Involuntary shifts of spatial attention contribute to behavioral crossmodal
distraction - Evidence from oscillatory alpha power and reaction time data

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Abstract

Imagine you are focusing on the traffic on a busy street to ride your bike safely when suddenly you hear an ambulance siren. This unexpected sound involuntarily captures your attention and interferes with ongoing performance. We tested whether this type of behavioral distraction involves a spatial shift of attention. We measured behavioral distraction and magnetoencephalographic alpha power during a crossmodal paradigm that combined an exogenous cueing task and a distraction task. In each trial, a task-irrelevant sound preceded a visual target (left or right). The sound was usually the same animal sound (i.e., standard sound). Rarely, it was replaced by an unexpected environmental sound (i.e., deviant sound). Fifty percent of the deviants occurred on the same side as the target, and 50% occurred on the opposite side. Participants responded to the location of the target. As expected, responses were slower to targets that followed a deviant compared to a standard, reflecting behavioral distraction. Crucially, this distraction was mitigated by the spatial relationship between the targets and the deviants: responses were faster when targets followed deviants on the same versus different side, indexing a spatial shift of attention. This was further corroborated by a posterior alpha power modulation that was higher in the hemisphere ipsilateral (vs. contralateral) to the location of the attention-capturing deviant. We suggest that this alpha power lateralization reflects a spatial attention bias. Overall, our data support the contention that spatial shifts of attention contribute to behavioral deviant distraction.

Keywords: alpha oscillations, deviant distraction, involuntary attention, magnetoencephalography, spatial attention
1. Introduction

Efficient cognitive functioning requires the ability to filter out task-irrelevant events to effectively perform a task at hand. On the other hand, it may also be adaptive to detect unexpected but behaviorally relevant changes in the soundscape. Hence, selective attention and change detection must be carefully balanced to limit the negative impact of task-irrelevant stimuli while allowing for potentially important stimuli to break through attentional filters. One upshot of this balance is the cognitive system’s vulnerability to distraction. For example, task-irrelevant events, such as an ambulance siren while you are riding your bike, may distract you and affect driving performance.

In an experimental setting, unexpected task-irrelevant events, so-called deviants, usually violate sensory predictions by deviating from an otherwise (more or less) regular sound sequence consisting of so-called standards. Distraction due to auditory deviants has been extensively studied via (e.g., Bendixen et al., 2007; Schröger & Wolff, 1998; Wetzel et al., 2012). It is well known that deviant distraction is accompanied by distinct event-related markers (for reviews, see Escera et al., 2000; Parmentier, 2014) and a behavioral effect (referred to as behavioral distraction hereafter) observed when unexpected sounds violate sensory predictions (e.g., Bendixen et al., 2008; Parmentier et al., 2011). Behavioral distraction is defined as the slowing of responses during the primary task following the presentation of a task-distracting event compared to standard events. This effect is thought to result from an involuntary (stimulus-driven, bottom-up, or exogenous) shift of attention from the task towards the deviant and back to the task, which results in a time penalty (Parmentier et al., 2008; Schröger, 1996) and appears to be related to a transient inhibition of action (Dutra et al., 2018; Vasilev et al., 2019; Wessel and Aron, 2017). In crossmodal scenarios in which the auditory deviant and the visual target occur at different locations, at least two determinants might contribute to distraction (Parmentier et al., 2008; Parmentier, 2014). 1) One is the shift of attention from one modality to the other, i.e. from vision to audition. 2) The second one is a shift of attention from one spatial location to another. Yet the contribution of a spatial shift of attention to distraction has so far not been tested systematically. Also, the exact mechanisms underpinning distraction remain relatively poorly understood. The spatial shift hypothesis posits that deviant sounds capture attention and trigger an
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involuntary shift of attention to the location of the deviant sound, and from there to the target’s location upon its appearance. Because spatial shifts require time to complete (Cheal & Gregory, 1997; Luck et al., 1996; Shiu & Pashler, 1994), they might contribute to behavioral distraction from deviant sounds (Parmentier, 2014).

Still, support for the hypothesis of a spatial shift of involuntary attention is currently lacking. The few reports relevant to this topic cannot rule out alternative (though not mutually exclusive) interpretations (Parmentier et al., 2008), or present inconsistent results (Corral & Escera, 2008). However, indirect support can be found in behavioral data outside of the deviant distraction literature. This is for example the case in studies that applied an exogenous spatial cueing paradigm in which a task-irrelevant sound, such as a noise burst, appeared lateralized (left or right) briefly before the presentation of an ipsi- or contralateral visual target. While the location of the sound does not predict the location or the type of upcoming visual target, results show a performance advantage when cue and visual target appeared on the same side (Dufour, 1999; Feng et al., 2014; McDonald et al., 2000; Spence & Driver, 1997). This pattern of results indicates that cues are effective in generating an involuntary shift of spatial attention.

Some of the studies showing behavioral evidence for an involuntary shift of spatial attention draw further support from electroencephalographic (EEG) data (Feng et al., 2017; Störmer et al., 2016). Focusing on oscillatory activity, the authors found a sound-induced modulation in posterior alpha power that built up rapidly and was rather short-lived, thus resembling the properties of involuntary attention (Corbetta & Shulman, 2002). Crucially, the posterior alpha power modulation was lateralized. That is, the alpha power is higher in the hemisphere ipsilateral to the location of the attention-capturing sound (compared to contralateral alpha power). In fact, the posterior alpha power lateralization is a well-known finding from research on voluntary (goal-directed, top-down, or endogenous) attention that has been linked to spatial attention bias (e.g., Banerjee et al., 2011; Deng et al., 2019; Rihs et al., 2009; Sauseng et al., 2005; Thut et al., 2006; Worden et al., 2000; Wöstmann et al., 2016). Research on voluntary attention typically uses endogenous spatial cueing tasks, in which cues require participants’ to shift their attention to certain target locations (left or right). Irrespective
of whether spatial attention is voluntarily directed to a visual or auditory target, sustained oscillatory
alpha power appears to be lateralized, especially over parieto-occipital areas. This lateralization is
often driven by an ipsilateral increase and/or a contralateral decrease in alpha power relative to the
attended side, thought to reflect the inhibition of brain regions that process task-irrelevant information
at unattended locations, or facilitate processing of task-relevant information at attended locations,
respectively (for reviews, see Hanslmayr, 2011; Klimesch, 2012; Peylo et al., in press).

**Current approach**

To test the possible role of a spatial shift of attention in deviance distraction, we employed a
crossmodal paradigm combining auditory distraction and an exogenous spatial cueing task (Figure 1).
In each trial, a task-irrelevant sound preceded a visual target. In most trials (80%), the same standard
sound was used. In the remaining trials (20%), novel sounds were used. As in an exogenous spatial
cueing task, the location of the deviant was non-informative about the spatial location of the
upcoming target and could either be congruent or incongruent with it (with equal probabilities). That
is, 50% of the deviants were presented on the same side as the upcoming visual target (Congruent
condition), while the remaining 50% of the deviants were presented on the side opposite to the
upcoming visual target (Incongruent condition). Participants were instructed to respond to the visual
target-s location (left vs right).

At the behavioral level, our predictions were two-fold. First, we expected to find longer reaction times
(RTs) to visual targets following a deviant compared to a standard, reflecting an involuntary shift to
and from the deviant (Escera et al., 1998; Parmentier et al., 2008; Parmentier, 2014; Ruhnau et al.,
2013). Secondly, and more crucially, we predicted a *spatial* congruence effect whereby RTs would be
shorter for congruent than incongruent trials.

To examine brain activity, we recorded MEG data to shed light on the neural oscillatory activity at
play in our task, with an emphasis on alpha power and its link to involuntary spatial attention. We
predicted that the deviant would induce a lateralized, short-lived, alpha power modulation in
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parieto-occipital areas that would build up rapidly after stimulus onset, thereby matching the characteristics of involuntary spatial attention (Feng et al., 2017; Störmer et al., 2016).

2. Methods

The study was approved by the local ethics committee of the Paris Lodron Universität Salzburg and adhered to the code of ethics of the World Medical Association (Declaration of Helsinki).

2.1 Participants

30 healthy young adults participated in the MEG experiment (12 males, mean age ± STD: 27 ± 6 yrs, 4 left-handed, 1 ambidextrous). Volunteers gave written, informed consent prior to their participation. All participants reported normal hearing and normal or corrected-to-normal vision, and none reported a history of neurological or psychiatric diseases. Volunteers received 10€/hour (h) or 1 credit point/h as compensation for their participation.

2.2 Stimuli and Procedure

The experiment was programmed in MATLAB 9.1 (The MathWorks, Natick, Massachusetts, U.S.A) using the open-source Psychophysics Toolbox (Brainard 1997; Kleiner et al. 2007) and o_ptb, an additional class-based abstraction layer on top of the Psychophysics Toolbox (Hartmann and Weisz, 2020; https://gitlab.com/thht/o_ptb). Stimuli were presented with precise timing using the VPixx System (DATAPixx2 display driver, PROPixx DLP LED Projector, TOUCHPixx response box by VPixx Technologies, Canada). The Blackbox2 Toolkit (The Black Box ToolKit Ltd, Sheffield, UK) was used to measure and correct for timing inaccuracies between triggers and the visual and auditory stimulation. Note that the tubes of the MEG-proof sound system caused a time delay between sound and trigger of 16.5 ms that was corrected within the stimulation protocol.
(A) Trial sequence

Left  ▪  S S D S S D S S S ...
Right ▪  S S S S S S D S

Legend

D  Deviant (10% left, 10% right)
S  Standard (80%, binaural)

(B) Exemplary deviant trial

(C) Locations of deviant and target in different conditions

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Figure 1: Cross-modal paradigm combining a deviant distraction task with an exogenous spatial cueing task. (A) Exemplary trial sequence. Trials could either be standard trials or deviant trials depending on whether the task-irrelevant sound was a binaurally presented standard sound (p = 0.8) or a lateralized deviant sound presented either left (p = 0.1) or right (p = 0.1). (B) Exemplary deviant trial. During each trial, the task-irrelevant sound - here a deviant sound, illustrated as an orange bar - preceded the task-relevant visual stimulus, illustrated as a green bar. Participants had to indicate whether the target occurred to the left or right of the fixation frog. (C) Visual representation of the two spatial conditions (Congruent and Incongruent) and the corresponding spatial locations (Left, Right) of deviants and targets.

We employed a cross-modal paradigm combining a distraction task with an exogenous spatial cueing task. Participants performed a visual-spatial two-alternative forced-choice task (see Figure 1).

Throughout the task, a frog was displayed in the center of the screen that participants were asked to
fixate on at all times. In every trial, a visual target was presented that consisted of the shape of a fly featured next to the frog. This visual target appeared on the left or on the right of the frog with equal probability (the order of these trials was random for every participant). Participants were instructed to hold their left and right thumbs over left/right response buttons respectively and to press these buttons to indicate the side where the visual target (fly) had been presented. The mapping between the response buttons and the target location was counterbalanced across participants. The target remained on the screen until a response was recorded or for a period of 1 s (whichever occurred first). A response within this time window terminated the current trial and initiated the next trial.

Apart from the visual target, each trial also involved the presentation of an auditory stimulus, which participants were instructed to ignore. This sound of 0.5 s duration (including 10 ms rise and fall times) was always presented after a 0.8 s silent interval at the start of each trial, and its onset preceded that of the visual target by 0.6 s. That is, a silent interval of 0.1 s separated the offset of the sound from the onset of the visual target. This timing was selected based on earlier work (Ruhnau et al., 2013) and because short stimulus onset asynchrony is best suited to study distraction (Bendixen et al., 2010). The sound was either a binaurally presented standard (buzzing mosquito, p = .8) or a monaurally presented deviant (p = .2). Deviants were chosen from a pool of 56 different environmental sounds (e.g., speech, animal voices, tool noises, etc.) selected from a commercial CD (1,111 Geräusche, Döbeler Cooperations, Hamburg, Germany). All sounds were equalized for overall intensity (RMS). The order of the trials was pseudorandomized such that deviant trials occurred randomly with the constraint that each deviant trial was preceded by at least two standard trials.

Two types of deviant trials, equiprobable (p = .1) were defined. In the Congruent condition, the deviant sound and visual target were presented on the same side (left or right, with equal probabilities). In the Incongruent condition, they were on opposite sides (left/right vs. right/left, with equal probabilities). Each of the 56 deviant sounds occurred only once within a block of trials, and a total of four times in the whole experiment - twice on the left (once in the Incongruent and once in the
Congruent condition) and twice on the right (once in the Incongruent and once in the Congruent condition).

The experiment consisted of seven blocks in total, each lasting approximately 5 min. and consisting of 160 trials (128 standards, 16 deviants presented on the left, 16 deviants presented on the right). Thus, the time spent on task (excluding breaks between blocks) could be up to approximately 35 min. The auditory stimuli were presented at 50 dB above the sensation threshold level. The auditory stimulation was delivered via MEG-compatible pneumatic in-ear headphones (SOUNDPixx, VPixx technologies, Canada). The visual stimuli were presented inside of the magnetically shielded room using a projector (PROPixx DLP LED Projector), via a back-projection screen and a mirror system.

Before the start of the experiment, each participant's individual sensation threshold was determined using a 200-ms 1000 Hertz (Hz) tone presented via a bayesian threshold method (Kontsevich & Tyler, 1999; Sanchez et al., 2016) utilizing the VBA toolbox (Daunizeau et al., 2014). The sensation threshold was measured separately for the right and left ears. Thereafter, participants practiced the task with fewer trials to familiarize themselves with the experiment. Novels were chosen randomly from the pool of environmental sounds (N=56). These were played backward so that participants did not become familiar with those sounds.

2.3 MEG recording and preprocessing

Before beginning the MEG recording, five head position indicator (HPI) coils were placed on the left and right mastoid, on the upper right and lower right forehead, and on the left middle forehead. Anatomical landmarks (nasion and left/right preauricular points), the HPI locations, and at least 300 headshape points were sampled using a Polhemus FASTTRAK digitizer. Six pre-gelled and self-adhesive Ag/AgCl electrodes (Ambu Neuroline 720) were attached to record the vertical and horizontal electrooculogram (EOG) and the electrocardiogram (ECG). One additional electrode - placed at the right shoulder - served as a ground.
Participants were in a seated position during data recording. A chin rest was used to avoid head movements during and across different blocks of recording. The eye, heart, and magnetic signals of the brain were recorded at 1000 Hz (hardware filters: 0.03 - 330 Hz) in a standard passive magnetically shielded room (AK3b, Vacuumschmelze, Germany) using a whole-head MEG (Elekta Neuromag Triux, Elekta Oy, Finland). Signals were captured by 102 magnetometers and 204 orthogonally placed planar gradiometers at 102 different positions.

MEG data were preprocessed and analyzed offline using Fieldtrip (Oostenveld et al., 2011), an open-source toolbox for Matlab (www.mathworks.com), and custom-written functions. Maxfilter (version 2.2.15, Elekta Neuromag, Finland) was applied to the continuous MEG raw data using a signal space separation algorithm (Taulu & Kajola, 2005; Taulu & Simola, 2006) provided by the MEG manufacturer. This allowed us to 1) automatically remove and interpolate the data of bad channels; 2) remove artifacts from the MEG signal (50 Hz line-noise and its harmonics, 16.7 Hz noise from a passing train, and muscle activity with origins beyond the head); and 3) compensate, offline, for changes in head position across blocks by realigning the data to a common standard head position (-trans default Maxfilter parameter) based on the head position measured at the beginning of each block. Thereafter, continuous data were visually inspected and sample points of extensive artifacts were stored separately for each block and participant.

Two different pipelines were used for further preprocessing of the MEG data. First, preprocessing of the data was optimized for independent component analysis (ICA). Therefore, MEG data were filtered offline using a 100 Hz lowpass filter (sinc FIR, kaiser window, order 3624, cutoff (-6 dB) 100 Hz) and a 1-Hz highpass filter (sinc FIR, kaiser window, order 908, cutoff (-6 dB) 1 Hz). After rejecting sample points covering extensive artifacts, runica ICA was applied in order to identify components originating from eye movements and heartbeat. The second preprocessing pipeline was optimized for further data analysis. Specifically, cleaned MEG data were filtered offline using a 35 Hz lowpass filter (sinc FIR, kaiser window, order 3624, cutoff (-6 dB) 35 Hz). Data were downsampled to 256 Hz.
Thereafter, a 0.1 Hz highpass filter was applied (sinc FIR, kaiser window, order 9056, cutoff (-6 dB) 0.1 Hz).

Filtered continuous data (originating from the second preprocessing pipeline) were cleaned from the ICs (on average two per participant). Data were segmented into 3-s epochs (-1.5 s to 1.5 s time-locked to the onset of the sound) and cleaned from artifacts that were identified via visual inspection (see above). For further analysis, we used deviant trials only. Trials in which deviants occurred on the left or right are, hereafter, referred to as Deviant Left or Deviant Right, respectively. Only those deviant trials for which participants responded once, correctly, and within the predefined response window (100 to 800 ms following the target), entered the analysis. This resulted in 106.53/112 (SD = 3.38) Deviant Left trials and 106.93/112 (SD = 2.89) Deviant Right trials, on average.

2.4 Data analyses

2.4.1 Behavioral Analysis

The following trials were excluded from the analysis of behavioral data: trials without responses, trials including responses faster than 100 ms or slower than 800 ms after target onset, and trials that included more than one button press. Individual RTs and hit rates were calculated. This resulted in ≥55.07/56 (SEM ≥ 0.90 ≤ 1.81) Congruent Left / Congruent Right / Incongruent Left /Incongruent Right trials and ≥109.83/112 (SEM ≥ 1.70 ≤ 2.44) Deviant Right / Standard-before-Deviant-Right / Deviant Left / Standard-before-Deviant-Left trials, on average.

2.4.2 Statistical analyses. We used jamovi (version 1.2.22.0, https://www.jamovi.org/) for the statistical analyses of behavioral data. To determine the behavioral distraction effect on RTs and hit rates, we applied two-way repeated-measures analyses of variance (ANOVAs) using the factors sound type (deviant, standard) and deviant location (left, right). Note that we used standards before the corresponding deviants for the analyses. To determine the behavioral effect of the spatial shift of attention on RTs and hit rates, we applied two-way repeated-measures ANOVAs using the factors
congruence type (congruent, incongruent) and deviant side (left, right). Note that for statistical purposes, hit rates were transformed to rationalized arcsine units (rau) to account for non-normality of proportion data (Studebaker, 1985). We reported the generalized eta-squared ($\eta^2_G$; Bakeman, 2005) as a measure of effect size. Where appropriate, we reported mean values (M) and the standard error of the mean (SEM) in the format $M \pm SEM$.

2.5 MEG Data Analyses

For the analyses of MEG data in sensor and source space, we solely used the signal recorded at the gradiometers. Data were detrended to remove slow frequency shifts.

2.5.1 Sensor space analyses.

2.5.1.1 Time-frequency analysis.

Time-frequency analysis was performed on the preprocessed single-trial data between 1 and 30 Hz (1 Hz steps) using Morlet wavelets with a width of 5 cycles, for each participant and condition, separately. The analysis time window was shifted in steps of 50 ms between -1.5 to 1.5 s relative to deviant onset. Note that power was first computed for each individual gradiometer. In a second step, the power for each gradiometer pair was summed in order to yield an orientation independent estimate. Thereafter, single-trial spectral power was log-transformed with a base of 10. For each participant, trials were averaged for Deviant Left and Deviant Right, separately.

In order to study the involuntary shift of spatial attention, we followed a methodological approach applied in research on voluntary spatial attention, in which alpha power was contrasted for the cued / attended target positions, namely attend left versus attend right (Banerjee et al., 2011; Deng et al., 2020; Haegens et al., 2011; Wöstmann et al., 2016). Accordingly, here we contrasted alpha power for the left and right attention-capturing, task-distracting deviants (Deviant Left versus Deviant Right). In order to statistically examine alpha power modulations dependent on the deviant side in both hemispheres, we performed non-parametric, cluster-based dependent-samples t-tests, with Monte-Carlo randomization (Maris & Oostenveld, 2007). The tests were restricted to the alpha
frequency range (8 – 14 Hz) and the time window ranging from 0.2 to 0.6 s following the deviant sound and preceding the target (cf. Feng et al., 2017; Störmer et al., 2016; Weise et al., 2016). Since we had a priori hypotheses concerning the direction of the lateralized alpha power modulations expected in each hemisphere, we applied paired-sample, one-tailed Student’s t-tests for the left-hemispheric channels and the right-hemispheric channels, separately. The Student’s t-tests were applied for each sensor-time-frequency bin separately for each hemisphere by testing the alpha power values of Deviant Left against the alpha power values of Deviant Right. Based on this, clusters were formed among neighboring sensors of the respective hemisphere and corresponding time-frequency bins that exceeded an a priori chosen threshold (p = 0.05). The sum of the t-values within the respective cluster was used as the cluster-level statistic. The cluster with the maximum sum was subsequently used as the test statistic. By permuting the data between the two conditions and recalculating the test statistic 10,000 times, we obtained a reference distribution of maximum cluster t-values to evaluate the significance of the to-be-tested contrast. The power distributions between the two conditions were considered to be significantly different from each other if no more than 5% of the sum-t-values of all of the 10,000-permuted data were larger than the sum-t-value of the to-be-tested cluster. Note that these tests were not performed on baseline-corrected data.

In addition, we ran the time-frequency analysis a second time using the same parameters as described above, but with the preceding step that the evoked signal averaged across trials of the respective condition was subtracted in the time domain from every single trial, to increase sensitivity to non-phase locked signals, and to reduce contamination by phase-locked signals. The results are comparable to those obtained via the time-frequency analysis of phase-locked activity and can be found in the supplementary material (Figure S1).

We further investigated the cause of the alpha power modulation observed in the data (Figure 3). Therefore, we analyzed the time courses of ipsi- and contralateral alpha power for the right centro-temporal and the left parieto-occipital region of interest (ROI), respectively, that had been identified via the preceding cluster-based permutation analysis (see above). Each ROI consisted of
three channels that showed prominent deviant-related effects in the deviant–target interval. Data were pooled within each ROI and across frequencies (8 – 14 Hz). Alpha power was submitted to non-parametric, cluster-based, dependent-samples t-tests with Monte-Carlo randomization (n = 10,000), contrasting the data at each sample point in the deviant-target interval (0.2 to 0.6 s following the deviant) with the mean alpha power of the baseline period (-0.4 to -0.2 s; Feng et al., 2017). Since we had no a priori hypotheses on the direction of the alpha power modulations expected in the respective ROI, we applied paired-sample, two-tailed Student’s t-tests for the cluster analysis (see above).

Note that for the analysis of alpha power we were exclusively interested in the activity following the deviant sound reflecting involuntary shifts of spatial attention towards the sound location. Accordingly, we did not expect differences between congruent and incongruent trials as the sounds were not predictive about the target location. Thus, there was no intent to contrast alpha activity between congruent and incongruent trials.

Also, note that the paradigm was not designed - and as a consequence is not suitable - to analyze alpha power between deviant and standard trials given that deviants were presented lateralized while standards were presented binaural.

2.5.2 Source space analyses.

2.5.2.1 Source projection of time series data.

For each participant, realistically shaped, single-shell head models (Nolte, 2003) were computed by co-registering the participants’ head shapes either with their structural MRI (18 participants) or —when no individual MRI was available (12 participants)— with a standard brain from the Montreal Neurological Institute (MNI, Montreal, Canada), warped to the individual head shape. A grid with 3-millimeter resolution based on an MNI template brain was morphed to fit the brain volume of each participant. The leadfield was calculated for each grid point and the forward model was computed for gradiometers only.
To project the preprocessed single-trial sensor data into source space (i.e., to the points of the grid), we applied the linearly constrained minimum variance (LCMV) spatial filter (Van Veen et al., 1997). We followed a procedure described for single virtual sensors (http://www.fieldtriptoolbox.org/tutorial/virtual_sensors/) and extended it to 146,400 points covering the gray matter of the brain. The covariance matrix across all trials (including Deviant Left and Deviant Right conditions) was calculated and used to obtain a LCMV beamformer spatial filter for each of the grid points (for a similar approach, see Neuling et al., 2015; Ruhnau et al., 2016). The covariance window for the calculation of the beamformer filter was based on a 2-s time window centered at deviant onset. To optimize the analysis in source space (i.e., increase spatial resolution using a high-definition grid and at the same time compensate for computation time), we divided the individual brain into 333 parcels (Gordon et al., 2016; http://www.nil.wustl.edu/labs/petersen/Resources.html) based on the ‘Anatomical Automatic Labeling’ [AAL] atlas (Tzourio-Mazoyer et al., 2002; provided by the fieldtrip toolbox). That is, for each anatomical area, the spatial filters of the corresponding grid points were averaged to obtain one spatial filter per parcel. The resulting ‘averaged filter’ was then multiplied with the sensor level data to obtain time-series data for each of the 333 parcels. A regularization parameter (lambda = 5%) was applied.

2.5.2.3 Time-frequency analysis.

For each participant, time-frequency analysis was performed on the single-trial data in source space between 1 and 30 Hz (1 Hz steps) using Morlet wavelets with a width of 5 cycles. The analysis time window was shifted in steps of 50 ms between -1.5 to 1.5 s relative to deviant onset. For each participant, trials were averaged separately for Deviant Left and Deviant Right.

To localize the deviant-induced sensor level effects in source space, alpha power was averaged over frequency (8 -14 Hz) and time (0.2 - 0.6 s). T values were calculated via a paired-sample, one-tailed Student’s t-test for each of the parcels and the corresponding time-frequency bin, contrasting the alpha
power values between Deviant Left and Deviant Right. The respective labels of localized brain regions were identified with an anatomical brain atlas (AAL atlas; Tzourio-Mazoyer et al., 2002) and a network parcellation atlas (Gordon et al., 2016).

As with the sensor space analysis, we ran the time-frequency analysis and corresponding statistics a second time using the same parameters described above, but with the preceding step that the evoked signal averaged across trials of the respective condition was subtracted in the time domain from every single trial. The results are comparable to those obtained via the time-frequency analysis of phase-locked activity and can be found in the supplementary material (Figure S1).

3. Results

3.1 Behavioral data

In terms of accuracy, task performance was generally very good, as indicated by the high hit rate (>0.97) for targets, regardless of whether they were preceded by deviants or standards (Figure 2A). The two-way repeated measures ANOVA on hit rates shows a significant main effect for factor stimulus type (F(1, 29) = 18.51, p < .001, $\eta^2_G = .087$), with higher hit rates for deviant trials (0.99 ± 0.003) than standard trials (0.97 ± 0.004). Neither was there a significant main effect for the factor deviant side (F(1, 29) = 1.09, p = .304, $\eta^2_G = .005$), or a significant interaction (F(1, 29) = .37, p = .548, $\eta^2_G = .001$). This result may be linked to an increase in nonspecific arousal upon the occurrence of a deviant (Näätänen, 1992), which nevertheless facilitates task performance. Hit rates did not differ when deviant and target occurred on the congruent vs. incongruent side (Figure 2B), as indicated by the two-way repeated-measures ANOVA yielding neither significant main effects for the factors congruence type (F(1, 29) = 2.44, p = .129, $\eta^2_G = .017$) and deviant side (F(1, 29) = .13, p = .726, $\eta^2_G = .001$), or a significant interaction (F(1, 29) = .06, p = .815, $\eta^2_G < .001$).

As depicted in Figure 2A, RTs were prolonged when targets followed a deviant versus a standard. This is substantiated by the two-way repeated measures ANOVA on RTs, yielding a significant main
effect for stimulus type ($F(1, 29) = 39.68, p < .001 \eta^2_G = .037$; deviants: 296 ± 9 ms; standards: 278 ± 8 ms). There was neither a significant main effect for the factor deviant side ($F(1, 29) = .78, p = .385, \eta^2_G < .001$), or a significant interaction ($F(1, 29) = 2.153, p = .153, \eta^2_G < .001$). Overall, these results point to a behavioral distraction effect due to an attention shift towards the task-distracting deviant. As depicted in Figure 2B, RTs were shorter when deviant and target occurred on the congruent versus incongruent side. This is substantiated by the two-way repeated measures ANOVA on RTs, yielding a significant main effect for congruence type ($F(1, 29) = 4.59, p = .041 \eta^2_G = .002$; congruent: 293 ± 9 ms; incongruent: 298 ± 10 ms). There was neither a significant main effect for the factor deviant side ($F(1, 29) = .00, p = .963, \eta^2_G < .001$), or a significant interaction ($F(1, 29) = .95, p = .337, \eta^2_G = .004$). Overall, these results point to a spatial shift of attention towards the task-distracting deviant sound.
**Figure 2:** Boxplots of RTs and hit rates in response to visual targets to determine a behavioral distraction effect (left) or a shift of spatial attention (right) caused by the deviant. Grey dots represent individual data. (Left) RTs and hit rates for targets following deviants presented on the left or right.
side versus the standard presented before the corresponding deviant. (Right) RTs and hit rate for targets following deviants on the same versus different side in the Incongruent and Congruent conditions. Data were analyzed with respect to the deviant side (left, right).

3.2 Alpha power analysis

We analyzed alpha-band (8 - 14 Hz) power to probe whether attentional reorienting towards the location of the deviant sound is also reflected in neural oscillatory activity, as indexed via the current RT data. We followed a methodological approach used to measure voluntary attention in which alpha power for the cued / attended target positions had been contrasted (attend left versus attend right; Banerjee et al., 2011; Deng et al., 2020; Haegens et al., 2011; Wöstmann et al., 2016). Accordingly, we contrasted alpha power for the left and right attention-capturing deviants (Deviant Left condition vs. Deviant Right condition). Since we had a priori hypotheses concerning the direction of the lateralized alpha power modulations expected in each hemisphere, we applied the corresponding, non-parametric, cluster-based permutation statistics (using paired-sample, one-tailed Student’s t-tests) on the left-hemispheric and the right-hemispheric channels, separately. Overall, the results clearly show that alpha power to lateralized deviants is modulated in a spatially selective manner (Figure 3).

Specifically, the results for the left hemispheric channels show that alpha power is significantly higher ipsilateral to the deviant location in the time window from 0.2 s to 0.6 s following the auditory deviant and preceding the visual target (see Figure 3 A). This effect is most pronounced over left parieto-occipital sensors (p = .023), as highlighted via the scalp distribution of the alpha power modulation (see Figure 3 C). The main generators of this sensor level effect are estimated in parieto-occipital regions (see Figure 3 D).

The results for the right hemispheric channels show that alpha power is significantly lower contralateral to the deviant location in the time window 0.2 s to 0.6 s following the auditory deviant and preceding the visual target (see Figure 3 B). This effect is most pronounced over right centro-temporal sensors (p = .012), as highlighted via the scalp distribution of the alpha power
modulation (see Figure 3 C). The main generators of the sensor level effect are estimated in fronto-parietal regions (see Figure 3 D).

Note that our analysis of non-phase-locked power yielded comparable results (Figure S1).

**Figure 3:** Log10-transformed alpha power modulation to lateralized deviants based on
group-averaged data for Deviant Left minus Deviant Right. (A and B) Time-frequency representations highlighting (A) high alpha power ipsilateral to the deviant location at left-hemispheric posterior gradiometers (average across three channels showing a prominent effect, marked in C with a star), and (B) low alpha power contralateral to the deviant location at right-hemispheric centro-temporal gradiometer (average across three channels showing a prominent effect, marked in C with a star). The black box marks the time-frequency range (time: 0.2 - 0.6 s, frequency: 8 - 14 Hz) used for the statistical cluster analysis. (C) Topoplot of the alpha power distribution in the 0.2 - 0.6 s time window following deviant onset. Stars indicate channels on which there was a prominent statistical effect. (D) Source estimation of the alpha power effects.

To test whether the observed alpha lateralization (see Figure 3) was driven by a decrease or increase in alpha-band power, we analyzed the time courses of alpha power for the right hemispheric centro-temporal ROI and the left hemispheric parieto-occipital ROI separately for the left and right deviant. We contrasted the alpha power following the deviant with the mean alpha power of the baseline period, via a non-parametric cluster-based permutation statistic. Since we had no a priori hypotheses on the direction of the alpha power modulations, we applied paired-sample, two-tailed Student’s t-tests. The results show that the right deviant caused a significant increase in alpha power in the right-hemispheric centro-temporal ROI, in the time range 0.2 to 0.4 s after sound onset (p = .017, see Figure 4 A). Furthermore, for the left deviant, we found a significant increase in alpha power in the left-hemispheric parieto-occipital ROI, in the time range 0.2 to 0.3 s after sound onset (p = .039, see Figure 4 B). Additionally, the right deviant caused a significant decrease in alpha power in the left-hemispheric parieto-occipital ROI, in the time range 0.5 to 0.6 s after sound onset (p = .044, see Figure 4 B). The left deviant neither caused a significant increase nor decrease in alpha power in the left-hemispheric posterior ROI.
Figure 4: Time courses of log10-transformed and group-averaged alpha power for (A) the right centro-temporal ROI and (B) the left parieto-occipital ROI, time-locked to the left deviant (red line) or the right deviant (blue line). Channels belonging to the ROI are marked with stars in the corresponding channel layout. Significant time samples are illustrated by the horizontal red and blue lines (*p < .05). Upward directed arrows indicate an alpha power increase, and the downward directed arrow indicates an alpha power decrease. Gray areas indicate the baseline interval (-0.4 to -0.2 s) and
the test interval (0.2 to 0.6 s). The orange and green bars indicate the presentation time of deviant and target, respectively.

4. Discussion

Our behavioral reaction time and magnetoencephalographic alpha power data support the notion that task-distracting deviant sounds cause a shift in spatial attention that contributes to behavioral distraction.

4.1 The shift of involuntary spatial attention contributes to behavioral distraction

As expected, our behavioral distraction effect - that is, the longer RTs to targets that followed a deviant versus a standard - shows that the deviant captured attention involuntarily. This is in line with numerous findings that were obtained via a deviant distraction task (Bendixen et al., 2007; Parmentier et al., 2008; Schröger & Wolff, 1998; Wetzel et al., 2012).

Importantly, we observed an advantage in response speed when targets followed deviants on the same versus opposite side. This clearly speaks in favor of a spatial shift of attention, which has been proposed, though not directly tested, earlier (Parmentier et al., 2008). That is, when the deviant and target occur on the same side, the time penalty due to the spatial shift is lower compared to when both events occur on opposite sides. This finding is in line with behavioral data obtained via the stimulus-driven spatial cueing paradigm (McDonald et al., 2000), and receives further support from our MEG data, as will be discussed below.

Note that the spatial component of the attention shift unlikely accounts for the overall behavioral distraction effect in full; in fact, several cognitive determinants may contribute to it. A second determinant that could be at play in the current crossmodal situation, is the involuntary shift of attention across sensory modalities (i.e., from vision to audition). In this case, the time penalty accumulated by a shift of attention from the visual to the auditory modality (i.e., upon the onset of the
deviant), and its re-orientation towards the visual modality (i.e., upon the onset of the target), would contribute to behavioral distraction (Parmentier, 2014). This idea receives support from behavioral studies showing that RTs are slower for a target in an unexpected modality compared to a target in an expected modality (Boulter, 1977; Spence et al., 2001).

At first glance, the RT data seem to be at odds with the hit rates. While the RTs indicate a decrease in performance for targets following a deviant versus a standard, the hit rates rather reflect an increase. However, this apparent contradiction can be reconciled by linking the RTs to the costs of the attention shift towards the deviant (Schröger & Wolff, 1998; Parmentier et al., 2008; for a review, see Parmentier 2014), and by linking the hit rates to an increase in nonspecific arousal that improves cognitive functioning (Näätänen, 1992). The current findings are well in line with previous research on deviant distraction, showing both costs and benefits (SanMiguel et al., 2010; Wetzel et al., 2012, Wetzel et al, 2019).

4.2 Posterior alpha power modulation reflects an involuntary shift of spatial attention

Our study is the first to show that a lateralized deviant sound modulates posterior alpha power in a spatially selective way. As expected, we observed a lateralized alpha power modulation over parieto-occipital sensors in the time window following the deviant sound and preceding the visual target. In line with our hypothesis, posterior alpha power modulation was higher in the hemisphere ipsilateral versus contralateral to the deviant location. That pattern of results was driven by an alpha power increase (relative to baseline) in the left hemisphere due to the left deviant (Figure 4 B) and likely reflects sound-induced spatial attention bias (involuntary spatial attention: Feng et al., 2017; Störmer et al., 2016; voluntary spatial attention: Deng et al., 2019; Deng et al., 2020; Frey et al., 2014; Wöstmann et al., 2016). This alpha power effect occurred relatively early and was rather short-lived, thereby closely matching the characteristics of involuntary attention both in timing and in the build-up phase (Corbetta & Shulman, 2002). The whole brain analyses further confirm the parieto-occipital localization of the effect (Figure 3 D).
From a functional point of view, our posterior alpha power increase is also an interesting finding. A lot of what we know about the functional role of oscillatory alpha activity in the context of spatial attention originates from voluntary attention research. Via endogenous spatial cueing tasks, attention was voluntarily guided to the location of an upcoming target (visual target: Kelly et al., 2006; Rihs et al., 2009; Worden et al., 2000; auditory target: Banerjee et al., 2011; ElShafei et al., 2018; Frey et al., 2014; Müller & Weisz, 2012). In those tasks, lateralized alpha power was larger over the ipsilateral than the contralateral hemisphere, relative to the attended target location. This alpha power modulation was driven by 1) an alpha power decrease in the contralateral hemisphere and/or an alpha power increase ipsilateral to the attended side (e.g., Haegens et al., 2011); 2) a bilateral alpha power decrease that was maximal over the hemisphere contralateral to the attended position (e.g., Thut et al., 2006); or 3) a bilateral alpha power increase that was maximal over the hemisphere ipsilateral to the attended side (e.g., Banerjee et al., 2011; Worden et al., 2000). The pattern that is observed, reflecting mechanisms of suppression or enhancement at play in attention biasing, probably relies on divergent task demands of the respective study (Kelly et al., 2006; Thut et al., 2006). In any case, based on those studies, it has been widely acknowledged that decreased alpha power reflects the processing of task-relevant information in the voluntarily attended space, and increased alpha power indicates the inhibition of processes associated with task irrelevant information in the unattended space (for a review, see Klimesch, 2012). In a similar vein, our left-hemispheric parieto-occipital alpha power increase due to the left deviant may reflect the suppression of irrelevant information in the unattended space, that is, where involuntary spatial attention was not captured. This active inhibition of visual brain areas may free up resources for brain areas contralateral to the deviant site that process relevant information (Jensen & Mazaheri, 2010; Meeuwissen et al., 2011), which are also consequently involved in attention reorienting towards the deviant location. This interpretation receives support from our behavioral data, suggesting that deviants capture spatial attention (Figure 2B). Furthermore, it is in line with findings suggesting that a sound can engage visual processing at the same location (Feng et al., 2014; Feng et al., 2017; McDonald et al., 2013; Störmer et al., 2016). Interestingly, earlier work focussing on involuntary spatial attention exclusively found posterior alpha power decreases (Störmer et al., 2016; Feng et al., 2017). In more detail, the attention-capturing sound
induced a bilateral alpha power decrease that was more prominent in the hemisphere contralateral to
the location of the sound. Based on these earlier data, the authors suggested that a shift in involuntary
attention exclusively facilitates target processing on the cued side, whereas target processing on the
opposite side is not negatively impacted (Feng et al., 2017; Störmer et al., 2016). This conclusion was
substantiated by the characteristic pattern of alpha power modulation preceding correct and incorrect
discriminations of valid and invalid targets (Feng et al., 2017). Our results - demonstrating an increase
of parieto-occipital alpha power - challenge that conclusion, as well as the prevalent notion that the
“lack of an early inhibitory influence may represent a fundamental difference between involuntary
orienting and voluntarily directed attention” (Feng et al., 2017, page 326). The difference between the
current and previous studies is that our’s did not only apply an exogenous spatial cueing task, but also
a distraction task, thus, likely resulting in different task demands (Kelly et al., 2006; Thut et al., 2006).

Considering the crossmodal nature of our task involving attention-capturing deviant sounds and visual
targets, one might argue that our posterior alpha power increase was rather linked to the attentional
disengagement from vision. Indeed, the outcome from intermodal attention tasks supports this view
(Fox et al., 1998; Fu et al., 2001). For example, it is known that directing attention to auditory (versus
visual) features of upcoming audio-visual targets increased posterior alpha power in the pre-target
interval (Fox et al., 1998). That finding was interpreted to reflect “a disengaged visual attentional
system in preparation for anticipated auditory input that is attentionally more relevant.” We cannot
rule out that alternative account. However, we consider the spatial shift of attention explanation more
likely. This receives support from our reaction time data from the congruent and incongruent
conditions. Further support for the spatial shift of attention interpretation comes from another yet
unpublished study (Weise et al., in preparation). In that purely auditory task we also found lateralized
posterior alpha power modulation that was driven by a deviant-induced alpha power increase.
Crucially, the main difference in the new task compared to the one used here was that it did not
contain visual but auditory targets. Thus, in that case the data cannot be linked to visual
disengagement.
4.3 Centro-temporal alpha power increase reflects disengagement of motor and auditory areas

In addition to the current evidence pointing to a spatial shift of attention (see above), we found a right-hemispheric alpha power modulation over centro-temporal sites (Figure 3). This right-hemispheric effect was driven by an increase in alpha power, relative to baseline, due to the right deviant (Figure 4 A). Supported by the source data, this may reflect the disengagement of motor areas (Figure 3 D).

Indeed, the disengagement of motor areas can be caused by unexpected events such as deviant sounds (Dutra et al, 2018; Vasilev et al., 2019; Wessel et al., 2016; for a review, see Wessel & Aron, 2017). According to that research line, deviants recruit a ‘stopping network’ including the prefrontal cortex/pre-supplementary motor area and temporarily interrupt the current action. The same network is also active following other unexpected events like unexpected action outcomes or action errors and is linked to motor slowing such as slowing of RTs and the impairment of ongoing cognitive processing. Its putative purpose is to put the task at hand on hold to free up processing resources and allow an attention shift towards the unexpected event. Our behavioral and MEG data nicely fit that view and might even extend it concerning the neuronal mechanisms engaged. Previous research linked an power increase in the delta (2–4 Hz) and low-theta (5–8 Hz) frequency bands to successful stopping (Huster et al., 2013; for a review, see Wessel & Aron, 2017), while a modulation in posterior alpha activity has been linked to attentional processing (Wessel & Aron, 2014). Our findings extend that view and suggest that central alpha power may also implement motor suppression. This receives support from research showing an alpha power increase during voluntary suppression of actions (Hummel et al., 2002; Sauseng et al., 2013).

Also, the current data add to the growing number of studies suggesting that distraction is more than a byproduct of attentional orienting. It includes transient motor suppression (Dutra et al., 2018; Vasilev et al., 2019). At that point, one might wonder whether the transient motor suppression itself might be responsible for our behavioral spatial attention effect. Yet this is unlikely. Even though the deviant-induced motor interruption might in general contribute to longer reaction times (compared to
standards), this should, however, not obscure our RT data supporting a spatial shift of attention. The motor interruption is induced by deviants irrespective of their location and becomes evident after about 150 ms (Wessel & Aron, 2013). Importantly, the effect of the spatial shift of attention exclusively depends on whether the target follows the deviant at the same or different side. Another question that might raise concerns is whether the observed central alpha power modulation can be explained by preparatory processes related to the task. We consider that far-fetched because: 1) Participants responded to the left and right target with both hands, and the button-response assignment was counterbalanced across participants; 2) The alpha power modulation occurred in the pre-target interval, thus before the participant knew which hand was required for the response.

We want to emphasize that our interpretation of the central alpha power modulation is mainly based on post-hoc speculation. Our study was not designed to test the hypothesis that central alpha power implements deviant-induced motor-stopping. Further research is needed to yield more insight into that aspect.

4.4 Conclusion

Here, we show that lateralized deviant sounds cause a shift of involuntary spatial attention and contribute to behavioral distraction. This is evident in our reaction time data, which demonstrates that responses are faster when the target follows an auditory deviant on the same side compared to the opposite sides. Additionally, our MEG data clearly show that oscillatory alpha power is shaped in a spatially selective manner. Most importantly, attention-capturing, task-distracting deviant sounds induced posterior alpha power that was higher in the left vs. right hemisphere. These attention-related changes were driven by alpha power increases due to the left deviant, suggesting disengagement of visual areas that process task-irrelevant information outside the locus of involuntary attention. Together with the behavioral data, our oscillatory data strengthen the view that alpha power reflects a shift of involuntary spatial attention.
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SOUNDS CAPTURE INVOLUNTARY SPATIAL ATTENTION


Author Note

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Data and code availability

The preprocessed MEG and behavioral data used to generate the figures and calculate the statistics are available at the OSF public repository (https://osf.io/qt95u/). The corresponding data analysis code is available at the corresponding author’s GitLab repository (https://gitlab.com/aweise/crossfrog-final-analysis). The original raw data are available, upon reasonable request, from the corresponding author.
Supplementary materials

**Figure S1:** Log10-transformed, non-phase locked alpha power modulation to lateralized deviants based on group-averaged data for Deviant Left minus Deviant Right. (A and B) Time-frequency representations highlighting (A) high alpha power ipsilateral to the deviant location at
left-hemispheric posterior gradiometers (average across three channels showing a prominent effect, marked in C with a star), and (B) low alpha power contralateral to the deviant location at right-hemispheric centro-temporal gradiometer (average across three channels showing a prominent effect, marked in C with a star). The black box marks the time-frequency range (time: 0.2 - 0.6 s, frequency: 8 - 14 Hz) used for the statistical cluster analysis. (C) Topoplot of the alpha power distribution in the 0.2 - 0.6 s time window following deviant onset. Stars indicate channels on which there was a prominent statistical effect. (D) Source estimation of the alpha power effects.