

Prior expectations induce prestimulus sensory templates

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Perception can be described as a process of inference, integrating bottom-up sensory inputs and top-down expectations. However, it is unclear how this process is neurally implemented. It has been proposed that expectations lead to prestimulus baseline increases in sensory neurons tuned to the expected stimulus, which in turn, affect the processing of subsequent stimuli. Recent fMRI studies have revealed stimulus-specific patterns of activation in sensory cortex as a result of expectation, but this method lacks the temporal resolution necessary to distinguish pre- from poststimulus processes. Here, we combined human magnetoencephalography (MEG) with multivariate decoding techniques to probe the representational content of neural signals in a time-resolved manner. We observed a representation of expected stimuli in the neural signal shortly before they were presented, showing that expectations indeed induce a preactivation of stimulus templates. The strength of these prestimulus expectation templates correlated with participants' behavioral improvement when the expected feature was task-relevant. These results suggest a mechanism for how predictive perception can be neurally implemented.

prediction | perceptual inference | predictive coding | feature-based expectation | feature-based attention

Perception is heavily influenced by prior knowledge (1–3). Accordingly, many theories cast perception as a process of inference, integrating bottom-up sensory inputs and top-down expectations (4–6). However, it is unclear how this integration is neurally implemented. It has been proposed that prior expectations lead to baseline increases in sensory neurons tuned to the expected stimulus (7–9), which in turn, leads to improved neural processing of matching stimuli (10, 11). In other words, expectations may induce stimulus templates in sensory cortex before the actual presentation of the stimulus. Alternatively, topdown influences in sensory cortex may exert their influence only after the bottom-up stimulus has been initially processed, and the integration of the two sources of information may become apparent only during later stages of sensory processing (12).

The evidence necessary to distinguish between these hypotheses has been lacking. fMRI studies have revealed stimulusspecific patterns of activation in sensory cortex as a result of expectation (9, 13), but this method lacks the temporal resolution necessary to distinguish pre- from poststimulus periods. Here, we combined magnetoencephalography (MEG) with multivariate decoding techniques to probe the representational content of neural signals in a time-resolved manner (14-17). The experimental paradigm was virtually identical to the ones used in our previous fMRI studies that studied how expectations modulate stimulus-specific patterns of activity in the primary visual cortex (9, 11). We trained a forward model to decode the orientation of task-irrelevant gratings from the MEG signal (18, 19) and applied this decoder to trials in which participants expected a grating of a particular orientation to be presented. This analysis revealed a neural representation of the expected grating that resembled the neural signal evoked by an actually presented grating. This representation was present already shortly before

stimulus presentation, showing that expectations can indeed induce the preactivation of stimulus templates.

Results

Participants (n = 23) were exposed to auditory cues that predicted the likely orientation $(45^{\circ} \text{ or } 135^{\circ})$ of an upcoming grating stimulus (Fig. 1 *A* and *B*). This grating was followed by a second grating that differed slightly from the first in terms of orientation and contrast. In separate runs of the MEG session, participants performed either an orientation or contrast discrimination task on the two gratings (details are in *Materials and Methods*).

Behavioral Results. Participants were able to discriminate small differences in orientation $(3.9^{\circ} \pm 0.5^{\circ})$, accuracy = 74.0 ± 1.6%, mean ± SEM) and contrast (4.6 ± 0.3%, accuracy = 76.6 ± 1.5%) of the cued gratings. There was no significant difference between the two tasks in terms of either accuracy ($F_{1,22} = 3.38$, P = 0.080) or reaction time (RT) (mean RT = 633 vs. 608 ms, $F_{1,22} = 2.89$, P = 0.10). Overall, accuracy and reaction times were not influenced by whether the cued grating had the expected or the unexpected orientation (accuracy: $F_{1,22} = 0.21$, P = 0.65; RT: $F_{1,22} < 0.01$, P = 0.93), and there was no interaction between task and expectation (accuracy: $F_{1,22} = 0.96$, P = 0.34; RT: $F_{1,22} = 0.09$, P = 0.77). Note that these discrimination tasks were orthogonal to the expectation manipulation in the sense that the expectation cue provided no information about the likely correct choice.

During the grating localizer (Fig. 1*C*; details are in *Materials* and *Methods*), participants correctly detected 91.2 \pm 1.6% (mean \pm SEM) of fixation flickers and incorrectly pressed the button on 0.2 \pm 0.1% of trials, suggesting that participants were successfully engaged by the fixation task.

Significance

The way that we perceive the world is partly shaped by what we expect to see at any given moment. However, it is unclear how this process is neurally implemented. Recently, it has been proposed that the brain generates stimulus templates in sensory cortex to preempt expected inputs. Here, we provide evidence that a representation of the expected stimulus is present in the neural signal shortly before it is presented, showing that expectations can indeed induce the preactivation of stimulus templates. Importantly, these expectation signals resembled the neural signal evoked by an actually presented stimulus, suggesting that expectations induce similar patterns of activations in visual cortex as sensory stimuli.

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Fig. 1. Experimental paradigm. (*A*) Each trial started with an auditory cue that predicted the orientation of the subsequent grating stimulus. This first grating was followed by a second one, which differed slightly from the first in terms of orientation and contrast. In separate runs, participants performed either an orientation or contrast discrimination task on the two gratings. (*B*) Throughout the experiment, two different tones were used as cues, each one predicting one of the two possible orientations (45° or 135°) with 75% validity. These contingencies were flipped halfway through the experiment. (*C*) In separate grating localizer runs, participants were exposed to task-irrelevant gratings while they performed a fixation dot dimming task.

MEG Results-Localizer Orientation Decoding. As mentioned, participants were exposed to auditory cues that predicted the likely orientation of an upcoming grating stimulus. The question that we wanted to answer was whether the expectations induced by these auditory cues would evoke templates of the visual stimuli before the presentation of the gratings. To be able to uncover such sensory templates, we trained a decoding model to reconstruct the orientation of (task-irrelevant) visual gratings (Fig. 1C) from the MEG signal in a time-resolved manner. We found that this model was highly accurate at reconstructing the orientation of such gratings from the MEG signal (Fig. 2). Grating orientation could be decoded across an extended period (from 40 to 655 ms poststimulus, P < 0.001 and from 685 to 730 ms, P = 0.018), peaking around 120-160 ms poststimulus (Fig. 2C). Furthermore, in the period around 100-330 ms poststimulus, orientation decoding generalized across time, meaning that a decoder trained on the evoked response at, for example, 120 ms poststimulus could reconstruct the grating orientation represented in the evoked response around 300 ms and vice versa (Fig. 2D). In other words, certain aspects of the representation of grating orientation were sustained over time.

MEG Results—Expectation Induces Stimulus Templates. Our main question pertained to the presence of visual grating templates induced by the auditory expectation cues during the main experiment. Therefore, we applied our model trained on task-irrelevant gratings to trials containing gratings that were either validly or invalidly predicted (Fig. 3*A*). In both conditions, the decoding model trained on task-irrelevant gratings succeeded in accurately reconstructing the orientation of the gratings presented in the main experiment (valid expectation: cluster from training time 60–410 ms and decoding time 60–400 ms, P < 0.001 and from training time 205–325 ms and decoding time 400–495 ms, P = 0.045; invalid expectation: cluster from training time 75–225 ms and decoding time 75–330 ms, P = 0.0012 and from training time 250–360 ms and decoding time 195–355 ms, P = 0.027).

If the cues induced sensory templates of the expected grating, one would expect these to be revealed in the difference in decoding between validly and invalidly predicted gratings (details of the subtraction logic are in *Materials and Methods*). Indeed, this analysis showed that the auditory expectation cues induce orientation-specific neural signals (Fig. 3A, *Bottom*). These signals were present already 40 ms before grating presentation and extended into the poststimulus period (from decoding time -40 to 230 ms, P = 0.0092 and from 300 to 530 ms, P = 0.016). Furthermore, these signals were uncovered when the decoder was trained on around 120–160 ms poststimulus during the grating localizer (Fig. 3B), suggesting that these cue-induced signals were similar to those evoked by taskirrelevant gratings. In other words, the auditory expectation cues evoked orientation-specific signals that were similar to sensory signals evoked by the corresponding actual grating stimuli (Fig. S1A).

In sum, expectations induced prestimulus sensory templates that influenced poststimulus representations as well; invalidly expected gratings had to "overcome" a prestimulus activation of the opposite orientation, while validly expected gratings were facilitated by a compatible prestimulus activation (Fig. S1B). The poststimulus carryover of these expectation signals lasted throughout the trial (Fig. S1C).

As in previous studies using a similar paradigm (11, 20), there was no interaction between the effects of the expectation cue and the task (orientation vs. contrast discrimination) that participants performed (no clusters with P < 0.05) (Fig. S2A). In other words, expectations evoked prestimulus orientation signals to a similar degree in both tasks (Fig. S2B). This suggests that influences of expectation on neural representations are relatively independent of the task relevance of the expected feature, in line with our previous fMRI study (11). Note, however, that, unlike in that study, there was no significant modulation of the orientation signal by task relevance (no clusters with P < 0.05) (Fig. S2A). The reason for this lack of difference is unclear, although it should be noted that there was a trend toward participants having higher accuracy and faster reaction times (see above) on the contrast task than on the orientation task. This may suggest that the two tasks were not optimally balanced in terms of difficulty, precluding a proper comparison of the effect of task set in this study.

In our previous fMRI study, we found a relationship between the effects of expectation on neural stimulus representations and



Fig. 2. Localizer orientation decoding. (A) The output of the decoder consisted of the responses of 32 hypothetical orientation channels; shown here are decoders trained and tested on the MEG signal 120–160 ms poststimulus during the grating localizer (cross-validated). Shaded region represents SEM. (B) Decoder output over time, trained and tested in 5-ms steps (sliding window of 29.2 ms), showing the temporal evolution of the orientation signal. (C) The response of the 32 orientation channels collapsed into a single metric of decoding performance (*SI Materials and Methods*) over time. Shaded region represents SEM; horizontal lines indicate significant clusters (P < 0.05). (D) Temporal generalization matrix of orientation decoding performance obtained by training decoders on each time point and testing all decoders on all time points (as above, steps of 5 ms and a sliding window of 29.2 ms). This method provides insight into the sustained vs. dynamical nature of orientation representations (15). Solid black lines indicate significant clusters (P < 0.05); dashed lines indicate grating onset (t = 0 s).

performance on the orientation discrimination task. Specifically, participants for whom valid expectations led to the largest improvement in neural stimulus representations also showed the strongest benefit of valid expectations on behavioral performance during the orientation discrimination task (11). This relationship was absent for the contrast discrimination task when grating orientation was task-irrelevant. This study allowed us to test for a similar relationship, with an important extension: here, we could test whether neural prestimulus expectation signals are related to behavioral performance improvements. We quantified the decoding of the expected orientation just before grating presentation (-50 to 0 ms, training window 120 to 160 ms) and correlated this with the difference in task accuracy for valid and invalid expectation trials, across participants. This analysis revealed that participants with a stronger prestimulus reflection of the expected orientation in their neural signal also had a greater benefit from valid expectations on performance on the orientation task (r = 0.44, P = 0.035) (Fig. 4, *Left*). No such relationship was found for the contrast task, where the orientation of the gratings was not taskrelevant (r = -0.13, P = 0.55) (Fig. 4, *Right*). This is exactly the pattern of results that we found in our previous fMRI study but with the important extension that it is the prestimulus expectation effect that is correlated with behavioral performance, whereas the previous study did not have the temporal resolution to distinguish prefrom poststimulus signals.

In this study, neural orientation signals were probed by applying a forward model that takes the noise covariance between MEG sensors into account (details are in *SI Materials and Methods*). This model was superior to a forward model that did not correct for the noise covariance (Fig. S3), suggesting that feature covariance is an important factor to take into account when applying multivariate methods to MEG data. Corroborating this notion, a two-class decoder that corrected for noise covariance (16) was able to reproduce our effects of interest (Fig. S4), showing that the expectation effects do not depend on a specific analysis technique as long as the covariance between MEG sensors is taken into account.

Finally, there was no difference in the overall amplitude of the neural response evoked between validly and invalidly expected gratings (no clusters with P < 0.4) (Fig. S5).

Discussion

Here, we show that expectations can induce sensory templates of the expected stimulus already before the stimulus appears. These results extend previous fMRI studies showing stimulus-specific patterns of activation in sensory cortex induced by expectations that could not resolve whether these templates indeed reflected prestimulus expectations or instead, stimulus specific error signals induced by the unexpected omission of a stimulus (9, 13). Furthermore, the strength of these prestimulus expectation signals correlated with the behavioral benefit of a valid expectation when the expected feature (i.e., orientation) was task-relevant (11). These results suggest that valid expectations facilitate perception by allowing sensory cortex to prepare for upcoming sensory signals. As in a previous fMRI study using a very similar experimental paradigm (11), the neural effects of orientation expectations reported here were independent of the task relevance of



Fig. 3. Expectation induces stimulus templates. (A) Temporal generalization matrices of orientation decoding during the main experiment. Decoders were trained on the grating localizer (training time on the *y* axis) and tested on the main experiment (time on the *x* axis; dashed vertical line indicates t = 0 s, onset of the first grating). Decoding shown separately for gratings preceded by a valid expectation (*Top*), an invalid expectation (*Middle*), and the subtraction of the two conditions (i.e., the expectation cue effect; *Bottom*). Solid black lines indicate significant clusters (*P* < 0.05). (*B*) Orientation decoding during the main task averaged over training time 120–160 ms poststimulus during the grating localizer. That is, *B* shows a horizontal slice through the temporal generalization matrices above at the training time for which we see a significant cluster of expected orientation decoding, for visualization. Shaded regions indicate SEM.

the orientation of the gratings, suggesting that the generation of expectation templates may be an automatic phenomenon.

The fact that expectation signals were revealed by a decoder trained on physically presented (but task-irrelevant) gratings suggests that these expectation signals resemble activity patterns induced by actual stimuli. The expectation signal remained present throughout the trial and extended into the poststimulus period, suggesting the tonic activation of a stimulus template. These results are in line with a recent monkey electrophysiology study (10), which showed that neurons in the face patch of inferior temporal cortex encode the prior expectation of a face appearing both before and after actual stimulus presentation. When the subsequently presented stimulus is noisy or ambiguous, such a prestimulus template could conceivably bias perception toward the expected stimulus (21–24).

What is the source of these cue-induced expectation signals? One candidate region is the hippocampus, which is known to be involved in encoding associations between previously unrelated discontiguous stimuli (25), such as the auditory tones and visual gratings used in this study. Furthermore, fMRI studies have revealed predictive signals in the hippocampus (13, 26, 27), and Reddy et al. (28) reported anticipatory firing to expected stimuli in the medial temporal lobe, including the hippocampus. One intriguing possibility is that predictive signals from the hippocampus are fed back to sensory cortex (13, 29, 30).

Previous studies have suggested, both on theoretical (31) and on empirical (32, 33) grounds, that top-down (prediction) and bottom-up (stimulus-driven or prediction error) signals are subserved by distinct frequency bands. Therefore, one highly interesting direction for future research would be to determine whether the expectation templates revealed here are specifically manifested in certain frequency bands (i.e., the alpha or beta band).

In addition to expectation, several other cognitive phenomena have been shown to induce stimulus templates in sensory cortex, such as preparatory attention (17, 34), mental imagery (35-37), and working memory (38, 39). In fact, explicit task preparation can also induce prestimulus sensory templates that last into the poststimulus period (17). Note that, in this study, the task did not require explicit use of the expectation cues, and the task response was, in fact, orthogonal to the expectation. Furthermore, there was no difference in the expectation signal between runs in which grating orientation was task-relevant (orientation discrimination task) and when it was irrelevant (contrast discrimination task); suggesting expectation may be a relatively automatic phenomenon (11, 40). In fact, neural modulations by expectation have even been observed during states of inattention (41), in sleep (42), and in patients experiencing disorders of consciousness (43). One important question for future research will be to establish whether the same neural mechanism underlies the different cognitive phenomena that are capable of inducing stimulus templates in sensory cortex or whether different top-down mechanisms are at work. Indeed, it has been suggested that expectation and attention, or task preparation, may have different underlying neural mechanisms (20, 44, 45). For instance, predictive coding theories suggest that attention may modulate sensory signals in the superficial layers of sensory cortex, while predictions modulate the response in deep layers (5, 46).

One may wonder why this study does not report a modulation of the overall neural response by expectation, while previous studies have found an increased neural response to unexpected stimuli (40, 47–51), including some using an almost identical paradigm as this study (11, 20). Of course, this study reports a null effect, from which it is hard to draw firm conclusions. However, it is possible that the type of measurement of neural activity plays a role in the absence of the effect. Most previous studies reporting expectation suppression in visual cortex used fMRI, whereas this study used MEG. It is possible that the blood oxygen level-dependent (BOLD) signal, a



Fig. 4. Correlation between neural expectation signals and behavioral improvement by expectation. Neural prestimulus expectation decoding (on the x axis) correlated with behavioral improvement induced by valid expectations (on the y axis) during the orientation discrimination task (*Left*). This correlation was absent during the contrast discrimination task (*Right*).

mass action signal that integrates synaptic and neural activity as well as integrating over time, is sensitive to certain neural effects that MEG, which is predominantly sensitive to synchronized activity in pyramidal neurons oriented perpendicular to the cortical surface, is not. It is even possible that, within MEG, different types of sensors (i.e., magnetometers, planar and axial gradiometers) differ in their sensitivity to expectation suppression (52).

Recent theories of sensory processing state that perception reflects the integration of bottom-up inputs and top-down expectations, but ideas diverge on whether the brain continuously generates stimulus templates in sensory cortex to preempt expected inputs (10, 23, 53, 54) or rather, engages in perceptual inference only after receiving sensory inputs (55, 56). Our results are in line with the brain being proactive and constantly forming predictions about future sensory inputs. These findings bring us closer to uncovering the neural mechanisms by which we integrate prior knowledge with sensory inputs to optimize perception.

Materials and Methods

Participants. Twenty-three (15 female, age 26 \pm 9 y old, mean \pm SD) healthy individuals participated in the MEG experiment. All participants were right-handed and had normal or corrected to normal vision. The study was approved by the local ethics committee [Commisie Mensgebonden Onderzoek (CMO) Arnhem-Nijmegen, The Netherlands] under the general ethics approval (Imaging Human Cognition, CMO 2014/288), and the experiment was conducted in accordance with these guidelines. All participants gave written informed consent according to the Declaration of Helsinki.

Experimental Design. Each trial consisted of an auditory cue followed by two consecutive grating stimuli (750-ms stimulus-onset asynchrony between auditory and first visual stimulus) (Fig. 1A). The two grating stimuli were presented for 250 ms each separated by a blank screen (500 ms). A central fixation bull's eye (0.7°) was presented throughout the trial as well as during the intertrial interval (ITI; 2,250 ms). The auditory cue consisted of either a low- (500 Hz) or high-frequency (1,000 Hz) tone, which predicted the orientation of the first grating stimulus (45° or 135°) with 75% validity (Fig. 1B). In the other 25% of trials, the first grating had the orthogonal orientation. Thus, the first grating had an orientation of either exactly 45° or 135° and a luminance contrast of 80%. The second grating differed slightly from the first in terms of both orientation and contrast (see below) as well as being in antiphase to the first grating (which had a random spatial phase). The contingencies between the auditory cues and grating orientations were flipped halfway through the experiment (i.e., after four runs), and the order was counterbalanced over subjects.

In separate runs (64 trials each, \sim 4.5 min), subjects performed either an orientation or a contrast discrimination task on the two gratings. When performing the orientation task, subjects had to judge whether the second grating was rotated clockwise or anticlockwise with respect to the first grating. In the contrast task, a judgment had to be made on whether the second grating had lower or higher contrast than the first one. These tasks

were explicitly designed to avoid a direct relationship between the perceptual expectation and the task response. Furthermore, as in a previous fMRI study (11), these two different tasks were designed to manipulate the task relevance of the grating orientations to investigate whether the effects of orientation expectations depend on the task relevance of the expected feature.

Interleaved with the main task runs, subjects performed eight runs of a grating localizer task (Fig. 1C). Each run (~2 min) consisted of 80 grating presentations (ITI uniformly jittered between 1,000 and 1,200 ms). The grating annuli were identical to those presented during the main task (80% contrast, 250-ms duration, 1.0 cycle per 1°, random spatial phase). Each grating had one of eight orientations (spanning the 180° space, starting at 0°, in steps of 22.5°), each of which was presented 10 times per run in pseudorandom order. A black fixation bull's eye (4 cd/m², 0.7° diameter, identical to the one presented during the main task runs) was presented throughout the run. On 10% of trials (counterbalanced across orientations), the black fixation point in the center of the bull's eye (0.2°, 4 cd/m²) briefly turned gray (324 cd/m²) during the first 50 ms of grating presentation. Participants' task was to press a button (response deadline: 500 ms) when they perceived this fixation flicker. This simple task was meant to ensure central fixation, while rendering the gratings task-irrelevant. Trials containing fixation flickers were excluded from additional analyses.

Orientation Decoding Analysis. To probe sensory representations in the visual cortex, we used a forward modeling approach to reconstruct the orientation of the grating stimuli from the MEG signal (17–19, 57). This method has been shown to be highly successful at reconstructing circular stimulus features, such as color (18), orientation (17, 19, 57), and motion direction (22), from neural signals. Neural representations in MEG signals have also been successfully investigated using binomial classifiers (58); however, when it comes to a continuous stimulus feature, such as orientation, forward model reconstructions provide a richer decoding signal than binomial classifier accuracy (59). We made certain changes to the forward model proposed by Brouwer and Heeger (18) (most notably, taking the noise covariance into account; details are in SI Materials and Methods) to optimize it for MEG data, given the high correlations between neighboring sensors, based on ref. 16. In sum, this previously published and theoretically motivated decoding model was optimally suited for recovering a continuous feature from MEG data. For our main analyses, the forward model was trained on the data from the localizer runs, in which the gratings were task-irrelevant, and then applied to the main task data to uncover sensory templates induced by prestimulus expectations (details are in SI Materials and Methods). Our effects of interest (Fig. 3 and Fig. S6) were reproduced using a two-class decoder (Fig. S4).

The full methods can be found in SI Materials and Methods.

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