Behavioral/Cognitive

Attentional Modulation of Alpha/Beta and Gamma Oscillations Reflect Functionally Distinct Processes

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The brain adapts to dynamic environments by adjusting the attentional gain or precision afforded to salient and predictable sensory input. Previous research suggests that this involves the regulation of cortical excitability (reflected in prestimulus alpha oscillations) before stimulus onset that modulates subsequent stimulus processing (reflected in stimulus-bound gamma oscillations). We present two spatial attention experiments in humans, where we first replicate the classic finding of prestimulus attentional alpha modulation and poststimulus gamma modulation. In the second experiment, the task-relevant target was a stimulus change that occurred after stimulus onset. This enabled us to show that attentional alpha modulation reflects the predictability (precision) of an upcoming sensory target, rather than an attenuation of alpha activity induced by neuronal excitation related to stimulus onset. In particular, we show that the strength of attentional alpha modulations increases with the predictability of the anticipated sensory target, regardless of current afferent drive. By contrast, we show that the poststimulus attentional gamma enhancement is stimulus-bound and decreases when the subsequent target becomes more predictable. Hence, this pattern suggests that the strength of gamma oscillations is not merely a function of cortical excitability, but also depends on the relative mismatch of predictions and sensory evidence. Together, these findings support recent theoretical proposals for distinct roles of alpha/beta and gamma oscillations in hierarchical perceptual inference and predictive coding.

Key words: alpha; attention; gamma; magnetoencephalography; oscillations; predictive coding

Introduction

A long tradition of research in cognitive neuroscience has addressed the question of the extent to which our brain processes the sensory environment selectively and the mechanisms by which this selection occurs (Luck et al., 2000; Carrasco, 2011). Neurophysiological correlates of attentional modulation in prestimulus alpha/beta and poststimulus gamma oscillations have supported more traditional findings of early and late attentional selection (for review, see Luck et al., 2000). More recent theoretical proposals have considered the nature of attentional selection from the viewpoint of predictive coding and formulate attention in terms of Bayesian inference with the idea that expectations about the salience or precision of sensory information shape perception (Friston, 2005; Feldman and Friston, 2010).

A distinct alpha modulation in the prestimulus period (Worden et al., 2000; Fu et al., 2001; Bauer et al., 2012b; Mazaheri et al., 2014) and gamma modulation in the poststimulus period (Fries et al., 2001; Bauer et al., 2006; Siegel et al., 2008) is seen in several lines of experimental work. This temporal dissociation begs the question as to whether these frequency-specific effects are functionally distinct or whether they reflect the same top-down modulation that is merely expressed differentially, depending on stimulus-driven neuronal activation. It is well known that low-frequency alpha (and usually beta) oscillations are suppressed by stimulation (Pfurtscheller and Lopes da Silva, 1999; Bauer et al., 2006), whereas gamma oscillations are enhanced (Hoogenboom et al., 2006; Donner and Siegel, 2011). In other words, high- and low-frequency oscillations show different responses to afferent input, and this might be a sufficient explanation for the temporal dissociation of their attentional modulation. An alternative explanation calls on distinct top-down signals where one is inhibitory (Klimesch et al., 2007; Jensen and Mazaheri, 2010) and the other is facilitative (Womelsdorf et al., 2007). Recently, on the basis of laminar-specific spike frequency coherence in high and low frequencies in primates (Maier et al., 2010; Buffalo et al., 2011), Bastos et al. (2012) suggested that low-frequency oscillations provide predictions via feedback connections, while gamma oscillations signal the mismatch of sensory input to these predictions (the prediction error) via feedforward connections to higher-level brain regions.

Here, we investigate this proposal in two visual spatial attention experiments. The first experiment follows a “classical design,” where participants had to discriminate the orientation of a static grating, a feature available from stimulus onset. In experiment 2, the task was to detect a speed change (at a cued location) that occurred several hundred milliseconds after stimulus onset. Hence, in experiment 2, in contrast to experiment 1, the timing of visual stimulation and onset of the attended feature were disso-
cated. If the temporal profiles of alpha/beta and gamma modulations can be explained by the stimulus-related suppression of alpha modulations and induction of gamma modulations, one would expect to see the same temporal pattern of alpha and gamma modulation in both experiments. However, if alpha modulation reflected a specific expectation signal before the onset of the task-relevant stimulus feature, one would expect the alpha modulation to persist beyond stimulus onset in experiment 2.

Materials and Methods

Participants. Eight subjects (3 males; mean age, 27.25 years; SD, 4.5 years) participated in experiment 1, and 10 subjects participated in experiment 2 (5 males; mean age, 24.2 years; SD, 4.8 years). Subjects were recruited from a local database and provided informed consent for the study protocol, which was endorsed by the local ethics committee.

Task. In both experiments (Fig. 1), stimuli were presented via a digital LCD projector on a screen (height, 32 cm; width, 42 cm; distance from participant, ~70 cm) inside a magnetically shielded room using Presentation software (Neurobehavioral Systems). In both experiments, a central visual cue (arrow pointing left or right, pseudo-randomized with equal frequency of occurrence, subtending 1° of visual angle) was presented for 500 ms, instructing participants to subsequently attend to the left or right of fixation. This cue was deterministic and informed participants of the behavioral relevance of the upcoming two stimuli (bilateral gratings, either left or right relevant), rather than predicting with a certain probability the location of a stimulus. This allowed us to compare brain activity in the two hemispheres as a mere function of spatial attention, while keeping the physical stimulus identical across experimental conditions. After a variable baseline interval (1300–1700 ms in experiment 1 and 1000–1500 ms in experiment 2), the bilateral stimuli were presented in the left and right hemifields.

In experiment 1 (Fig. 1A), two Gabor-like patches (sinusoidal stimuli with a carrier frequency of 0.5 cycles/° multiplied with a Tukey window size of 9°) were presented with a slight tilt from the vertical axis. Participants were asked to indicate the direction of the tilt with respect to the vertical axis (left index finger counter-clockwise tilt–right index finger clockwise tilt). Participants were trained on the task, and the tilt angle was adjusted using a block-wise staircase procedure (to minimize influences of lapses) that increased the tilt angle if the participant committed >20% errors and decreased if they committed <10% errors. This procedure familiarized the subjects with the task requirements and made the task sufficiently engaging (with performance levels at ~80–90% accuracy). During the experiment, the gratings were presented at eccentricities of either 6° or 8° of visual angle. This factor was not of relevance to the question at hand (since MEG does not measure responses at the receptive field level), so we pooled trials across both eccentricities.

In experiment 2 (Fig. 1B), stimuli comprised concentric circles adapted from Schoffelen et al. (2005) and Hoogenboom et al. (2006) with a contracting motion (standard speed, ~0.5 cycles/s). Stimuli were always presented bilaterally at a (symmetric) eccentricity of 8° of visual angle at two different contrast levels (100% or 66%, the same for both stimuli). Again, we averaged across this factor. Stimuli were modulated using a Tukey window of size 9°. For the full set of stimuli, a cycle was divided into 250 phases (to enable smooth motion at various speeds); that is, for the progression of one circle to the nearest inward position (spatial cycle), there were 250 frames available in total (of which only a subset was presented on successive frames). After a stimulus onset asynchrony (SOA) of 382 ms, the speed of this motion became faster or slower at a time point selected randomly (in discrete steps of ~16.6 ms) until 1362 ms after stimulus onset, independently on either side. The speed change could therefore occur on either side at random timing, but only the cued side was task relevant. In 20% of trials, no speed change occurred on either side. Participants were asked to indicate the direction of speed change (faster or slower) by means of a button press with the left (slower) or right (faster) index finger. Participants were trained on this task with a staircase procedure, increasing the level of the speed change if their performance fell below 80%, and decreasing, if it rose above 90%.

MEG recording and analysis. Data were recorded continuously from 274 axial gradiometers using a CTF Omega whole-head system at a sampling rate of 600 Hz in third-order gradient configuration. MEG data analysis was implemented in SPM8 (Litvak et al., 2011) and Fieldtrip (Oostenveld et al., 2011). Data preprocessing comprised the following steps: data were converted to SPM8 format and notch filtered from 49 to 51 Hz (on the entire unepoched dataset). Eye blinks were detected using a semiautomatic thresholding procedure, a z-transform of the absolute value of the 5–15 Hz bandpass-filtered Hilbert transform, with interactive threshold setting (this was done blind to the condition and for all trials of the session simultaneously). These data were then inspected for excessive artifacts, and a principal component analysis was computed over the eye-blink periods. In all participants, the first spatial component reflected the eye blink topography (an obvious “bipolar pattern” over frontal and temporal sensors). The data were then epoched: for experi-
ment 1 (short stimulus presentation) a window of −2000 to 1200 ms around stimulus onset was used, whereas for experiment 2, the epoch ranged from −2000 to 2500 ms after stimulus onset. Eye-blink artifacts were then suppressed by removing the first principal component above using signal space projection. Finally, epochs were visually inspected (using ft_reject_visual.m), and trials with excessively high power (summed over all channels) were removed. For source analysis, individual electromagnetic forward models were constructed based either on an individual structural T1-weighted MRI (when available) or on the default MNI brain coordinates (when not). For those participants with an individual structural MRI (5 of 8 participants in experiment 1 and 8 of 10 participants in experiment 2), the T1 MNI brain was first warped on their individual structural image, and this transform was then applied to the (canonical) cortical surface mesh, as described in the study by Mattout et al. (2007) (see also Henson et al., 2009). For beamformer solutions, covariance matrices were calculated for the low and high frequencies. For low frequencies, the data were low-pass filtered (35 Hz cutoff frequency), and the covariance matrix was calculated over the entire epoch. The same approach was used for high frequencies, where the data were bandpass filtered at 0.15–150 Hz data (were low pass filtered during recording with a 130 Hz antialiasing filter). Beamformer filters were then calculated for 413 cortical vertices on a (downsampled) cortical mesh (Van Veen et al., 1997). Time series data were then projected through the resulting beamformer coefficients to produce time courses for subsequent time frequency analysis. Separate reconstruction of high-frequency responses reflects the fact that low-frequency responses require a narrower bandwidth, whereas higher frequencies have a larger bandwidth. To account for this, we used a multitaper approach for high-frequency responses (Mitra and Pesaran, 1999) that allows smoothing in both time and frequency. For low frequencies, time windows of 400 ms (from −1.95 to 0.8 s in experiment 1, and from −1.95 to 2.5 s in experiment 2, both in steps of 0.1 s) were multiplied with a Hanning window and then multiplied with complex exponentials at frequencies from 2.5 to 30 Hz in steps of 2.5 Hz. For the high frequencies, time windows of 400 ms (from −1.95 to 0.8 in experiment 1 and to 1.5 in experiment 2, both in steps of 0.05 s) were used with seven discrete prolate Slepian tapers multiplied with complex exponentials at frequencies from 40 to 150 Hz in steps of 5 Hz (with an effective smoothing of 10 Hz in both directions). The resulting cortical power spectra for each trial were then log transformed, and outliers (above and below 2 SDs from the mean) were removed for each time–frequency bin. Finally, condition-specific responses were summarized with statistics (mean and SEM over trials within the condition) for “attend left” and “attend right” separately. The resulting condition–specific power spectra were subtracted from baseline values (from −1.95 to −1.85 s) and smoothed using a moving average over three neighboring time–frequency bins to accommodate well known interindividual differences in induced responses (Muthukumaraswamy et al., 2009; van Pelt et al., 2012).

Statistical analysis. To quantify attentional effects, we first calculated the attentional lateralization (AL) index for condition-specific power estimates as follows:

\[ AL(r, f, t) = \frac{1}{2} \left( S(r, f, t)_{\text{attend}} - S(r, f, t)_{\text{attend}} \right) \left( |\text{abs}(S(r, f, t)_{\text{attend}})| + |\text{abs}(S(r, f, t)_{\text{attend}})| \right), \]

where \( S(r, f, t) \) is the power spectral density at a cortical vertex \( r \) for frequency \( f \) at time \( t \) (absolute values for denominators to accommodate for rare cases where the sum of baseline-corrected values in individual bins for AttendLeft versus AttendRight gave near-zero values for denominator).

We determined the nearest homologous vertex in the left hemisphere for each vertex in the right hemisphere. We then subtracted the right AL values from their corresponding left hemisphere values, effectively summarizing mirror-symmetrical attentional modulation (Bauer et al., 2012). The resulting data were subjected to a cluster-based permutation test (Maris and Oostenveld, 2007) using a one-sample t test (of attentional lateralization against zero) to find significant attentional effects. This procedure corrects for multiple (nonindependent) comparisons over cortical vertices, time, and frequency. We performed cluster-based inference separately for the low and high frequencies, and focused on the prestimulus period before the target and the poststimulus period. We therefore searched over 5–25 Hz (for low-frequency modulation effects) and 40–150 Hz (for high-frequency effects). The time window considered was −500 to 800 ms after stimulus onset (for experiment 1) and −500 to 1500 ms (for experiment 2), because our hypotheses were about the neuronal dynamics around stimulus onset and during subsequent processing.

Dependency of behavioral and physiological responses on hazard rate. To investigate the effect of expectations in experiment 2 on behavioral and electrophysiological responses, we collapsed trials over nine time bins according to the SOA between stimulus onset and the target (speed change). The cumulative probability of a speed change occurring within a certain bin (the hazard rate) as well as negative and inverse reaction time (as two different measures of behavioral response speed) were averaged, and the SD of reaction time (RT) was calculated for each bin. This revealed a large negative correlation between reaction time and the cumulative probability (both averaged in bins of trials) of \( r = -0.98 \) across participants, as well as a negative correlation of cumulative probability with the SD of binned reaction times (\( r = -0.71 \)), indicating faster and less variable responses when the stimulus change was expected. Inverse reaction time and hazard rate were also highly correlated (\( r = 0.9 \)). Since cumulative probability and negative response time showed a rather strong linear relationship (see Fig. 3), we chose the cumulative probability as the relevant metric of stimulus expectation (in contrast to the hazard rate; Schoffelen et al., 2005).

To characterize the time courses of attentional lateralization in terms of stimulus expectation, a weighted mean of attentional lateralization in each participant was calculated from all significant vertices (grid points in the cortical mesh). The weights assigned to each vertex were the proportion of significant time–frequency bins over all time–frequency bins. To average the time courses of attentional lateralization over frequencies, we calculated again a weighted average where the time courses obtained for each frequency were weighted by the proportion of (multiple comparison corrected) significant bins for each frequency. This measure precludes biased selection of frequencies or vertices and identifies responses that survive correction for multiple comparisons, weighted by the proportion of significant effects in time and frequency bins. These individual time courses were then used for a mixed-effects ANOVA to test for differential peristimulus attentional lateralization effects in the two experiments (using prestimulus vs poststimulus times as a repeated-measures factor and experiment 1 vs 2 as a between-subject factor).

To investigate more precisely how the attentional lateralization in low and high frequencies depends on target (speed change) expectation, it was necessary to separate the trials with early and late speed changes. This is because trials with early targets would otherwise contribute to the estimation of responses induced by late targets. Therefore, we time locked the spectral estimates in each trial to the target (speed change). The cumulative probability of a speed change occurring within a certain bin (the hazard rate) as well as negative and inverse reaction time (for experiment 2), because our hypotheses were about the neuronal dynamics around stimulus onset and target onset. These categories reflect different target timings and therefore specific values of the underlying hazard function that determines the strength (or precision) of expectations. We then regressed the attentional lateralization index (entire time frequency matrix from −500 to 500 ms around target onset) on the timing category and report the associated t values (thresholded at \( p < 0.05 \), two-tailed; see Fig. 4).

Correlation of gamma to reaction time. Finally, we conducted a control analysis to ensure that we could reproduce previously reported correlations of gamma modulation with reaction time. To this end, we split the trials of each subject into seven reaction time bins (for the same approach, see Bauer et al., 2009) and then regressed the binned response on reaction time, reporting the associated \( t \) values. This analysis was performed over an average of vertices within 3 cm of the right calcarine sulcus. The results confirmed that stronger induced gamma is associated with faster responses (see Fig. 5).

Results

Participants in experiment 1 gave correct responses in a mean of 88.9% of the trials (SD, 9.3%) with an average response latency of 659 ms (SD, 126 ms). In experiment 2, participants responded with an average latency of 827 ms (SD, 119 ms) with a mean accuracy of 80.5% (SD, 12.7%). We had no hypotheses concerning
the effect of the experimental conditions (or comparisons between experiments) regarding the behavioral data. We discuss the correlations of reaction times with the predictability of targets in greater detail below.

**Attentional modulation of low and high frequencies**

Figure 2 shows the stimulus-induced responses and the effects of spatial attention in both experiments at the cortical and source level, in Figure 2A–F for the alpha/beta band and in Figure 2G–L for the gamma band. Figure 2, A and D, shows the stimulus-induced responses (averaged z values for the comparison between peri-stimulus time and precue baseline) in occipital cortex for experiment 1 (gratings) and experiment 2 (moving circles), respectively. Note the suppression of low-frequency activity at stimulus onset in both experiments, as would be predicted from many previous studies (Bauer et al., 2006; Hoogenboom et al., 2006). Figure 2B shows the comparison of alpha lateralization for “attend-left” – “attend-right” trials in experiment 1. This shows the relative suppression of alpha/beta power in the hemisphere contralateral to the attended hemifield (ipsilateral enhancement) and originates predominantly from the parieto-occipital sulcus, which is in agreement with previous findings (Fig. 2G; Bauer et al., 2012b). Note that this effect occurs predominantly in the period preceding the stimulus, consistent with numerous previous studies (Worden et al., 2000; Fu et al., 2001; see Discussion). Figure 2E shows the attentional lateralization in experiment 2. Note that, here, attentional suppression of alpha power appears in the poststimulus period. It is also present before stimulus onset, but is less pronounced than in the poststimulus period, particularly toward the end of the trial. Figure 2F shows the thresholded topography of this effect, indicating that it is more distributed; involving occipitotemporal areas, the temporoparietal junction (TPJ) and premotor areas (e.g., the frontal eye field). Crucially, in experiment 2, the peak attentional alpha/beta modulation was seen in the poststimulus period, after stimulus onset, whereas in experiment 1 it was found predominantly in the prestimulus period (Fig. 3, time courses of both effects).

Figure 2, G and J, shows the stimulus-induced gamma oscillations for both experiments. In experiment 1 (Fig. 2G), we see an increase in gamma band oscillations ~500 ms, which then return to baseline. The later lower-frequency increase is
a result of the so-called beta rebound (Jensen et al., 2005; Bauer et al., 2006; Fig. 2A) that extends into higher frequencies, partly due to the spectral smoothing characteristics of the multitaper frequency analysis. In Figure 2I, we see the same for experiment 2 with the sustained motion stimulus, initially high frequency-induced gamma oscillations that then turn into a sustained gamma band response at a slightly lower frequency. Figure 2H shows the attentional enhancement of gamma-band power in contralateral visual cortex for experiment 1. The peak of this effect is seen clearly in the poststimulus period, although some sparse modulations at (and above) 100 Hz appear at stimulus onset. Figure 2I shows that the topography of this effect is dominated by extrastriate occipital cortex. Figure 2K shows the thresholded spectrum of the attentional modulation for experiment 2—with a similar pattern to that shown in Figure 2H, although somewhat more sustained, given the nature of the task. Figure 2L shows its topography, which again extends slightly more toward temporal areas, presumably due to the nature of the speed change detection task.

Figure 3 shows the time courses of these attentional modulations. These time courses reflect weighted averages of the attentional lateralization index (i.e., the normalized power difference between hemispheres corresponding to the attended vs the unattended hemifield). These results replicate the basic finding in Figure 2: in experiment 1, the peak of the lateralized alpha-beta modulation (Fig. 3A) is in the prestimulus period, then slowly returns to baseline (and might rebound toward the end of the trial in opposite direction). Conversely, in experiment 2, while the attentional lateralization is already observable in the pre-stimulus period, the peak emerges toward the end of the trial, in the poststimulus period. Our main aim was to investigate whether, in a paradigm in which the task-relevant stimulus (change) occurs after stimulus onset (experiment 2), alpha/beta lateralization was observable in the later poststimulus period. Figures 2 and 3 illustrate that this is indeed the case. We nevertheless tested explicitly for a significant time shift of the dominant period of alpha lateralization in experiment 2 compared with experiment 1 (the latter reflecting the temporal pattern found in many previous studies). A 2 × 2 ANOVA was performed on the attentional lateralization with repeated-measures factor “time interval” (prestimulus vs poststimulus; Figs. 2, 3, time intervals) and between-subjects factor “experiment” (dynamic vs static gratings). The interaction was significant ($F_{(1,1,16)} = 11.17, p < 0.01$), confirming the dissociation induced by the two experimental paradigms with respect to timing.

Relation of stimulus change probability, behavior, and physiology (experiment 2)

We separated trials into nine equally spaced bins (from 433 to 1433 ms) according to the SOA of the stimulus change (target) to
the display onset and examined the correlation between the average RT and the hazard rate for these bins. In accord with previous studies, participants appear to have tracked this probabilistic structure rather well, as reflected in a correlation of $r = 0.9$ ($p < 0.01$). The hazard rate (a conditional probability) increases relatively sharply toward the end of the trial. In contrast, the cumulative probability indicates the probability—across all trials—that a stimulus change will have occurred at any given point in post-stimulus time. Due to the uniform probability distribution, this is a linear function. To make these measures easier to compare, Figure 3, B and D, shows the normalized (and sign-inverted) cumulative probability of stimulus change as a function of SOA (aligned with the time axis for the attentional lateralization) and the average reaction times for the respective time bins (individual bins are connected by line segments). These results illustrate the tight relationship between these measures ($r = -0.98$ for averaged bins and RT, $p < 0.01$), indicating that participants learned this “probabilistic structure” and form precise expectations about the occurrence of the target (change in speed). The precision of stimulus expectancy should also be reflected in the SD of reaction times. Indeed, there was also a negative correlation between cumulative probability and the SD of reaction times ($r = -0.71$, $p < 0.01$), confirming that strong expectations reduce not only the average response latency, but also their variance.

Figure 3B shows another salient feature; namely, the fact that the strength of attentional alpha lateralization seems to follow the cumulative probability (or rather temporally precede it by a few hundred milliseconds). The correlation of alpha lateralization with cumulative probability was moderate, although significant across time bins and participants ($r = -0.25$, $p < 0.05$). For the gamma lateralization, the pattern is reversed (note that “maximal absolute alpha lateralization” is inverted, placing the largest magnitude of contralateral suppression at the bottom); gamma lateralization is maximal just after the earliest possible time the target can appear. With an increasing cumulative probability for target appearance, gamma lateralization is clearly reduced in amplitude ($r = -0.43$, $p < 0.05$), hence showing the opposite pattern with respect to the absolute strength of attentional modulation.

There is, however, one shortcoming with the above analysis: the average calculated across all trials (with the spectra giving estimates of lateralized power) includes attentional lateralization after the stimulus change has already occurred for the later time bins. To avoid this possible confound, we conducted an additional analysis in which we focused on the responses locked to the speed change. Spectral estimates of attentional lateralization were time locked to the speed change, and trials were then sorted into four categories of SOAs between stimulus and target onset. Figure 4A shows the $t$ values of the regression of alpha lateralization on SOA (significant time–frequency windows marked; $p < 0.05$ two-tailed, uncorrected). This takes the form of a stronger suppression of contralateral alpha lateralization when the stimulus change occurs late in the trial (i.e., is more likely to occur). Figure 4B shows the result of the same regression analysis, but now for the attentional gamma modulation [note that we inverted the sign of the predictor variable for this analysis (negative SOA), which makes a direct comparison of Figs. 4A (contralateral alpha decrease by attention), B (contralateral gamma enhancement by attention) more intuitive, given the inverted sign of these effects]. This regression of the gamma modulation on (negative) SOA shows the opposite of the SOA effect on alpha modulation in Figure 4A; there is increased gamma lateralization at short SOAs, clearly peaking after the stimulus change (~150 ms), and a negative relationship between SOA and gamma lateralization. In summary, the results of this analysis confirm the results of Figure 3 that more precise predictions increase the magnitude of the attentional alpha modulation and decrease gamma lateralization.

High-frequency gamma responses and reaction time

The above result implies a positive correlation between gamma lateralization and response time, when calculated over averaged binned trials sorted by target SOA. Since this result (implying stronger gamma for slow responses) is at odds with those in several reports in the literature (Womelsdorf et al., 2006; Hoogenboom et al., 2010), we characterized the relationship between induced gamma oscillations and RT when not averaging within bins sorted by SOA (which eliminates intrinsic variability in response times). The result of a regression analysis of stimulus-induced gamma oscillations in occipital cortex (ROI around the calcarine sulcus, response averaged in seven bins of trials sorted by reaction time) on reaction time per se clearly shows (Fig. 5) that our study replicates previous results by showing larger stimulus-induced gamma oscillations for faster responses.

Discussion

In summary, we found that the typical prestimulus nature of attentional alpha/beta lateralization is not simply explained by an attenuation caused by afferent drive following the onset of a sensory stimulus. In experiment 2, where the task-relevant stimulus feature was delayed with respect to stimulus onset, the alpha/beta modulation peaks during strong afferent stimulation by a high-contrast motion stimulus. Instead, our results suggest that attentional alpha/beta modulation reflects the precision of expectations about impending stimulus events (as indexed by the objective hazard function and behavior; Figs. 3B, 4A). Thus, greater alpha modulation is seen when subjects are more certain that the event is about to occur and the modulation returns to baseline when the actual evidence is presented. Conversely, attentional lateralization of gamma os-
oscillations is stimulus bound, and its strength depends on the mismatch between expectations and physically presented stimuli. In other words, the strength of lateralized attentional gamma modulation predominantly postdates task-relevant stimulus features and is greater when these are unpredicted (or more surprising). So far, these results are therefore consistent with the notion that prestimulus modulations of alpha activity set the context (sensory) for expected stimuli, while stimulus-bound gamma responses signal prediction errors induced by stimulus changes, particularly when the stimuli are less predicted.

The fact that alpha/beta frequency modulation reflects anticipatory predictions about stimulus events has been shown in the visual (Cravo et al., 2011, 2013; Rohenkohl et al., 2012), somatosensory (van Ede et al., 2011, 2014) and motor (Schoffelen et al., 2005) domains (see also Arnal and Giraud, 2012). These data, together with our results, are therefore consistent with suggestions that low-frequency oscillations mediate feedback or descending predictions that include predictions about the precision of specific processing channels or stimulus features that establish an attentional context for subsequent sensory processing (von Stein et al., 2000; Bauer, 2008; Bastos et al., 2012), a point we discuss in greater detail below. What is new with respect to alpha oscillations in our study is the finding that this lateralized attentional modulation can be seen under strong afferent stimulation (von Stein et al., 2000; Bauer et al., 2012a); and in a study in the placebo condition by Bauer et al. (2012b). In the study by Bauer (2008, chapter 3), we present a reanalysis of data from a previous study (Bauer et al., 2006), in which no stimulus-induced attentional alpha/beta lateralization was reported. This reanalysis confirmed that a significant, lateralized alpha/beta modulation in somatosensory cortex was found before stimulus onset and returned to zero during the poststimulus period, then rebounding before the onset of the next stimulus. Interestingly, there was no lateralization after the onset (and during the processing) of the tactile stimulus. This finding has very recently been confirmed, using very similar stimuli, by van Ede et al. (2014). Fries et al., 2008 showed (with invasive recordings from monkey V4 in a speed change detection task) that the alpha/beta attentional modulation was dominant in the prestimulus period (their Fig. 5) and much less pronounced in the poststimulus period (their Fig. 7). Hence, there is clear evidence that the peak of alpha/beta lateralization is typically found prestimulus, and that the peak gamma lateralization emerges in the poststimulus period.

With respect to the attentional modulation of gamma oscillations, our findings are more controversial. A previous study, using invasive recordings and intra-areal and interareal measures of synchronization in cats (von Stein et al., 2000), reports that low-frequency oscillations reflect the strength of a prediction signal and gamma oscillations reflect a prediction error. Hence, these results are similar to our results, which are based on intra-areal synchronization measured with MEG. A study by Lima et al. (2011) specifically investigated the effect of expectation on stimulus-induced gamma oscillations in two monkeys using tasks where the stimulus could occur at either of two discrete time points, cued or uncued, with different probabilities. In one monkey, Lima et al. (2011) found a significant reaction time effect, but both monkeys showed increased stimulus-induced gamma oscillations with higher expectations, to some extent contrary to our findings.

There are several differences between the study by Lima et al. (2011) and ours. First, we looked at the impact of a spatially selective top-down signal on occipital activity, indexed by lateralized responses, whereas they described a global stimulation effect. Second, in our study, human subjects performed a task with a continuous probability distribution. The fact that we observe a strong correlation between cumulative probability (expectation) and average binned response time of \( r = -0.98 \) across subjects (\( r = -0.71 \) for the SD of the response time) shows that human observers form and use their subjective precision or confidence in their predictions. Schoffelen et al. (2005) used a task similar to the one we used (in fact, our experiment 2 is a modification of their study) and showed that, at least in the motor system, corticospinal coherence in the gamma range showed a positive correlation with the hazard function, similar to the findings of Lima et al. (2011). We did not examine stimulus-induced responses for two reasons. First, our focus was on the lateralized attentional effects on alpha/beta and gamma oscillations. Second, characterizing stimulus-induced responses to a stimulus change is confounded by adaptation to the stimulus per se. Surprise-related effects on stimulus-induced gamma oscillations may be better examined using separate presentations of stimuli following a probability distribution, as opposed to predictable and unpredictable changes in an ongoing stimulus. The literature on the relation of gamma oscillations to stimulus expectations is currently based on different measures of neuronal activity; nevertheless, there seems to be an interesting heterogeneity in the relationship between gamma oscillations and the strength of predictions, which we discuss in a more principled way below.

In the context of predictive coding, it has recently been proposed that low-frequency oscillations provide prediction signals via descending (feedback) connections to a lower level areas,
whereas gamma oscillations (reflecting feedforward propagation of neuronal activity) convey the prediction error encoded by superficial pyramidal cells (Bastos et al., 2012), consistent with our own findings and those of von Stein et al., 2000. However, the relation among predictability, attention, and the expected strength of a feedforward signal on downstream neurons is more intricate. Under generalized predictive coding (Feldman and Friston, 2010; Brown and Friston, 2013), attention is mechanistically conceptualized as the top-down regulation of the precision (inverse variance) of sensory signals through synaptic gain modulation and is typically thought to follow the anticipation (or expectation) of behaviorally relevant stimulus events. It is assumed that while predictions of a stimulus attenuate its feedforward imprint by inhibitory or subtractive effects on prediction error units in superficial layers (that are thought to be the predominant source of MEG signals), the predictability of a behaviorally relevant stimulus can also augment synaptic gain in superficial layers that may reverse the suppressive effects of the prediction per se. This gain control corresponds to an encoding of precision or confidence in prediction errors that boosts their influence on processing in the next level of the cortical hierarchy (Feldman and Friston, 2010; Brown and Friston, 2013). The ensuing changes in the membrane potential from the synaptic gain control is supposed to change the temporal properties of the neurons and lead to a shift from low frequencies to higher frequencies (Chawla et al., 1999), hence boosting gamma oscillations (reflecting the prediction error) at the expense of alpha oscillations, as has been regularly observed (Hoogenboom et al., 2006; Potes et al., 2014).

Hence, there are two mechanisms with antagonistic effects on gamma oscillations in operation, as follows: (1) top-down predictions of a stimulus event (as in classic predictive coding schemes) are proposed to suppress the prediction errors per se and therefore, according to our hypotheses (Bastos et al., 2012), gamma oscillations; and (2) attentional synaptic gain regulation can boost the efficacy of prediction errors. The hypothesized interactions between both mechanisms (suppressive effects of prediction and attentional enhancement through gain modulation) have recently been supported in two studies. A study by Jiang et al. (2013) found that attention can boost the prediction error, such that the difference between the neuronal responses to expected and unexpected stimuli is amplified when these are attended. Kok et al. (2012) have found that attention can override the suppressive effects of top-down prediction signals, such that expected stimuli that are attended yield the largest responses in early sensory cortex. Results like those by Lima et al. (2011) and Schoffelen et al. (2005) can be interpreted as manifestations of the second mechanism we have described above (gain enhancement), whereas the suppressive effects seem less effective there. Schoffelen et al. (2005) focused on motor gamma oscillations where one would not expect a suppressive effect in the first place, while the discrete timing in a relatively long interval in the study by Lima et al. (2011) may similarly diminish the suppression of prediction errors.

In our study, attention was operationalized through the manipulation of the behavioral relevance of two simultaneously presented bilateral stimuli with a deterministic cue. Hence, only one side was of behavioral relevance and therefore the effect of target prediction should be dominant in the hemisphere contralateral to the preceded hemifield. The predictive coding perspective allows us to partition cortical responses into two components. The first corresponds to (prestimulus) activity due to anticipatory processing, while the second (poststimulus) response is induced by a target when it appears. In simple terms, the prestimulus activity can only reflect the confidence or precision that a target will appear in a future. We assumed that this is reflected in the desynchronization of alpha/beta activity, such that increasing precision or anticipation of an impending target increases cortical excitability and suppresses (desynchronizes) alpha activity. The second component is associated with the generation of prediction errors during the processing of the stimulus, which we associate with the amplitude of induced gamma responses. With this dissociation in mind, one would hypothesize that a target that was anticipated (later in peristimulus time) would be preceded by a desynchronization of alpha/beta power and a subsequent reduction in gamma activity, as prediction errors are explained away more efficiently. These dissociable effects reflect the increasing precision or confidence that a target will appear and the suppression of prediction errors when the target appears, respectively. These hazard rate or anticipation-dependent effects of the lateralized gamma oscillations are exactly what we observed. Nevertheless, in the light of the diversity of findings in the literature summarized here, we appreciate that more research is needed to complete our understanding of the specific interactions between suppressive and gain-enhancing effects of top-down influences within the context of predictive coding.

Furthermore, with respect to the systems generating top-down predictions of stimulus content and precision, the theoretical literature focuses on canonical microcircuits in a hierarchical network; however, dedicated brain structures have been associated with functions related to goal-directed attentional selection and stimulus-driven adaptive responses. For example, it is possible that the same mechanisms are implicated in the volitional control of selective attention and subjective precision (Corbetta and Shulman, 2002; Vossel et al., 2012). These studies suggest (dorsal) prefrontal and parietal sources of descending gain control that increase with stimulus anticipation. Conversely, if the stimulus appears relatively unexpectedly, the ventral system may boost forward propagation of unexpected stimulus changes through the hierarchy to enable a (relatively) rapid response, and this may then be reflected in the post-stimulus augmentation of gamma oscillations.

Thus, we have shown that lateralized attentional alpha modulation may reflect a (prestimulus) top-down precision or gain control that persists until the anticipated event occurs and correlates with the subjective confidence or precision of the anticipated sensory signals. Furthermore, our results give support to the idea that (poststimulus) gamma oscillations are implicated in forwarding prediction errors, and their strength may reflect the degree of surprise induced by predicted sensory information rather than simply reflecting the state of neuronal excitation driven by afferent stimulation.

References


