Missier, 2017; Morgan et al., 2013; Todorov, Del Missier, & Mäntylä, 2014; Todorov, Kubik, Carelli, Del Missier, & Mäntylä, 2018). Moreover, it was shown that multitasking, compared to dual-tasking, involves an incremental contribution of spatial ability (Kubik, Zimmermann, Del Missier, Frick, & Mäntylä, 2018, 2019), and that concurrent spatial load selectively impairs multitasking performance (Mäntylä et al., 2017; Todorov et al., 2018).

The aim of this study was to test a central assumption of the spatial offloading hypothesis, namely that spatial processes are involved when keeping track of multiple overlapping task

timelines. Accordingly, we expected neural areas involved in spatial processing, especially superior and inferior parietal cortex (Cohen et al., 1996; Halari et al., 2006; Jordan, Heinze, Lutz, Kanowski, & Jäncke, 2001; Kesner & Long, 1998; Richter, Ugurbil, Georgopoulos, & Kim, 1997; Sack, 2009; Zacks, 2008), and the hippocampus (Eichenbaum, 2017) to be incrementally engaged when the demands on time monitoring are increased using parametric analyses. Additionally, a model-based approach was used to investigate whether the spatial offloading hypothesis is specific to multitasking situations (see Mäntylä, 2013), or generally valid for time monitoring tasks in a wider range of situations (see also Kubik et al., 2018, 2019). Specifically, we tested whether recruitment of spatial processing areas increases gradually with time monitoring demands (approximated by a linear increase model, reflecting general processes), or whether spatial processing areas are selectively recruited when the demands on time monitoring reach a threshold, for example only during multitasking situations (i.e. following a step-like pattern).

#### Methods

#### **Participants**

Twenty-four healthy, right-handed participants between 18–35 years (*M* [*SD*]; 27.9 [4.1]; 12 female) took part in the study. Participants were recruited from a large-scale behavioral study, in which they completed a series of cognitive tasks, including tests of mental rotation, working memory, and different versions of time-based monitoring tasks (see also Kubik et al., 2018, 2019). The behavioral study was completed approximately six months before the fMRI study. All participants were screened for claustrophobia, neurological, and psychiatric medications, MRI contraindication, and all had normal or corrected to normal vision using scanner compatible glasses or contact lenses. One participant was excluded from the analysis due to excessive head

movements (>3 mm) during all scanning sessions. Participants received financial compensation of 600 SEK for participation. The study was approved by the Regional Ethical Review Board in Stockholm and written consent was obtained from all participants before the study commenced.

#### Tasks

#### Countdown task

Monitoring of multiple timelines was tested with an adapted version of the *counter-task* paradigm (Mäntylä, 2013; Todorov et al., 2014). In the original version, participants monitor a number of counters (clocks) and respond to predefined targets (e.g. Mäntylä, 2013). For the adapted version used here, forward-running counters of numbers were replaced with backward-running letter-loops (letter countdowns) with fixed targets (Figure 1). Therefore, in the current task, inherent spatial processing was reduced by removing the need to update and maintain (numerical) target values. and thus reducing possible spatial associations of the stimuli (further spatial associations due to stimulus-response mappings are controlled for using a control task; see below). This adapted task was recently shown to be highly correlated to and reveal similar associations with the cognitive reference measures (e.g. executive functioning, spatial ability) as the original counter task (Kubik et al., 2018, 2019). Time monitoring demands were manipulated by varying the number of letter countdowns that had to be monitored and responded to. Specifically, the task included conditions with one to four letter countdowns at a time. Letter countdowns consisted of loops of letter series that followed the alphabet backwards (e.g. H-G-F-E-...), until A, at which point (after a 2-second pause) the loop started again (here referred to as a single countdown series). The letter A was defined as a target letter for each countdown series. Participants had to respond to the occurrence

of the target letter as accurately as possible by pressing a *target button*. Multiple countdown series formed a trial, during which a number of countdowns, varying from one to four (referred to as conditions), had to be monitored and responded to continuously for 60 s. Importantly, letter countdowns were not continuously visible; instead, participants needed to uncover them by pressing one of four *monitor buttons*. This displayed the corresponding countdown for 1 s, while no other countdowns could be displayed. Target-button presses were assigned to the latest monitored countdown. Thus, participants had to make sure that, prior to indicating a target occurrence, the corresponding countdown was displayed last. The target button could, but did not have to, be pressed while the corresponding countdown was displayed. Upon pressing the target button, the monitored countdown changed into hidden state again, allowing for displaying of other countdowns. Monitor buttons and target buttons were assigned to MR compatible button boxes, with four monitor buttons to be pressed with the fingers of the right hand, and a single target button to be pressed with the left index finger. Countdowns (after pressing the color-matched monitor button) were presented centrally on the screen to reduce eye movements and changes in spatial layout.

To prevent cross-task monitoring (i.e., predicting the state of another countdown), and to match the number of targets between trials with varying numbers of countdowns, each letter series progressed at different rates and covered different series lengths (e.g., *F* to *A* at a slow rate, or *L* to *A* at faster rate). Updating rates (i.e. the time between two letter changes in a series of a given countdown) were adjusted for the number of countdowns. For example, for the 4-countdown trials, the updating rates were 816, 1376, 1536, and 1568 ms, respectively. For trials with less than 4 countdowns, updating rates were randomly selected from the 4 possible updating rates, and adjusted to  $\frac{3}{4}$ ,  $\frac{1}{2}$ , or  $\frac{1}{4}$  of the updating rate intervals for 4 countdown trials, for trials with

respectively 3, 2, and 1 countdowns. Correspondingly, to warrant similar number of targets per countdown, countdowns started (and restarted after reaching target state) at varying letters, with series lengths of 25, 16, 11, and 10 letters, for the fastest to slowest updating rates, respectively. Using these parameters, each trial of 60 seconds contained 8–10 evenly distributed targets, independently of the number of countdowns, and independently of the randomly selected countdown settings (for trials with less than 4 countdowns). The assignment between monitor buttons and countdowns (in terms of updating rate/series lengths) was randomized for each trial.

## << Insert Figure 1 about here >>

A control task was introduced to control for spatial mapping of stimulus–response associations, visual stimulation, and motor output. Critically, the control task did not require (or enable) time monitoring as both monitoring and target occurrence was random to the participant. Similar to the experimental task, the control task was presented in trials consisting of 1 to 4 random letter series (here referred to as control countdowns). Control countdowns contained random letters (A–Z) that were presented automatically for 1 second (comparable to actively monitored countdown series during the experimental task). Participants were asked to press the corresponding monitor button upon presentation of a control countdown appearances was based on the active monitoring behavior (experimental task) of a pilot participant. Therefore, the control task provides condition-matched control conditions (i.e. in terms of number of simultaneous countdowns and monitor buttons per condition), visual input, and motor output (both monitor- and target-button presses, reflecting timing and complexity/number of response options).

The experiment consisted of five runs, including three runs of the experimental task, intermitted by two runs of the control task (exp - con - exp- con - exp). Experimental task runs

consisted of 2 blocks, and each block contained 4 trials, one per condition (1–4 countdowns). Control-task runs consisted of 1 block with 4 trials each, one per condition (1–4 control countdowns). Trial order within each block was randomized. Each trial lasted 60 seconds, during which 8–10 targets appeared. A 20-second break was given between two successive trials/blocks during which a cross hair was displayed centrally on the screen. After each run (experimental or control task), the scanner was stopped and a variable break was given to the participant. Participants remained inside the scanner between runs. Participants were informed verbally and visually about the upcoming task (experimental or control), and the experiment continued when participants indicated they were ready to do so.

The main behavioral outcome measure was *monitoring accuracy*, calculated as the ratio of successful attempts to indicate the occurrence of target items, relative to all target occurrences. The temporal discrepancy between target occurrence and response was taken as *temporal (in)accuracy*. This included early responses (i.e. before the target occurred) as well as late responses (response after target occurred). For further analyses, only absolute values (i.e. temporal offset) were used. Overall monitoring accuracy was calculated over all countdowns within a condition. Moreover, monitoring accuracy was calculated at a countdown level, that is – for each trial – the accuracy on every single countdown. These *countdown-specific monitoring accuracy* was sorted by performance into best, 2<sup>nd</sup> best, 3<sup>rd</sup> best, and 4<sup>th</sup> best countdown (as far as applicable for conditions with less than 4 countdowns), such that performance of, for example, the best countdowns could be compared between conditions (i.e. depending on the number of additionally monitored countdowns). Only the first target-button press was recorded for each countdown series (i.e. between two targets), and data from additional target button presses to the same countdown/target were not considered further. To allow for early and late responses, target button

presses that occurred shortly after a target appearance (up to 20% of the target interval), were assigned to the previous target (i.e. late responses), and the remaining target button presses were assigned to the upcoming target (i.e. early responses). Target-button presses that occurred more than 3 seconds from the target were removed from the behavioral data analysis since these responses could not reliably be assigned to a specific target or counter (< 1%). Therefore, only target button presses close to a target occurrence were analyzed. The interquartile range (IQR; quartile 3 – quartile 1) rule of IQR  $\times$  1.5 (Tukey, 1977) was used for detecting the presence of outliers in target button presses. Accordingly, target button presses with discrepancies that were below Q1 – 1.5 IQR or above Q3 + 1.5 IQR (Tukey, 1977) were removed as outliers (5.22%). A secondary behavioral outcome measure was *monitoring behavior*, specifically the distribution of Zie monitoring actions over time.

#### *Spatial ability and working memory*

Spatial processing and working memory skills were tested in all participants within the largescaled behavioral study reported elsewhere (Kubik et al., 2018, 2019). From these data, measures of spatial ability and working memory performance were used in the current study.

Spatial ability was measured with the redrawn version of the mental rotation test (MRT, version A; Peters et al., 1995; Vandenberg & Kuse, 1978). This pen-and-paper task contained 12 items. Each item was made up of a target cube construction and four probes. Two of the probes matched the target cube construction, however, rotated around multiple axes relative to the target cube construction; two other probes did not match the target cube construction. The matching cube constructions had to be identified by the participants. It is assumed that participants mentally rotate the figures and try to match them to each other. Participants had three minutes to solve as many items as possible. For each participant, we counted the number of correctly answered items (with

"correct" referring to having identified both matching cube constructions, and none of the nonmatching probes), within the three minute period.

*Working memory* was measured using verbal and spatial binding tasks (Wilhelm, Hildebrandt, & Oberauer, 2013). The verbal (letter–color) binding task consisted of 15 trials, in which a short list of letter–color pairs were sequentially presented for 1 s each (ISI = 3 s). Similarly, the spatial (letter–location) binding task consisted of 14 trials, during which a list of letters were presented sequentially for 1.5 s (ISI = 0.5 s) at specific locations within a 3 × 3 grid. Load levels ranged from 2 to 6 pairs. After each trial, participants were asked to remember the letter–color and letter–location pairs in a cued recall test. In half of the test trials, participants were probed with the letters and had to select/indicate the corresponding color/location; for the other half, participants were probed with the color/location, and had to select the corresponding letter (for more details see Kubik et al., 2018, 2019). The average performance (proportion correct responses) over both tasks was used as measure for participants' ability of working-memory binding.

#### Behavioral data analysis

Behavioral data obtained during the multitasking (countdown) task was analyzed in terms of performance (monitoring accuracy, countdown-specific monitoring accuracy, temporal accuracy) and monitoring behavior. Performance was measured by overall target detection (monitoring accuracy; in line with previous studies Kubik et al., 2018, 2019; Mäntylä, 2013; Todorov et al., 2014), and temporal accuracy, a more sensitive measure (see "Countdown task"). Monitoring accuracy and temporal accuracy were analyzed using one-way ANOVAs, with condition as a within-subjects factor, followed by pairwise comparisons between conditions where applicable.

Furthermore, countdown-specific monitoring accuracy (see "Countdown task") was compared between the single best countdowns of each condition (as well as additional counters) between conditions. To test and demonstrate possible relations to spatial processing (as in a spatial offloading mechanism), monitoring accuracy was correlated with participants' spatial abilities (mental rotation task), both for overall performance, and for performance on the best counters for each condition. Moreover, following previous studies showing that spatial abilities (MRT) explain multitasking performance (monitoring accuracy) beyond working-memory skills and executive functioning (Kubik et al., 2018, 2019), monitoring accuracy was correlated with mental rotation abilities after correction for working-memory abilities. Therefore, partial correlations were computed between monitoring accuracy, spatial ability, and working memory.

*Monitoring behavior* was analyzed in terms of monitoring actions per condition, overall (i.e. number of monitoring actions per trial), and over time. One purpose of analyzing monitoring behavior was to verify and demonstrate that participants proactively plan their monitoring behavior. Moreover, by comparing monitoring behavior between conditions, we aimed to demonstrate that demands on time monitoring increase with an increasing number of monitored countdown series. For overall monitoring behavior, monitoring actions were counted for each condition, and compared using one-way ANOVAs, followed by posthoc comparisons where applicable. Furthermore, monitoring behavior was analyzed in a time-resolved manner: monitoring actions were time-locked to target onsets on a countdown-series specific basis (e.g. timing of monitoring of the red countdown relative to the target occurrence of the red countdown), and split in bins of 500 ms duration. Only the final 6 seconds (12 bins) before target onset were considered during this analysis, such that all conditions covered the analyzed period (based on target-to-target intervals). Time bins were categorized into 'early' bins (6–3 seconds before target onset) and 'late'

bins (1.5–0 seconds before target onset). Time-resolved monitoring behavior was analyzed in a two-way ANOVA with factors time (early, late) and condition (number of countdowns). A significant Time × Condition interaction was followed up by time-specific one-way ANOVAs (testing for effects of condition within each epoch; followed by pairwise posthoc comparisons between conditions where applicable). Ratios between early and late monitoring behavior were calculated by dividing the (average) number of monitoring actions during early time bins by the (average) number of monitoring the late time bins for each condition. Ratios were compared between conditions using repeated measures ANOVA.

#### MR data acquisition and preprocessing

A 3T MRI scanner (GE Discovery MR 750; General Electric, Boston, MA, USA) with an 8-channel head coil was used for acquiring functional and structural MR images. Anatomical images were acquired using a T1-weighted sequence (TR/TE = 6400/2800 ms, voxel size:  $1 \times 1 \times 1$  mm). Blood-oxygen-level-dependent (BOLD) signal was recorded with a whole-brain T2\*-weighted echo-planar imaging sequence (TR/TR = 2000/30 ms; voxel size:  $3 \times 3 \times 3$  mm, gap size: 0.5 mm). The task was divided into 3 experimental runs of 11.2 minutes each and 2 control runs of 5.7 minutes each during which 336 and 172 volumes were recorded, respectively. Additional diffusion-tensor imaging and resting-state fMRI scans were acquired after the task fMRI scans. This data is not part of the current report.

Images were preprocessed using SPM12 (Wellcome Centre for Human Neuroimaging, University College London, UK), and ICA-AROMA (Pruim et al., 2015) as part of FSL v5.0 (FMRIB's Software Library). SPM-based preprocessing included spatial realignment, slice time

correction and co-registration of the functional images to the structural image of each participant. All brains were normalized to the MNI template using the unified segmentation procedure, and resampled to  $2 \times 2 \times 2$  mm voxel size. Functional images were smoothed using a 6 mm full width at half maximum (FWHM) Gaussian kernel. Next, AROMA was used to remove movement related artifacts. BOLD signals from white matter, cortico-spinal fluid and out of brain areas, obtained from the segmentation in SPM, were extracted, and removed from the fMRI time using linear regression, and the data was filtered using a 128 s high pass filter.

#### fMRI data analysis

Sur. For single subject analyses, separate general linear models were constructed for each run, modelling trials using boxcar functions, with separate regressors for each condition (i.e. number of countdowns), with a length corresponding to the block length. The purpose of these regressors was to capture underlying state activation during monitoring of one, two, three or four overlapping task timelines. Additional regressors were constructed to model button presses and visual stimulation. Using FEAT (part of FSL), we first estimated statistical parametric maps corresponding to each condition within each run. Next, in a second level fixed-effect analysis, we calculated the contrast of BOLD signal during the experimental task and the control task, separately for each condition/number of countdowns. This resulted in four whole-brain maps per participants, corresponding to the BOLD signal difference between experimental and control task for 1, 2, 3 and 4 letter series (i.e., exp1 > con1, exp2 > con2, exp3 > con3, exp4 > con4; in the remainder referred to as *conditions*), which were used for group-level random-effect analyses.

We used complementary approaches for *group-level random effect analyses* of the functional imaging data. In a *contrast-based approach*, we analyzed the activation corresponding to each of the experimental conditions (i.e. monitoring 1-4 deadlines), compared to the corresponding control conditions, and then directly compared activation maps corresponding to different conditions/numbers of parallel countdowns. In a complementary *model-based approach*, we identified brain regions that followed linear and step-function increases with the number of parallel countdown series, and compared the model fits. These analyses are described in detail below.

# Part I: Contrast-based analyses

We obtained neural activation maps for each experimental condition of the multitask paradigm, corrected for activation during the corresponding control-task condition. Specifically, for each condition (i.e. number of parallel countdowns, 1–4), we compared the whole-brain activation maps between the experimental and the control task in a group analysis, treating subjects as random effects using nonparametric paired *t*-tests in PALM (Permutation Analysis of Linear Models; Winkler, Ridgway, Webster, Smith, & Nichols, 2014). The resulting condition-specific activation maps were compared between the different conditions (i.e. number of component tasks, corrected for condition-specific control-task activation) using nonparametric *F*- and *t*-tests. First, we performed a one-way within-subject (repeated measures) ANOVA (*F*-test) with 4 levels (reflecting the 4 conditions). The *F*-test was followed up with condition-specific comparisons (nonparametric paired *t*-tests), using the significant clusters obtained from the *F*-test as mask. Second, we performed whole-brain, condition-specific comparisons. For this comparison, 1-tailed one-sample *t*-tests of the difference maps (i.e. 2y1, 3y1, 4y1, 3y2, 4y2, 4y3) were used to identify increased

brain activation as a function of increased numbers of parallel monitored countdowns. All inferential statistics were performed using permutation testing (sign flipping) with 1000 random permutations, combined with gamma acceleration to improve *p*-value estimates. Significant clusters were identified using a cluster-defining threshold of Z > 3.1 (corresponding to  $p_{(unc)} < .001$ ), and cluster-corrected threshold of  $p_{(FWE)} < .05$  (Worsley, 2001).

#### Part II: Model-based analyses

For the model-based approach, we identified brain regions that showed activation patterns that corresponded to four proposed theoretical models (Figure 2). Models followed a priori predictions how activation depends on the number of counters in different parts of the brain, following different versions of the spatial offloading hypothesis. Specifically, a linear increase model (Figure 2A), where activity increases linearly with the number of monitored countdowns, a *multi-task (3+) step model* (Figure 2C), with high activation levels for 3 and more tasks, relative to 1 and 2 tasks, and finally, a *dual-task step model* (Figure 2B), with a high activation level for 2 and more monitored series, relative to 1 series. Multi- (or dual-) task specific spatiotemporal processes should follow a multi- (or dual-) task step model activation pattern, whereas general spatiotemporal processes should follow the linear increase model. Importantly, given that these models are implicitly tested against a null model (i.e. no systematic activation differences depending on the number of monitored countdowns), for spatiotemporal offloading mechanisms to be supported, a model needs to explain activation differences in cortical regions related to spatial processing. The pattern of this activation (e.g. linear vs multi-task step increase) will provide information about the specificity (with respect to dual- or multi-task) of such a spatiotemporal offloading process.

<< Insert Figure 2 about here >>

For each participant, we used FEAT to obtain model-specific activation maps from the [experimental – control task] contrast images for each number of component tasks (see *Part I: Contrast-based analyses*). Group average model activation maps were obtained using PALM, using 1000 permutations, gamma acceleration (Winkler, Ridgway, Douaud, Nichols, & Smith, 2016), and whole-brain masking. Moreover, we obtained differences between model activation maps by pairwise comparison of model-specific activation maps, using PALM (1000 permutation, gamma acceleration, whole-brain masking). We used a cluster-defining threshold of Z > 3.1 (corresponding to  $p_{(unc)} < .001$ ), and cluster corrected threshold of  $p_{(FWE)} < .05$  (Worsley, 2001) to identify significant clusters. Only positive relationships (i.e. activation increases following the models) were considered. For visualization purposes, we extracted parameter estimates from the identified affected regions (i.e. significant clusters in any of the model-based analyses) for each condition-specific neural activation map (i.e. number of countdowns). Mean parameter estimates were extracted from regions of interest centered on the peak coordinates of affected regions (sphere-shaped, 5 mm radius).

#### Region of interest analysis: hippocampus

In addition to whole-brain analyses, we analyzed activation in two regions of interest (ROI), the left and right hippocampus. ROIs were defined based on a probabilistic atlas (Harvard-Oxford subcortical structural atlas, part of FSL). ROIs for left and right hippocampus were based on maximum probability for left and right hippocampus, with a lower limit requirement of 50% probability. Parameter estimates of the experimental – control task contrasts were extracted for both ROIs. Parameter estimates were compared between conditions using 1-way ANOVAs, and fitted to a set of pre-defined models (see Figure 2). Model-fitting was performed using the FITLM function in MatLab to obtain the Akaike information criterion (AIC; Akaike, 1974) as well as R-squared values for each model/ROI.

# Brain-behavior correlations

To investigate whether inter-individual differences in behavioral performance during multitasking can be directly related to inter-individual differences in brain function, we correlated measures of behavioral performance with measures of brain activation in the set of affected regions. Specifically, we created regions of interest for each affected region (based on any of the model-based analyses, see "Model-based analyses"), using sphere-shaped regions of interest (5 mm radius) centered on the peak coordinates of the affected regions. Parameter estimates of the 4-countdown condition (i.e. multitask-specific activation), as well as the difference between 4- and 2-countdown conditions (i.e. activation during multitasking corrected for dual-tasking), were correlated with corresponding measures of behavioral performance (i.e. monitoring accuracy during multitasking divided by monitoring accuracy during dual-tasking condition, respectively) and measures of spatial ability (MRT scores), using Spearman's rank correlations.

#### Results

#### Behavioral results

Monitoring accuracy (proportion correctly identified target letters) was not significantly affected by the number of monitored countdown series (F(3,69) = 1.17, p = .329,  $\eta^2 = .003$ ). Specifically, participants correctly identified a high number of countdown targets independent of the number of monitored countdown series (1 countdown:  $0.83 \pm 0.21$  (mean  $\pm$  SD); 2 countdowns:  $0.83 \pm 0.19$ ; 3 countdowns:  $0.81 \pm 0.19$ ; 4 countdowns:  $0.81 \pm 0.18$ ; Figure 3A). However, an analysis of *countdown-specific monitoring accuracy* (i.e. monitoring accuracy for individual countdown series within each trial) revealed that performance of the best countdown series increased with the number of monitored countdown series (Figure 3B). The same pattern was observed for the 2<sup>nd</sup> and 3<sup>rd</sup> best countdown series of the dual- and multi-tasking conditions. Whereas overall monitoring accuracy did not show significant correlations with mental rotation scores (all p > .10), accuracy for the best countdown in each trial did correlate significantly with MRT scores for the multitasking conditions (3 countdowns: r = .459, p = .032; 4 countdowns: r =.511, p = .015), however, not significantly with the single-task condition (r = .373, p = .087) and dual-task condition (r = .420, p = .052). Partial correlations between MRT and monitoring accuracy, corrected for individual working memory skills, were significant for the best countdown in the 4-countdown condition (r = .467, p = .033). Partial correlations for best countdowns of other conditions were not significant (3 countdowns: r = .429, p = .053; 2 countdowns: r = .405, p =.069; 1 countdown: r = .366, p = .103). Performance for added countdowns (i.e. 2<sup>nd</sup> best, 3<sup>rd</sup> best,  $4^{\text{th}}$  best countdown series) did not correlate with MRT scores (all p > .10) and none of the partial correlations for  $2^{nd}$ ,  $3^{rd}$  and  $4^{th}$  best countdowns were significant (all p > .10).

*Temporal accuracy* was significantly affected by the number of monitored countdown series (F(3.69) = 26.08, p < .001,  $\eta^2 = .056$ ). Specifically, the temporal accuracy at which targets

were marked decreased with a higher number of monitored countdowns (Figure 3C). In detail, temporal accuracy was higher for 1 countdown ( $390 \pm 207 \text{ ms}$ ) compared to 2 countdowns ( $518 \pm 256 \text{ ms}$ ; t(23) = 6.00, p < .001), 3 countdowns ( $536 \pm 288 \text{ ms}$ ; t(23) = 5.48, p < .001) and 4 countdowns ( $534 \pm 256 \text{ ms}$ ; t(23) = 7.81, p < .001). There were no significant differences between temporal accuracy for 2, 3 and 4 countdowns (all p > .10; all Bonferroni corrected).

#### < Insert Figure 3 about here >

Monitoring behavior was significantly affected by the number of monitored countdown series. Generally, participants performed more monitoring actions with increasing number of monitored countdown series (F(3,69) = 116.19, p < .001,  $\eta^2 = .465$ ). Specifically, monitoring actions increased with each condition (i.e. from 2 to 1 countdown: t(23) = 13.63, p < .001; 3 to 2 countdowns: t(23) = 4.49, p = .001; 4 to 3 countdowns: t(23) = 5.16, p < .001). Average number of monitoring actions per trial was  $31.60 \pm 7.6$  during the 1-countdown condition,  $40.76 \pm 5.79$ during the 2-countdown condition,  $43.43 \pm 4.54$  during the 3-countdown condition, and  $45.04 \pm$ 4.24 during the 4-countdown condition (Figure 4A). A temporal analysis of monitoring behavior (Figure 4B) suggested that monitoring of a specific countdown series (relative to its target occurrences) was similar with regards to the temporal proximity to the target, but differed in the time before target occurrence. This pattern was confirmed by a significant interaction of monitoring actions between time (early vs late within a countdown series) and condition (number of monitored countdown series) (F(3, 69) = 44.64, p < .001). The interaction was driven by a significant difference between conditions during the early epoch (time bins 1-6; F(3, 69) = 48.32, p < .001), and the absence of a significant difference during the late epoch (time bins 10-12; F(3, 1)) (69) = 0.39, p = .761). During the early epoch, pairwise comparisons of monitoring behavior indicated significant decreases in monitoring activity with each added countdown series (2 vs 1

countdown: t(23) = 4.36, p = .001; 3 vs 2 countdowns: t(23) = 7.22, p < .001; 4 vs 3 countdowns: t(23) = 3.89, p = .002; all Bonferroni corrected).

#### <Insert Figure 4 about here>

For each participant and condition, we calculated the ratio between early and late monitoring actions to measure the impact of increased demands on time monitoring on monitoring behavior. The ratio of early monitoring actions relative to late monitoring actions decreased with the number of monitored counters (Figure 4C), meaning that participants perform more monitoring actions when they monitored fewer countdowns, relative to monitoring actions that occurred in close temporal proximity to the target. The decrease can be described by linear regression ( $\beta = -$ .758, F(1,23) = 127.817, p < .001,  $\eta^2 = .847$ ; quadratic: F(1,23) = 9.342, p = .006,  $\eta^2 = .289$ ). 

#### Brain imaging results

#### Part I: Condition-specific brain activation

First, to identify the neural networks involved in monitoring of task time- and deadlines during single-, dual-, and multitasking, we obtained neural activation maps for each condition of the multitasking paradigm, corrected for activation during the corresponding control-task condition (see "Methods – Part I"). For the single-countdown condition, the experimental task, relative to the control task, activated middle cingulate cortex and the supplementary motor area (SMA), bilateral middle frontal gyrus, bilateral superior temporal gyrus and pole, the right inferior parietal cortex, left superior frontal gyrus, and parts of the cerebellum (Figure 5A). With two monitored countdowns, the experimental task, relative to the control task, activated extensively the visual cortex bilaterally, as well as left and right postcentral gyri, middle and superior frontal

gyri, SMA and fusiform gyri, as well as left superior temporal gyrus, and the left parahippocampal gyrus (Figure 5B). Similar results were obtained for the condition with 3 countdowns, activating extensively the visual cortex, middle and inferior frontal gyri, SMA and midcingulate area, preand postcentral cortex bilaterally, as well as left superior and inferior temporal gyri (Figure 5C). The four-countdown condition activated extensively the visual cortex, fusiform gyrus, SMA, middle temporal gyri and poles bilaterally, and middle and frontal gyri (Figure 5D). None of the reverse contrasts ([control – experimental] for each of the conditions) resulted in significant clusters.

<< Insert Figure 5 about here >>

2.04

#### Condition-specific comparisons

Next, activation was compared between conditions. We observed a significant cluster in the precuneus in a one-way ANOVA including all four conditions testing for overall differences between the four conditions (MNI: [2 -54 34]; F = 543.24,  $p_{(FWE)} = .04$ ). Follow-up comparisons using the significant cluster as mask revealed no step-wise differences (i.e. 2 > 1, 3 > 2, 4 > 3; all p > .05; however, all p < .10; all FWE-corrected). However, there were significant differences in comparisons between 3- and 1-countdown conditions (p = .003), 4- and 1-countdown conditions (p < .001), and between 4- and 2-countdown conditions (p = .003). Similarly, whole-brain paired *t*-tests revealed no significant clusters for step-wise comparisons (i.e. 2 > 1, 3 > 2, 4 > 3). However, a significant cluster in the precuneus was observed when comparing multitask conditions with 3 and 4 parallel countdown series with the single countdown condition (Table 1). When comparing

the 4-countdown multitask condition with the single countdown condition, additional clusters were observed in bilateral inferior parietal cortex/intraparietal sulcus, bilateral middle temporal gyrus, medial frontal cortex, bilateral occipital cortex, posterior cingulate, and the right parahippocampal gyrus (Table 1). In the whole-brain comparisons, no differences were observed between the multitask conditions (3 and 4 countdowns) and the two-countdown condition.

<< Insert Table 1 about here >>

## Part II: Model specific activation maps

In a complementary approach, we identified brain regions in accordance to model-based activation patterns. These results are presented in Figure 6 and Table 2. The *linear increase model* (Figure 2A) was associated with activation in the left IPL (extending into the angular gyrus; p = .013), and non-significant clusters in the bilateral precuneus (p = .056), and the right middle temporal gyrus (p = .063). Similarly, the *dual-task step model* (2+; Figure 2B) predicted significant clusters of increasing activation in the left IPL (p = .039), and non-significant clusters in the left and right precuneus (p = .060) and the left inferior medial frontal lobe (p = .082). The *multi-task* (3+; Figure 2C) *step model* predicted activation in the precuneus (p = .023) and left IPL (p = .045), and non-significant clusters in the right middle temporal gyrus (p = .057). Generally, there was extensive overlap between the activation maps of all models, given the similarity of the models.

<< Insert Figure 6 about here >>

<< Insert Table 2 about here >>

Regions of interest (ROIs) were generated based on the significant clusters obtained from the model-specific activation maps (Table 2) for illustration purposes (Figure 7) and for calculation of model fits. These ROIs represented the left inferior parietal cortex (center MNI [-34 -68 42]; significant cluster in the linear increase model, dual-task step model, and multi-task (3+) step model), and the precuneus (MNI [0 -54 36]; significant cluster in multi-task (3+) step model). Visual inspection of extracted parameter estimates generally showed increasing activation with increased number of monitored countdowns in each ROI. Model fit for the inferior parietal ROI was highest for the linear increase model (AIC = 11.26,  $R^2 = .95$ ), followed by the dual-task step model (AIC = 17.86,  $R^2 = .72$ ) and the multi-task step model (AIC = 19.33,  $R^2 = .59$ ). For the precuneus ROI, model fit was highest for the linear increase model (AIC = 6.37,  $R^2 = .99$ ), followed by the multi-task step model (AIC = 19.45,  $R^2 = .73$ ) and dual-task step model (AIC = 20.07,  $R^2 = .68$ ). No correlations between behavioral measures (multi-task performance, mental rotation ability) and brain activation within these ROIs were detected (all p > .10).

<< Insert Figure 7 about here >>

Model comparisons

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Whereas the cluster in the precuneus was significant only for the multi-task (3+) step model (Table 2), no significant differences between model-activation maps of the *multitask-step model*, the *dualtask-step model*, and the *linear-increase model* were observed.

#### Region of interest: hippocampus

In addition to the whole brain, model- and contrast-based analyses, we analyzed conditiondependent differences in the hippocampal gyri, based on its role on spatial processing (Eichenbaum, 2017). No significant differences between conditions were observed for left or right hippocampus (1-way ANOVA; all p > .10; Figure 8). However, for both left and right hippocampus, model fit was best for the dual-task (2+) step models (left: AIC = 4.91,  $R^2 = .77$ ; right: AIC = 4.72;  $R^2 = .80$ ), followed by the linear increase models (left: AIC = 5.57,  $R^2 = .73$ ; right: AIC = 8.49,  $R^2 = .48$ ), and the multi-task step models (left: AIC = 9.43,  $R^2 = .30$ ; right: AIC= 10.64,  $R^2 = .11$ ). No correlations with behavioral performance were observed (all p > .10).

<< Insert Figure 8 about here >>

#### Discussion

The main goal of this study was to identify the neural correlates underlying multiple-task performance. Specifically, we tested a central assumption of the spatial offloading hypothesis, namely whether keeping track of multiple task deadlines relies on spatial processes.

Overall monitoring accuracy was constantly high across the different numbers of countdowns, but detailed analyses suggested that participants adapt their behavior in response to

changing task demands. Specifically, when the number of countdowns in the task increased. performance for the best countdown in each task condition increased. Performance on additional countdowns (2<sup>nd</sup>, 3<sup>rd</sup>, 4<sup>th</sup> best) was reduced, resulting in similar overall performance across task conditions (Figure 3B). This pattern may be the result of strategic choices of the participants, in which they increased effort with more countdowns, while at the same time focusing on a subset of component tasks. Such behavioral facilitation from increasing task demands were recently demonstrated within the context of working memory and executive control (Samrani, Marklund, Engström, Broman, & Persson, 2018). Moreover, and partly in line with previous studies (Kubik et al., 2018, 2019; Mäntylä, 2013; Todorov et al., 2014), performance on this best countdown was predicted by participants' spatial (mental rotation) abilities, even after correction for working memory (binding) skills. Importantly, this relation became stronger as a function of increased number of monitored countdowns, suggesting that spatial abilities become increasingly important in more complex multitasking situations, which require a higher degree of time monitoring due to multiple overlapping task timelines. An analysis of participants' monitoring behavior further showed that, despite an overall increased monitoring activity, participants reduced their monitoring of each single countdown a relatively long time before the target occurred, whereas monitoring of specific countdowns was not affected by the number of (additionally) monitored countdowns in close temporal proximity to the target (Figure 4). This pattern of monitoring behavior suggests that participants had to rely more on internal representations of the countdowns (rather than monitoring/using external sources of information to retrieve the state of a countdown), when more countdowns had to be monitored. As hypothesized in the introduction, such internal representations may be offloaded onto spatial processes in order to achieve the same level of performance.

Importantly, the behavioral performance data were accompanied by increased neural activation in the precuneus and IPL, in which the BOLD signal increased as a function of task complexity. Both these regions were more strongly activated when participants monitored and responded to a larger number of parallel tasks. Together, these findings support the idea that multiple task timelines are represented spatially, and are in line with the *spatial offloading hypothesis*, by showing that (a) maintaining high behavioral performance is related to individuals' spatial abilities, especially in situations with multiple overlapping task timelines (as in multitask situations), and (b) keeping track of multiple overlapping task timelines involves cortical regions critical for spatial processing (see below).

A secondary goal of this study was to investigate whether processing of task timelines using spatial representations is a general mechanism for time monitoring, or whether such spatial offloading is a mechanism involved in keeping track of task timelines specifically in multitasking situations (Mäntylä, 2013). Here, using a model-based approach, we did not observe significant differences between various models designed to distinguish between predicted neural consequences of these possibilities. In fact, dual- and multi-task specific models, as well as a general (linear increase) model were all associated with increased activation in a similar set of brain regions. For the identified core regions in the inferior parietal cortex and precuneus, a linear increase was the best fit for the data, however, no specific pattern of brain activation could be *exclusively* related to a multitasking-specific mechanism. Therefore, our results do not provide conclusive support for the spatial offloading hypothesis as a mechanism for time monitoring specifically during multitasking. Rather, a spatial offloading should be assumed to be a general mechanism for time monitoring in tasks.

#### Temporal states and spatiotemporal offloading in the posterior parietal cortex and hippocampus

There is much evidence that the posterior parietal cortex is important for spatial processing, including sensory and motor control, spatial cognition, and spatial transformations (Cohen et al., 1996; Goodale & Milner, 1992; Halari et al., 2006; Jordan et al., 2001; Kesner & Long, 1998; Milner & Goodale, 2008; Richter et al., 1997; Sack, 2009; Zacks, 2008). However, the role of the posterior parietal cortex is not restricted to spatial processing per se. In fact, the intraparietal lobe is also critical for integrating spatial and temporal information (Assmus et al., 2003), and overlapping areas in IPL have been shown to process spatial and temporal information (Gijssels et al., 2013). Moreover, processes underlying common magnitude systems are thought to be located in the lateral posterior parietal cortex (Sokolowski, Fias, Bosah Ononye, & Ansari, 2017). Based on these findings, and the observation that cortical activation in IPL increases with the number of tasks performed in parallel, IPL may be involved in spatiotemporal offloading. Specifically, temporal information about the states of multiple tasks may be represented in general magnitude systems, based on spatial processing areas in the lateral posterior parietal cortex.

Temporal information about single, ongoing tasks may be processed and/or maintained in the precuneus. Indeed, there is evidence for a role of precuneus in temporal processing from studies showing that this region is linked to time estimation over long intervals (Morillon, Kell, & Giraud, 2009). Specifically, the precuneus, as part of the default mode network, has been associated with time estimation for relatively long event durations exceeding 2 seconds, whereas shorter intervals are estimated by the motor system (Morillon et al., 2009). Increasing activation in the precuneus with the number of tasks, as observed in the current study, suggests that the precuneus estimates the temporal progress of multiple tasks when necessary.

In addition to regions in the posterior parietal cortex, the hippocampus also showed increased activation during dual- and multitask conditions, compared to single-task condition (Figure 3, 6). In contrast to IPL and precuneus, however, activation in the hippocampi does not increase with additional tasks. The hippocampus is known to be involved in spatial processes including memory, navigation, and mapping (Eichenbaum, 2017).

Taken together, we suggest that the precuneus, left IPL, and the hippocampus are core regions for processing of multiple tasks with overlapping timelines. The precuneus, in line with studies relating the precuneus to temporal processing of long (> 2 s) intervals (Morillon et al., 2009), may be involved in temporal processing of multiple task timelines. The IPL may then form relations between different task timelines, using spatial representations and transformations, thus shifting cognitive load of temporal processing onto spatial processes through spatiotemporal offloading (Mäntylä, 2013; Risko & Gilbert, 2016). The hippocampus, based on the spatiotemporal transformation in IPL, may generate and/or maintain a spatial map of the temporal relations within a multitasking situation, and/or may be involved in storage of multiple of such maps over the course of the experiment. The causal contribution of each of these regions remains to be tested experimentally.

#### Contributions of medial frontal, occipital and temporal cortex

We also observed increased activation outside spatial processing areas of the posterior parietal cortex during the experimental task conditions compared to the condition-specific control conditions, especially in the medial frontal lobe and the hippocampi (Figure 3, 6). However, here we did not observe that activation in frontal cortex scaled with the number of task instances (i.e. monitored countdowns). This may be the case because in the present study participants did not switch between different task rules and task sets, or stimulus features, as is commonly necessary

in dual- and multi-task research (von Bastian & Druey, 2017). Furthermore, we observed increased activation in temporal and occipital regions during the experimental task, compared to the control task (Figure 3). Activation of the temporal cortices may be related to language processing (specifically, processing of letters) and alphabetically structured stimuli (Heinzel et al., 2008; Thesen et al., 2012), considering that temporal areas were activated consistently over all complexity levels, without systematic increases. Whereas the control condition included letters, these did not have to be processed to the same extent. In fact, during the control task, participants only had to recognize the letter *A*, and could ignore any other letter. Finally, stronger activation in visual areas during the experimental task conditions, compared to their respective control conditions, is most likely related to differences in visual attention to the (matched) visual input.

2.

## Interpretational considerations and limitations

Posterior parietal regions such as precuneus and IPL have been related to spatial attention and motor control (Lee et al., 2013; Singh-Curry & Husain, 2009). Even though spatial features were largely controlled for, compared to previous studies on this topic, it can be argued that the different letter series were mapped onto four different fingers. Moreover, in the current task, and in particular in conditions with higher number of parallel countdowns, may have increased the demands for spatial motor coordination in the sense that participants had to perform the task with two hands, and up to four fingers on one of the hands. However, two aspects argue against such an explanation. First, we introduced condition-specific control tasks, which were similar in terms of motoric complexity in the sense that both hands had to be used, and varying numbers of button presses were executed depending on the control condition. Overall, motor output was similar between the experimental and control tasks. Another important aspect that argues against a motor

complexity explanation of posterior parietal activation increases is the lack of activation in other known motor areas, such as premotor and primary motor cortices.

Moreover, it can be argued that the involvement of spatial processes in this task (both in terms of neural activation and correlation with measures of spatial abilities, i.e. mental rotation) were a direct consequence of the task design and the participants' strategies to perform the task. Specifically, one option is that, given the association between (spatially distinct) response buttons and task deadlines/countdowns, task timelines were represented in these spatially distinct locations (e.g. finger locations). To this end, the current study cannot clearly distinguish between a direct neurally implemented spatial offloading mechanism and an alternative embodied offloading strategy. That is, participants may strategically chose to represent the different timelines spatially onto their fingers used for responding. However, we argue that both implementations should be considered forms of a spatial offloading mechanism. Importantly, both mechanism would be realized through recruitment of cortical spatial processing areas, as observed in this study.

The interpretation in this study, that spatial processing areas are involved in keeping track of multiple timelines, is based upon observation of increased activation in areas commonly known for spatial processing. However, given that participants did not complete a spatial processing task (such as the mental rotation task) during fMRI scanning, we were not able to examine whether spatial tasks and the deadline monitoring task used in the current study engage overlapping brain regions. Moreover, the posterior parietal cortex has also been associated with working memory processes, especially working memory updating (Borst & Anderson, 2013; Wager & Smith, 2003). It is likely that working memory demands also increase with the number of monitored countdowns (i.e. participants need to keep track of a higher number of letters). Therefore, additional research is required where spatial tasks and working memory tasks are tested in the same participants as time monitoring tasks using functional neuroimaging techniques.

#### Conclusion

This study provides evidence that the precuneus represents timelines of multiple tasks and the IPL is involved in building spatial representations of the temporal relations of these overlapping task timelines. These results are in line with the spatiotemporal offloading hypothesis of multitasking, suggesting that spatial processes are used to resolve temporally demanding tasks when dealing with overlapping task timelines. Therefore, this study enhances our understanding of the neurocognitive mechanisms underlying everyday performance with multiple tasks and overlapping deadlines.

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to Review Only



Figure 1: Experimental task. Panel A: Timeline of a single countdown series and corresponding responses. Participants responded on two separate response boxes, one with four monitor buttons (right hand), and one with a single target button (left hand). The task is presented on a screen. The lower part of the screen contains information about the active countdowns (colored boxes). Countdowns are run down the alphabet, but are generally hidden from view. Letter countdowns were displayed in the upper part of the screen for 1 second upon pressing of one of the four corresponding monitor buttons (screen 2; green series displayed upon pressing of the green monitor button, at countdown state "H''). After 1 second, the countdown is covered again (3rd screen), until a monitor button is pressed again (5th screen, "A"). When the countdown reaches the target letter (A), participants need to press the target button (last screen, single button response box). Panel B: Schematic representation of a task sequence. Up to four letter series ran, hidden from view (represented as grey letters), at different updating rates and with varying starting letters. Pressing a color-coded monitor button (4-button response box, right) displays the corresponding series for a short period (color-coded frames, black letters). When a series runs down to A, the target button (single button response box, right) needs to be pressed. In this example, targets of the blue and yellow letter countdowns were missed (red circles) while a target of the red letter countdown was correctly indicated (green circle).

176x124mm (300 x 300 DPI)



**Figure 2: Models used to identify brain regions following specific activation patterns depending on the number of parallel tasks.** Predicted BOLD signal intensity depending on the number of parallel tasks for different tested models, i.e. linear increase model (A), dual-task step model (B; brain regions recruited for dual-task+ situations), multi-task (3+) step model (C; brain regions recruited for multitasking situations with 3 and more tasks, but not dual-tasking).





**Figure 4: Monitoring behavior** (A) Monitoring actions by condition/number of monitored counters (B) Monitoring behavior relative to target occurrence, depending on number of monitored countdowns. Thick lines represent monitoring actions binned in 500 ms windows, relative to target occurrence (up to 6 seconds before target occurrence). Thin lines represent standard errors of the mean. Asterisks indicate time bins with significant differences between conditions (F-test, 1-way ANOVA; p < .01). (C) Ratio between early [bin 1-6 (5.5-3 seconds before target occurrence)] and late [bin 10-12, 1.5-0 seconds before target occurrence] time bins, depending on number of monitored countdowns. Error bars indicate standard error of the mean.

1293x347mm (96 x 96 DPI)



(4 countdowns). The reverse contrasts [control – experimental] did not result in any significant clusters.

176x152mm (300 x 300 DPI)





#### Figure 7: Parameter estimates of selected regions from model activation maps for illustration.

Post-hoc comparisons of the experimental vs control task condition (i.e. parameter estimates <> 0) indicate that none of the parameter estimates was significantly smaller than zero (all p>.05, uncorrected), indicating that the control condition did not result in stronger activation than the experimental condition for any of the conditions, within the ROIs. Error bars indicate standard errors of the mean.

170x53mm (300 x 300 DPI)



#### Tables

**Table 1: Condition-specific comparisons.** Clusters for comparisons with significant activation differences are listed (cluster forming threshold: z=3.1; measure: cluster mass). FWE-corrected cluster level p-values are shown, based on permutation testing (PALM, see methods).

Contrast/region	voxels	Z max	X [mm]	Y [mm]	Z [mm]	Cluster p
4>1						
L anterior IPS/IPL	96	4.932	-34	-62	35	.018
R post. cingulate gyrus	78	4.279	2	-36	32	.019
R frontal medial cortex	64	5.299	4	52	-20	.021
R middle temporal gyrus	60	5.031	54	-4	-20	.021
L/R precuneus	60	4.310	-2	-54	38	.022
L occipital pole (V2)	58	4.624	-26	-100	-10	.022
L occipital pole (V2)	50	4.519	-10	-88	32	.024
R anterior IPS/IPL	47	4.147	36	-64	44	.024
R post. cingulate gyrus	46	4.625	4	-50	8	.024
R hippocampus	35	4.857	26	-26	-18	.028
L middle temporal gyrus	32	4.605	-62	-42	4	.030
L IPL	32	5.080	-52	-74	26	.030
L post, cingulate gyrus	29	4.556	-6	-44	0	.033
R occipital pole (V2)	28	5.248	18	-88	-10	.031
R middle temporal gyrus	25	4.083	64	-4	-24	.042
3>1				5		
L/R precuneus	80	4.758	2	-54	34	.050
				T		

Table 2: model specific activation patterns. Clusters for each model are listed (cluster forming threshold: z=3.1; measure: cluster mass). FWE-corrected cluster level p-values are shown, based on permutation testing (PALM, see methods). Significant clusters (p<.05) are printed in bold.

#### Table 2a: Linear increase model

Region	voxels	Z max	X [mm]	Y [mm]	Z [mm]	Cluster p
L inf parietal	308	6.81	-34	-62	36	0.013
L/R precuneus	132	4.88	-2	-54	38	0.056
R mid temporal	118	5.22	54	-6	-20	0.063
R mid cingulum	75	4.26	2	-36	32	0.101
R inf medial frontal lobe	60	5.21	4	52	-20	0.117
R angular g	54	4.47	36	-64	44	0.131
L angular g	50	6.91	-52	-74	26	0.133
cerebellum vermis	26	4.27	2	-52	10	0.226
R lingual	14	5.46	18	-88	-8	0.441

## Table 2b: dual-task step model

K illigual	τT	5.40	10	00	0	0.771				
Table 2b: dual-task step model										
Region	voxels	Z max	X [mm]	Y [mm]	Z [mm]	Cluster p				
L inf parietal	104	5.50	-32	-72	46	0.039				
L/R precuneus	54	4.29	-2	-54	38	0.060				
L inf medial frontal lobe	33	4.76	-2	46	-26	0.082				
R mid temporal	24	5.19	54	2	-26	0.106				
R post cingulum	24	4.15	2	-34	30	0.117				

## *Table 2c: multi-task (3+) step model*

Region	voxels	Z max	X [mm]	Y [mm]	Z [mm]	Cluster p
L/R precuneus	257	5.08	4	-54	34	0.023
L inf parietal	148	4.79	-34	-74	44	0.045
R mid temporal	116	5.01	52	-10	-18	0.057
R post cingulum	52	4.40	10	-36	30	0.112
R mid frontal g	50	4.41	36	22	44	0.114

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L inf tri frontal	45	5.00	-54	34	10	0.118
L angular	39	5.89	-52	-74	26	0.132
L mid frontal	32	4.43	-22	20	38	0.156
R angular	22	3.97	42	-68	48	0.24

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