

1 **Prior expectations induce pre-stimulus sensory templates**

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16 analysed the data, P.K., P.M. and F.d.L. wrote the paper.

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35 **Abstract**

36

37 Perception can be described as a process of inference, integrating bottom-up sensory inputs and top-
38 down expectations. However, it is unclear how this process is neurally implemented. It has been
39 proposed that expectations lead to pre-stimulus baseline increases in sensory neurons tuned to the
40 expected stimulus, which in turn affects the processing of subsequent stimuli. Recent fMRI studies have
41 revealed stimulus-specific patterns of activation in sensory cortex as a result of expectation, but this
42 method lacks the temporal resolution necessary to distinguish pre- from post-stimulus processes. Here,
43 we combined human MEG with multivariate decoding techniques to probe the representational content
44 of neural signals in a time-resolved manner. We observed a representation of expected stimuli in the
45 neural signal well before they were presented, demonstrating that expectations indeed induce a pre-
46 activation of stimulus templates. These results suggest a mechanism for how predictive perception can
47 be neurally implemented.

48 **Introduction**

49

50 Perception is heavily influenced by prior knowledge (Helmholtz, 1866; Gregory, 1997; Kersten et al.,
51 2004). Accordingly, many theories cast perception as a process of inference, integrating bottom-up
52 sensory inputs and top-down expectations (Lee and Mumford, 2003; Friston, 2005; Summerfield and de
53 Lange, 2014). However, it is unclear how this integration is neurally implemented. It has been proposed
54 that prior expectations lead to baseline increases in sensory neurons tuned to the expected stimulus
55 (Wyart et al., 2012; SanMiguel et al., 2013; Kok et al., 2014), which in turn leads to improved neural
56 processing of matching stimuli (Kok et al., 2012; Bell et al., 2016). In other words, expectations may
57 induce stimulus templates in sensory cortex, prior to the actual presentation of the stimulus.
58 Alternatively, top-down influences in sensory cortex may exert their influence only after the bottom-up
59 stimulus has been initially processed, and the integration of the two sources of information may become
60 apparent only during later stages of sensory processing (Rao et al., 2012).

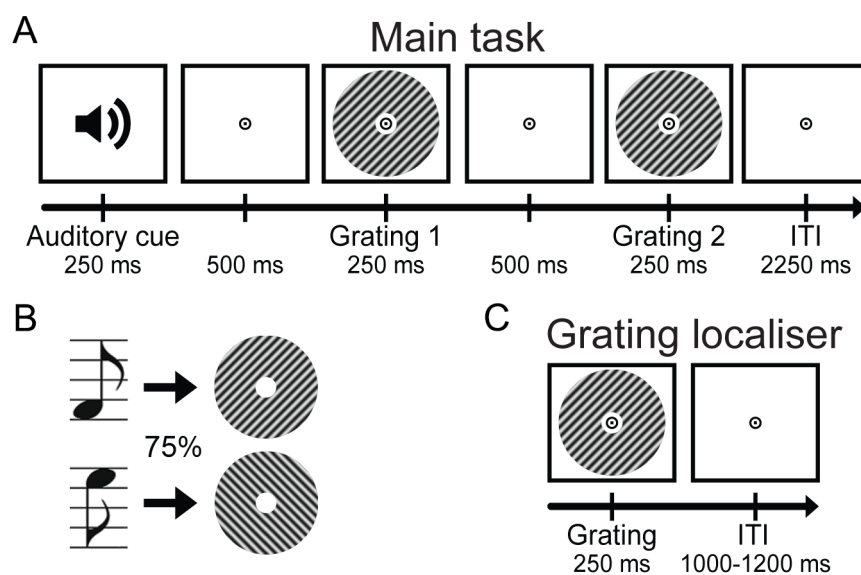
61 The evidence necessary to distinguish between these hypotheses has been lacking. fMRI studies
62 have revealed stimulus-specific patterns of activation in sensory cortex as a result of expectation (Kok et
63 al., 2014; Hindy et al., 2016), but this method lacks the temporal resolution necessary to distinguish pre-
64 from post-stimulus periods. Here, we combined MEG with multivariate decoding techniques to probe
65 the representational content of neural signals in a time-resolved manner (Cichy et al., 2014; King and
66 Dehaene, 2014; Mostert et al., 2015; Myers et al., 2015). We trained a forward model to decode the
67 orientation of task-irrelevant gratings from the MEG signal (Brouwer and Heeger, 2009, 2011), and
68 applied this decoder to trials in which participants expected a grating of a particular orientation to be
69 presented. This analysis revealed a neural representation of the expected grating that resembled the
70 neural signal evoked by an actually presented grating. This representation was present already before
71 stimulus presentation, demonstrating that expectations can indeed induce the pre-activation of stimulus
72 templates.

73 Results

74

75 Participants were exposed to auditory cues that predicted the likely orientation (45° or 135°) of an
76 upcoming grating stimulus (Figure 1A-B). This grating was followed by a second grating that differed
77 slightly from the first in terms of orientation and contrast. In separate runs of the MEG session,
78 participants performed either an orientation or contrast discrimination task on the two gratings (see
79 Materials and Methods for details).

80



81

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

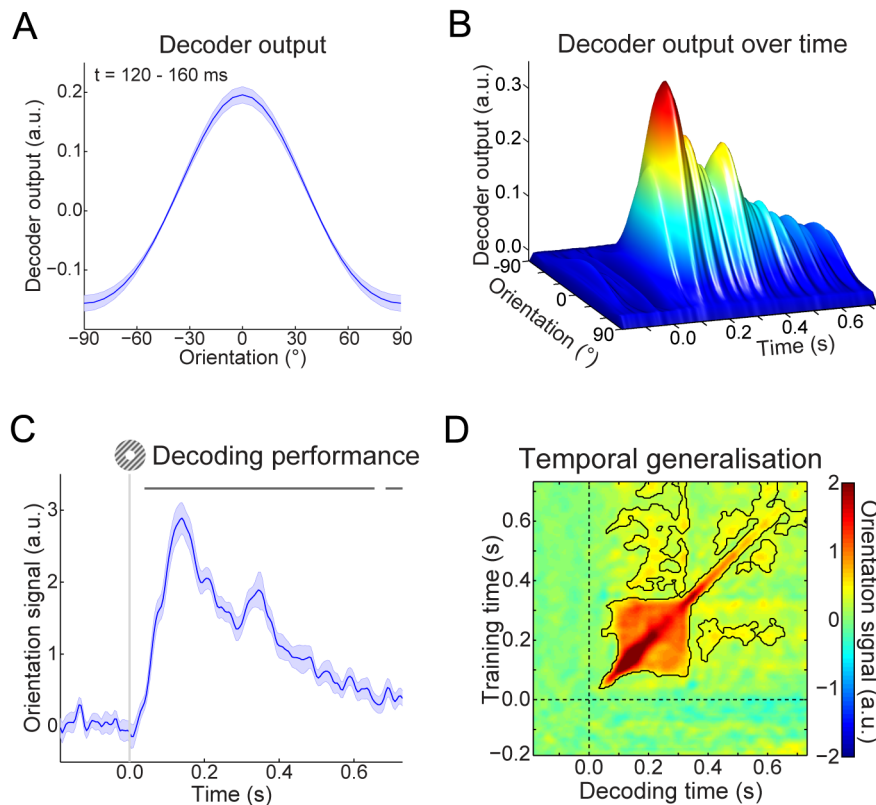
90

91 **Behavioural results.** Participants were able to discriminate small differences in orientation ($3.9^\circ \pm 0.5^\circ$,
92 accuracy = $74.0\% \pm 1.6\%$, mean \pm sem) and contrast ($4.6\% \pm 0.3\%$, accuracy = $76.6\% \pm 1.5\%$) of the cued
93 gratings. There was no significant difference between the two tasks in terms of either accuracy ($F_{1,22} =$
94 3.38 , $p = 0.080$) or reaction time (mean RT = 621 ms vs. 603 ms, $F_{1,22} = 1.46$, $p = 0.24$). Overall, accuracy
95 and reaction times were not influenced by whether the cued grating had the expected or the
96 unexpected orientation (accuracy: $F_{1,22} = 0.21$, $p = 0.65$; RT: $F_{1,22} = 0.03$, $p = 0.87$), nor was there an
97 interaction between task and expectation (accuracy: $F_{1,22} = 0.96$, $p = 0.34$; RT: $F_{1,22} = 0.42$, $p = 0.52$). Note
98 that these discrimination tasks were orthogonal to the expectation manipulation, in the sense that the
99 expectation cue provided no information about the likely correct choice.

100 During the grating localiser (Figure 1C, see Materials and Methods for details), participants
101 correctly detected $91.2\% \pm 1.6\%$ (mean \pm sem) of fixation flickers, and incorrectly pressed the button on
102 $0.2\% \pm 0.1\%$ of trials, suggesting that participants were successfully engaged by the fixation task.

103

104 **MEG results – Localiser orientation decoding.** As mentioned, participants were exposed to auditory
105 cues that predicted the likely orientation of an upcoming grating stimulus. The question we wanted to
106 answer was whether the expectations induced by these auditory cues would evoke templates of the
107 visual stimuli prior to the presentation of the gratings. To be able to uncover such sensory templates, we
108 trained a decoding model to reconstruct the orientation of (task-irrelevant) visual gratings (Figure 1C)
109 from the MEG signal, in a time-resolved manner. First, we found that this model was highly accurate at
110 reconstructing the orientation of such gratings from the MEG signal (Figure 2). Grating orientation could
111 be decoded across an extended period of time (from 40 to 655 ms post-stimulus, $p < 0.001$, and from
112 685 to 730 ms, $p = 0.018$), peaking around 120-160 ms post-stimulus (Figure 2C). Furthermore, in the
113 period around 100 to 330 ms post-stimulus, orientation decoding generalised across time, meaning that
114 a decoder trained on the evoked response at, for example, 120 ms post-stimulus could reconstruct the
115 grating orientation represented in the evoked response around 300 ms, and vice versa (Figure 2D). In
116 other words, certain aspects of the representation of grating orientation were sustained over time.



117

118 **Figure 2. Localiser orientation decoding.** (A) The output of the decoder consisted of the responses of 32
119 hypothetical orientation channels, shown here decoders trained and tested on the MEG signal 120-160
120 ms post-stimulus during the grating localiser (cross-validated). Shaded region represent SEM. (B)
121 Decoder output over time, trained and tested in 5 ms steps (sliding window of 29.2 ms), showing the
122 temporal evolution of the orientation signal. (C) The response of the 32 orientation channels collapsed
123 into a single metric of decoding performance (see Materials and Methods), over time. Shaded region
124 represent SEM, horizontal lines indicate significant clusters ($p < 0.05$). (D) Temporal generalisation
125 matrix of orientation decoding performance, obtained by training decoders on each time point, and
126 testing all decoders on all time points (as above, steps of 5 ms and a sliding window of 29.2 ms). This
127 method provides insight into the sustained versus dynamical nature of orientation representations (King
128 and Dehaene, 2014). Solid black lines indicate significant clusters ($p < 0.05$), dashed lines indicate grating
129 onset ($t = 0s$).

130

131

132 **MEG results – Expectation induces stimulus templates.** Our main question pertained to the presence of
133 visual grating templates induced by the auditory expectation cues during the main experiment.
134 Therefore, we applied our model trained on task-irrelevant gratings to trials containing gratings that
135 were either validly or invalidly predicted, respectively (Figure 3A). In both conditions, the decoding
136 model trained on task-irrelevant gratings succeeded in accurately reconstructing the orientation of the
137 gratings presented in the main experiment (valid expectation: cluster from training time 60 to 410 ms
138 and decoding time 60 to 400 ms, $p < 0.001$, and from training time 205 to 325 ms and decoding time 400
139 to 495 ms, $p = 0.045$; invalid expectation: cluster from training time 75 to 225 ms and decoding time 75
140 to 330 ms, $p = 0.0012$, and from training time 250 to 360 ms and decoding time 195 to 355 ms, $p =$
141 0.027).

142 If the cues induced sensory templates of the expected grating, one would expect these to be
143 revealed in the difference in decoding between valid and invalidly predicted gratings (see Material and
144 Methods for details of the subtraction logic). Indeed, this subtraction/analyses demonstrates that the
145 auditory expectation cues induce orientation-specific neural signals (Figure 3A, bottom panel). These
146 signals were present already 40 ms before grating presentation, and extended into the post-stimulus
147 period (from decoding time -40 to 230 ms, $p = 0.0092$, and from 300 to 530 ms, $p = 0.016$). Furthermore,
148 these signals were uncovered when the decoder was trained on around 120 to 160 ms post-stimulus
149 during the grating localiser (Figure 3B), suggesting that these cue-induced signals were similar to those
150 evoked by task-irrelevant gratings. In other words, the auditory expectation cues evoked orientation-
151 specific signals that were similar to sensory signals evoked by the corresponding actual grating stimuli.

152 In sum, expectations induced pre-stimulus sensory templates that influenced post-stimulus
153 representations as well; invalidly expected gratings had to ‘overcome’ a pre-stimulus activation of the
154 opposite orientation, while validly expected gratings were facilitated by a compatible pre-stimulus
155 activation (Figure 3 – figure supplement 2A). The post-stimulus carryover of these expectation signals
156 lasted throughout the trial (Figure 3 – figure supplement 2B).

157 As in previous studies using a similar paradigm (Kok et al., 2012, 2016b), there was no
158 interaction between the effects of the expectation cue and the task (orientation vs. contrast

159 discrimination) participants performed (no clusters with $p < 0.4$).

160 In the current study, there was no difference in the overall amplitude of the neural response
161 evoked between validly and invalidly expected gratings (no clusters with $p < 0.4$, Figure 3 – figure
162 supplement 3).

163

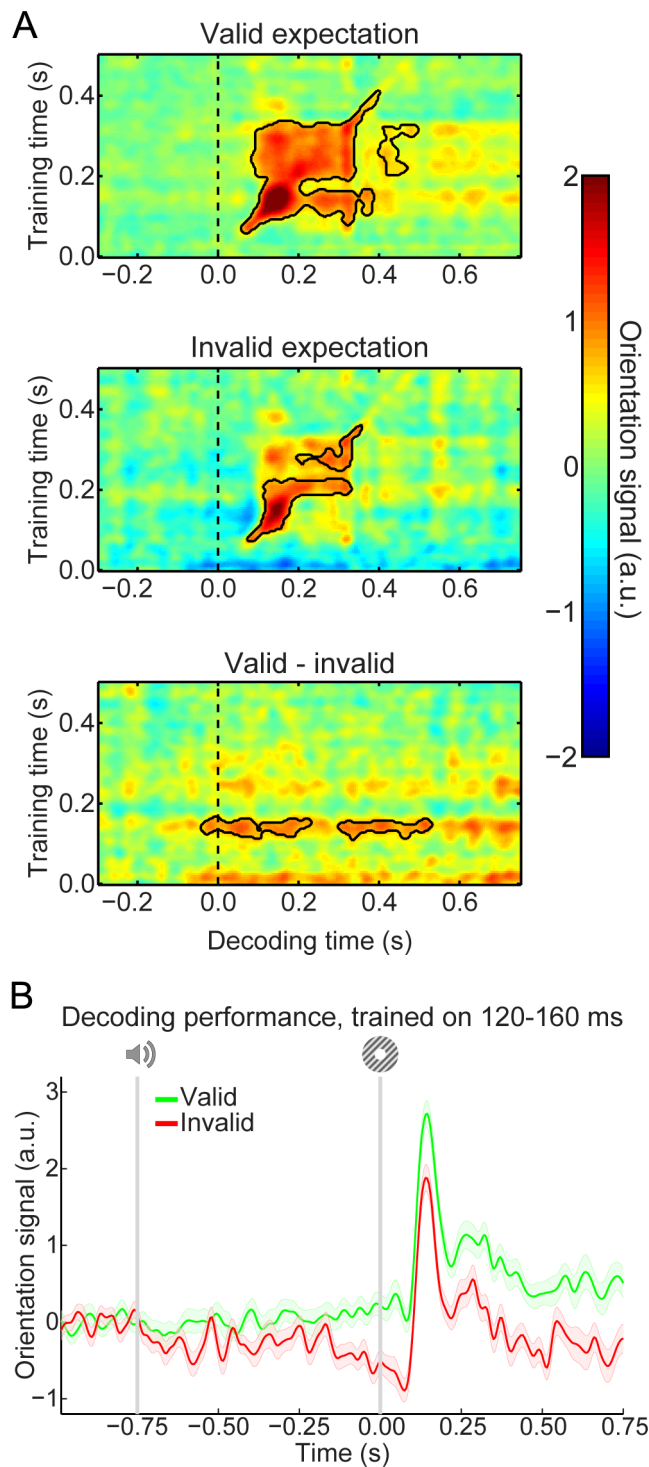


Figure 3. Expectation induces stimulus templates. (A) Temporal generalisation matrices of orientation decoding during the main experiment. Decoders were trained on the grating localiser (training time on the y-axis) and tested on the main experiment (time on the x-axis; dashed vertical line indicates $t = 0$ s, onset of the first grating). Decoding shown separately for gratings preceded by a valid expectation (top row), invalid expectation (middle row), and the subtraction of the two conditions (i.e., the expectation cue effect, bottom row). Solid black lines indicate significant clusters ($p < 0.05$). (B) Orientation decoding during the main task, averaged over training time 120 – 160 ms post-stimulus during the grating localiser. That is, a horizontal slice through the temporal generalisation matrices above at the training time for which we see a significant cluster of expected orientation decoding, for visualisation. Shaded regions indicate SEM.

186 The following figure supplements are available for figure 3:

187 **Figure supplement 1. Expectation effects in the absence of low-pass filtering.**

188 **Figure supplement 2. Expectation affects orientation decoding throughout the trial.**

189 **Figure supplement 3. Event related fields.**

190

191 **Discussion**

192

193 Here, we show that expectations can induce sensory templates of the expected stimulus already before
194 the stimulus appears. These results extend previous fMRI studies demonstrating stimulus-specific
195 patterns of activation in sensory cortex induced by expectations, which could not resolve whether these
196 templates indeed reflected pre-stimulus expectations, or instead stimulus specific error signals induced
197 by the unexpected omission of a stimulus (Kok et al., 2014; Hindy et al., 2016).

198 The fact that expectation signals were revealed by a decoder trained on physically presented
199 (but task-irrelevant) gratings suggests that these expectation signals resemble activity patterns induced
200 by actual stimuli. The expectation signal remained present throughout the trial, extending into the post-
201 stimulus period, suggesting the tonic activation of a stimulus template. These results are in line with a
202 recent monkey electrophysiology study (Bell et al., 2016), which showed that neurons in the face patch
203 of IT cortex encode the prior expectation of a face appearing, both prior to and following actual stimulus
204 presentation. When the subsequently presented stimulus is noisy or ambiguous, such a pre-stimulus
205 template could conceivably bias perception towards the expected stimulus (Chalk et al., 2010; Kok et al.,
206 2013; Pajani et al., 2015; John-Saaltink et al., 2016).

207 What is the source of these cue-induced expectation signals? One candidate region is the
208 hippocampus, which is known to be involved in encoding associations between previously unrelated,
209 discontinuous stimuli (Wallenstein et al., 1998), such as the auditory tones and visual gratings used in
210 the present study. Furthermore, fMRI studies have revealed predictive signals in the hippocampus
211 (Schapiro et al., 2012; Davachi and DuBrow, 2015; Hindy et al., 2016), and Reddy and colleagues (Reddy
212 et al., 2015) reported anticipatory firing to expected stimuli in the medial temporal lobe, including the

213 hippocampus. One intriguing possibility is that predictive signals from the hippocampus are fed back to
214 sensory cortex (Lavenex and Amaral, 2000; Bosch et al., 2014; Hindy et al., 2016).

215 In addition to expectation, several other cognitive phenomena have been shown to induce
216 stimulus templates in sensory cortex, such as preparatory attention (Stokes et al., 2009b; Myers et al.,
217 2015), mental imagery (Stokes et al., 2009a; Lee et al., 2012; Albers et al., 2013), and working memory
218 (Harrison and Tong, 2009; Serences et al., 2009). In fact, explicit task preparation can also induce pre-
219 stimulus sensory templates that last into the post-stimulus period (Myers et al., 2015). Note that in the
220 current study the task did not require explicit use of the expectation cues, the task response was in fact
221 orthogonal to the expectation. Furthermore, there was no difference in the expectation signal between
222 runs in which grating orientation was task-relevant (orientation discrimination task) and when it was
223 irrelevant (contrast discrimination task), suggestion expectation may be a relatively automatic
224 phenomenon (den Ouden et al., 2009; Kok et al., 2012). In fact, neural modulations by expectation have
225 even been observed during states of inattention (Näätänen, 1990), sleep (Nakano et al., 2008) and in
226 patients experiencing disorders of consciousness (Bekinschtein et al., 2009). One important question for
227 future research will be to establish whether the same neural mechanism underlies the different
228 cognitive phenomena that are capable of inducing stimulus templates in sensory cortex, or whether
229 different top-down mechanisms are at work. Indeed, it has been suggested that expectation and
230 attention, or task preparation, may have different underlying neural mechanisms (Summerfield and
231 Egnér, 2009; Kok et al., 2016b; Summerfield and Egnér, 2016). For instance, predictive coding theories
232 suggest that attention may modulate sensory signals in the superficial layers of sensory cortex, while
233 predictions modulate the response in deep layers (Friston, 2005; Kok et al., 2016a).

234 One may wonder why the current study does not report a modulation of the overall neural
235 response by expectation, while previous studies have found an increased neural response to unexpected
236 stimuli (Summerfield et al., 2008; den Ouden et al., 2009; Alink et al., 2010; Meyer and Olson, 2011;
237 Todorovic et al., 2011; Wacongne et al., 2011), including some using an almost identical paradigm as the
238 current study (Kok et al., 2012, 2016b). Of course, the current study reports a null effect, from which it is
239 hard to draw firm conclusions. However, it is possible that the type of measurement of neural activity

240 plays a role in the absence of the effect. Most previous studies reporting expectation suppression in
241 visual cortex used fMRI, while the current study used MEG. It is possible that the BOLD signal, a mass-
242 action signal that integrates synaptic and neural activity, as well as integrating over time, is sensitive to
243 certain neural effects that MEG, which is predominantly sensitive to synchronised activity in pyramidal
244 neurons oriented perpendicular to the cortical surface, is not. It is even possible that within MEG,
245 different types of sensors (i.e. magnetometers, planar and axial gradiometers) differ in their sensitivity
246 to expectation suppression (Cashdollar et al., 2016).

247 Recent theories of sensory processing state that perception reflects the integration of bottom-
248 up inputs and top-down expectations, but ideas diverge on whether the brain continuously generates
249 stimulus templates in sensory cortex to pre-empt expected inputs (Berkes et al., 2011; Pajani et al., 2015;
250 Bell et al., 2016; Fiser et al., 2016), or rather engages in perceptual inference only after receiving sensory
251 inputs (Rao and Ballard, 1999; Bar et al., 2006). Our results are in line with the brain being proactive,
252 constantly forming predictions about future sensory inputs. These findings bring us closer to uncovering
253 the neural mechanisms by which we integrate prior knowledge with sensory inputs to optimise
254 perception.

255 **Materials and Methods**

256

257 **Participants.** Twenty-three (15 female, age 26 ± 9 , mean \pm SD) healthy individuals participated in the
258 experiment. All participants were right-handed and had normal or corrected-to-normal vision. The study
259 was approved by the local ethics committee (CMO Arnhem-Nijmegen, The Netherlands) under the
260 general ethics approval (“Imaging Human Cognition”, CMO 2014/288), and the experiment was
261 conducted in accordance with these guidelines. All participants gave written informed consent
262 according to the declaration of Helsinki.

263

264 **Stimuli.** Grayscale luminance-defined sinusoidal grating stimuli (spatial frequency: 1.0 cycles/°) were
265 generated using MATLAB (MathWorks, Natick, MA) in conjunction with the Psychophysics Toolbox
266 (Brainard, 1997). Gratings were displayed in an annulus (outer diameter: 15° of visual angle, inner
267 diameter: 1°), surrounding a black fixation bull’s eye (4 cd/m^2), on a gray (580 cd/m^2) background. The
268 visual stimuli were presented with an LCD projector (1024 × 768 resolution, 60 Hz refresh rate)
269 positioned outside the magnetically shielded room, and projected on a translucent screen via two front-
270 silvered mirrors. The projector lag was measured at 36 ms, which was corrected for by shifting the time
271 axis of the data accordingly. The auditory cue consisted of a pure tone (500 or 1000 Hz, 250 ms duration,
272 including 10 ms on and off-ramp time), presented over MEG-compatible earphones.

273

274 **Experimental design.** Each trial consisted of an auditory cue, followed by two consecutive grating stimuli
275 (750 ms SOA between auditory and first visual stimulus) (Figure 1A). The two grating stimuli were
276 presented for 250 ms each, separated by a blank screen (500 ms). A central fixation bull’s eye (0.7°) was
277 presented throughout the trial, as well as during the intertrial interval (ITI, 2250 ms). The auditory cue
278 consisted of either a low- (500 Hz) or high-frequency (1000 Hz) tone, which predicted the orientation of
279 the first grating stimulus (45° or 135°) with 75% validity (Figure 1B). In the other 25% of trials, the first
280 grating had the orthogonal orientation. Thus, the first grating had an orientation of either exactly 45° or
281 135°, and a luminance contrast of 80%. The second grating differed slightly from the first in terms of

282 both orientation and contrast (see below), as well as being in antiphase to the first grating (which had a
283 random spatial phase). The contingencies between the auditory cues and grating orientations were
284 flipped halfway through the experiment (i.e., after four runs), and the order was counterbalanced over
285 subjects.

286 In separate runs (64 trials each, ~4.5 minutes), subjects performed either an orientation or a
287 contrast discrimination task on the two gratings. When performing the orientation task, subjects had to
288 judge whether the second grating was rotated clockwise or anticlockwise with respect to the first
289 grating. In the contrast task, a judgment had to be made on whether the second grating had lower or
290 higher contrast than the first one. These tasks were explicitly designed to avoid a direct relationship
291 between the perceptual expectation and the task response. Subjects indicated their response (response
292 deadline: 750 ms after offset of the second grating) using an MEG-compatible button box. The
293 orientation and contrast differences between the two gratings were determined by an adaptive
294 staircase procedure (Watson and Pelli, 1983), being updated after each trial. This was done to yield
295 comparable task difficulty and performance (~ 75% correct) for the different tasks. Staircase thresholds
296 obtained during one task were used to set the stimulus differences during the other task, in order to
297 make the stimuli as similar as possible in both contexts. As in previous studies using a similar paradigm
298 (Kok et al., 2012, 2016b), there was no interaction between the effects of the expectation cue and the
299 task participants performed, and therefore we collapsed over the two tasks in all MEG analyses.

300 All subjects completed eight runs (four of each task, alternating every two runs, order was
301 counterbalanced over subjects) of the experiment, yielding a total of 512 trials. The staircases were kept
302 running throughout the experiment. Before the first run, as well as in between runs four and five, when
303 the contingencies between cue and stimuli were flipped, subjects performed a short practice run
304 containing 32 trials of both tasks (~4.5 minutes).

305 Interleaved with the main task runs, subjects performed eight runs of a grating localiser task
306 (Figure 1C). Each run (~2 min) consisted of 80 grating presentations (ITI uniformly jittered between 1000
307 and 1200 ms). The grating annuli were identical to those presented during the main task (80% contrast,
308 250 ms duration, 1.0 cycles/°, random spatial phase). Each grating had one of eight orientations

309 (spanning the 180° space, starting at 0°, in steps of 22.5°), each of which was presented ten times per
310 run in pseudorandom order. A black fixation bull's eye (4 cd/m², 0.7° diameter, identical to the one
311 presented during the main task runs) was presented throughout the run. On 10% of trials
312 (counterbalanced across orientations), the black fixation point in the centre of the bull's eye (0.2°, 4
313 cd/m²) briefly turned gray (324 cd/m²) during the first 50 ms of grating presentation. Participants task
314 was to press a button (response deadline: 500 ms) when they perceived this fixation flicker. This simple
315 task was meant to ensure central fixation, while rendering the gratings task-irrelevant. Trials containing
316 fixation flickers were excluded from further analyses.

317 Finally, participants were exposed to a tone localiser (~1.5 min), presented at the start, end, and
318 halfway through the MEG session. These runs consisted of 81 presentations of the two tones used in the
319 main experiment. Data from these runs were not analysed further.

320 Prior to the MEG session (1–3 days), all participants completed a behavioural session. The aim
321 of this session was to familiarise participants with the tasks and to initialise the staircase values for both
322 the orientation and the contrast discrimination task (see above). The behavioural session consisted of
323 written instructions and 32 practice trials of each task, followed by four runs (~4.5 min each) of the main
324 experiment (each task twice, alternating between runs, cue contingencies switching between the
325 second and third run). Finally, participants were exposed to one run each of the grating and tone
326 localiser, to familiarise them with the procedure.

327

328 **MEG recording and preprocessing.** Whole-head neural recordings were obtained using a 275-channel
329 MEG system with axial gradiometers (CTF Systems, Coquitlam, BC, Canada) located in a magnetically
330 shielded room. Throughout the experiment, head position was monitored online, and corrected if
331 necessary, using three fiducial coils that were placed on the nasion and on earplugs in both ears (Stolk et
332 al., 2013). If subjects had moved their head more than 5 mm from the starting position they were
333 repositioned during block breaks. Furthermore, both horizontal and vertical electrooculograms (EOGs),
334 as well as an electrocardiogram (ECG) were recorded to facilitate removal of eye- and heart-related

335 artifacts. The ground electrode was placed at the left mastoid. All signals were sampled at a rate of 1200
336 Hz.

337 The data were preprocessed offline using FieldTrip (Oostenveld et al., 2011)
338 (www.fieldtriptoolbox.org). In order to identify artifacts, the variance (collapsed over channels and time)
339 was calculated for each trial. Trials with large variances were subsequently selected for manual
340 inspection and removed if they contained excessive and irregular artifacts. Independent component
341 analysis was subsequently used to remove regular artifacts, such as heartbeats and eye blinks.
342 Specifically, for each subject, the independent components were correlated to both EOGs and the ECG
343 to identify potentially contaminating components, and these were subsequently inspected manually
344 before removal. For the main analyses, data were low-pass filtered using a two-pass Butterworth filter
345 with a filter order of 6 and a frequency cutoff of 40 Hz. To rule out that the temporal smoothing caused
346 by low-pass filtering may have artificially decreased the onset latency of neural signals, we repeated the
347 decoding analyses (see below) on data that were not low-pass filtered (Figure 3 - figure supplement 1).
348 Here, only notch filters were applied at 50, 100 and 150 Hz to remove line noise and its harmonics.
349 Finally, main task data were baseline corrected on the interval of -250 to 0 ms relative to auditory cue
350 onset, and grating localiser data were baseline corrected on the interval of -200 to 0 ms relative to visual
351 grating onset.

352

353 **Event-related field analysis.** Event-related fields (ERFs) were calculated per participant, and subjected
354 to a planar gradient transformation (Bastiaansen and Knösche, 2000) before averaging across
355 participants. The planar transformation simplifies the interpretation of the sensor-level data because it
356 typically places the maximal signal above the source. To avoid differences in the amount of noise when
357 comparing conditions with different numbers of trials, we matched the trial count by randomly selecting
358 a subsample of trials from the conditions with more trials (i.e., valid expectations).

359

360 **Orientation decoding analysis.** To probe sensory representations in the visual cortex, we used a forward
361 modelling approach to reconstruct the orientation of the grating stimuli from the MEG signal (Brouwer

362 and Heeger, 2009, 2011; Garcia et al., 2013; Myers et al., 2015). The forward modelling approach was
363 two-fold. First, a theoretical forward model was postulated that described the measured activity in the
364 MEG sensors, given the orientation of the presented grating. Second, this forward model was used to
365 obtain an inverse model that specified the transformation from MEG sensor space to orientation space.
366 The forward and inverse models were estimated on the basis of the grating localiser data. The inverse
367 model was then applied to the data from the main experiment, in order to generalise from sensory
368 signals evoked by task-irrelevant gratings to the gratings and expectation signals evoked in the main task.
369 To test the performance of the model we also applied it to the localiser data itself, using a cross-
370 validation approach in which in each iteration one trial of each orientation was used at the test set, and
371 the remaining data were used as the training set.

372 The forward model was based on work by Brouwer and Heeger (Brouwer and Heeger, 2009,
373 2011) and involved 32 hypothetical channels, each with an idealised orientation tuning curve. Each
374 channel consisted of a half-wave-rectified sinusoid raised to the fifth power, and the 32 channels were
375 spaced evenly within the 180° orientation space, such that a tuning curve with any possible orientation
376 preference could be expressed exactly as a weighted sum of the channels. Arranging the hypothesised
377 channel activities for each trial along the columns of a matrix \mathbf{C} (32 channels \times n trials), the observed
378 data could be described by the following linear model:

$$379 \quad \mathbf{B} = \mathbf{WC} + \mathbf{N}$$

380 where \mathbf{B} are the (m sensors \times n trials) MEG data, \mathbf{W} is a weight matrix (m sensors \times 32 channels) that
381 specifies how channel activity is transformed into sensory activity, and \mathbf{N} are the residuals (i.e., noise).

382 In order to obtain the inverse model, we estimated an array of spatial filters that, when applied
383 to the data, aimed to reconstruct the underlying channel activities as accurately as possible. In doing so,
384 we extended Brouwer and Heeger's (Brouwer and Heeger, 2009, 2011) approach in three respects. First,
385 since the MEG signal in (nearby) sensors is correlated, we took into account the correlational structure
386 of the noise. Second, we estimated a spatial filter for each orientation channel independently. As a
387 result, the number of channels used in our model was not constrained, whereas the maximum number

388 of channels would otherwise be dependent on the number of presented orientations. In practice, this
 389 resulted in smoothing in orientation space, because the channels were not truly independent. Third,
 390 each filter was normalised such that the magnitude of its output matched the magnitude of the
 391 underlying channel activity it was designed to recover. Prior to estimating the inverse model, \mathbf{B} and \mathbf{C}
 392 were demeaned such that their average over trials equalled zero, for each sensor and channel,
 393 respectively.

394 As stated above, the inverse model was estimated on the basis of the grating localiser data. On
 395 each localiser trial, one of eight orientations was presented (see above), and the hypothetical responses
 396 of each of the channels could thus be calculated for each trial, resulting in the response column vector
 397 $\mathbf{c}_{train,i}$, of length n_{train} trials, for each channel i . The weights on the sensors \mathbf{w}_i could now be obtained
 398 through least squares estimation, for each channel:

$$399 \quad \mathbf{w}_i = \mathbf{B}_{train} \mathbf{c}_{train,i}^T \left(\mathbf{c}_{train,i} \mathbf{c}_{train,i}^T \right)^{-1}$$

400 where \mathbf{B}_{train} are the (m sensors \times n_{train} trials) localiser MEG data. Subsequently, the optimal spatial
 401 filter \mathbf{v}_i to recover the activity of the i -th channel was obtained as follows (Mostert et al., 2015):

$$402 \quad \mathbf{v}_i = \frac{\tilde{\Sigma}_i^{-1} \mathbf{w}_i}{\mathbf{w}_i^T \tilde{\Sigma}_i^{-1} \mathbf{w}_i}$$

403 where $\tilde{\Sigma}_i$ is the regularised covariance matrix for channel i . Incorporating the noise covariance in the
 404 filter estimation leads to the suppression of noise that arises from correlations between sensors. The
 405 noise covariance was estimated as follows:

$$406 \quad \hat{\Sigma}_i = \frac{1}{n_{train} - 1} \boldsymbol{\varepsilon}_i \boldsymbol{\varepsilon}_i^T$$

$$407 \quad \boldsymbol{\varepsilon}_i = \mathbf{B}_{train} - \mathbf{w}_i \mathbf{c}_{train,i}^T$$

408 where n_{train} is the number of training trials. For optimal noise suppression, we improved this estimation
 409 by means of regularization by shrinkage, using the analytically determined optimal shrinkage parameter
 410 (for details, see (Blankertz et al., 2011)), yielding the regularised covariance matrix $\tilde{\Sigma}_i$.

411 Such a spatial filter was estimated for each hypothetical channel, yielding an m sensors \times 32
 412 channel filter matrix \mathbf{V} . Given that we performed our decoding analysis in a time-resolved manner, \mathbf{V}

413 was estimated at each time point of the training data, in steps of 5 ms, resulting in array of filter
414 matrices, or decoders. To improve the signal-to-noise ratio, the data were first averaged within a
415 window of 29.2 ms centred on the time point of interest. The window length of 29.2 ms was based on
416 an a priori chosen length of 30 ms, but minus one sample such that the window contained an odd
417 number of samples for symmetric centring (Mostert et al., 2015). These filter matrices could now be
418 applied to estimate the orientation channel responses in independent data – in this case, the trials from
419 the main experiment:

$$\mathbf{C}_{test} = \mathbf{V}^T \mathbf{B}_{test}$$

420
421 where \mathbf{B}_{test} are the (m sensors \times n_{test} trials) main experiment data. These channel responses were
422 estimated at each time point of the test data, in steps of 5 ms, with the data being averaged within a
423 window of 29.2 ms at each step. This procedure resulted in a four-dimensional (training time \times testing
424 time \times 32 channel \times n_{test}) matrix of estimated channel responses for each trial in the main experiment.
425 Each trials' channel responses were shifted such that the channel with its hypothetical peak response at
426 the orientation presented on that trial (i.e. 45° or 135°) ended up in the position of the 0° channel,
427 before averaging over trials within each condition (i.e., valid vs. invalid expectation). Thus, the presented
428 orientation was defined as 0°, by convention. Note that for 3D surface plots that show the evolution of
429 channel responses over time (e.g., Figure 2B), the response of the 90° channel (i.e., orthogonal to the
430 presented orientation) was used as a baseline, to avoid negative numbers for visualisation purposes.

431 To quantify decoding performance, the channel responses for a given condition were converted
432 into polar form and projected onto a vector with angle 0° (the presented orientation, see above).

$$r = |z| \cos(\arg(z)), \quad z = \mathbf{c}^T e^{2i\phi}$$

433
434 where \mathbf{c} is a vector of estimated channel responses, and ϕ is the vector of angles at which the channels
435 peak (multiplied by 2 to project the 180° orientation space onto the full 360° space). The scalar
436 projection r indicates the strength of the decoder signal for the orientation presented on screen. (Note
437 that subtracting the estimated response of the 90° channel from that of the 0° channel yielded virtually

438 identical results, data not shown.) This quantification yielded (training time \times testing time) temporal
439 generalisation matrices of orientation decoding performance.

440 In order to isolate any orientation-specific neural signals evoked by the expectation cues, we
441 applied the following subtraction logic. On valid expectation trials, the expected and presented
442 orientations are identical, and thus the orientation signal induced by both the cue and stimulus be
443 expected to be positive, by convention. On invalid expectation trials on the other hand, the expected
444 and presented orientations are orthogonal, and thus the orientation signal induced by the stimulus
445 would be positive and the signal induced by cue would be expected to be negative. Thus, subtracting the
446 orientation decoding signal on invalid trials from that on valid trials would subtract out the stimulus-
447 evoked signal while revealing any cue-induced orientation signal.

448

449 **Statistical testing.** Neural signals evoked by the different conditions were statistically tested using
450 nonparametric cluster-based permutation tests (Maris and Oostenveld, 2007). For ERF analyses, we
451 averaged over the spatial (sensor) dimension, on the basis of independent localisation of the 10 sensors
452 that showed the strongest visual-evoked activity during the grating localiser between 50 and 150 ms
453 post-stimulus. Therefore, our statistical analysis considered one-dimensional (temporal) clusters. For
454 orientation decoding analyses, the data consisted of two-dimensional (training time \times testing time)
455 decoding performance matrices, and the statistical analysis thus considered two-dimensional clusters.
456 For both one- and two-dimensional data, univariate t -statistics were calculated for the entire matrix and
457 neighbouring elements that passed a threshold value corresponding to a p -value of 0.01 (two-tailed)
458 were collected into separate negative and positive clusters. Elements were considered neighbours if
459 they were directly adjacent, either cardinally or diagonally. Cluster-level test statistics consisted of the
460 sum of t -values within each cluster, and these were compared to a null distribution of test statistics
461 created by drawing 10,000 random permutations of the observed data. A cluster was considered
462 significant when its p -value was below 0.05 (two-tailed).

463 **Figure supplements**

464

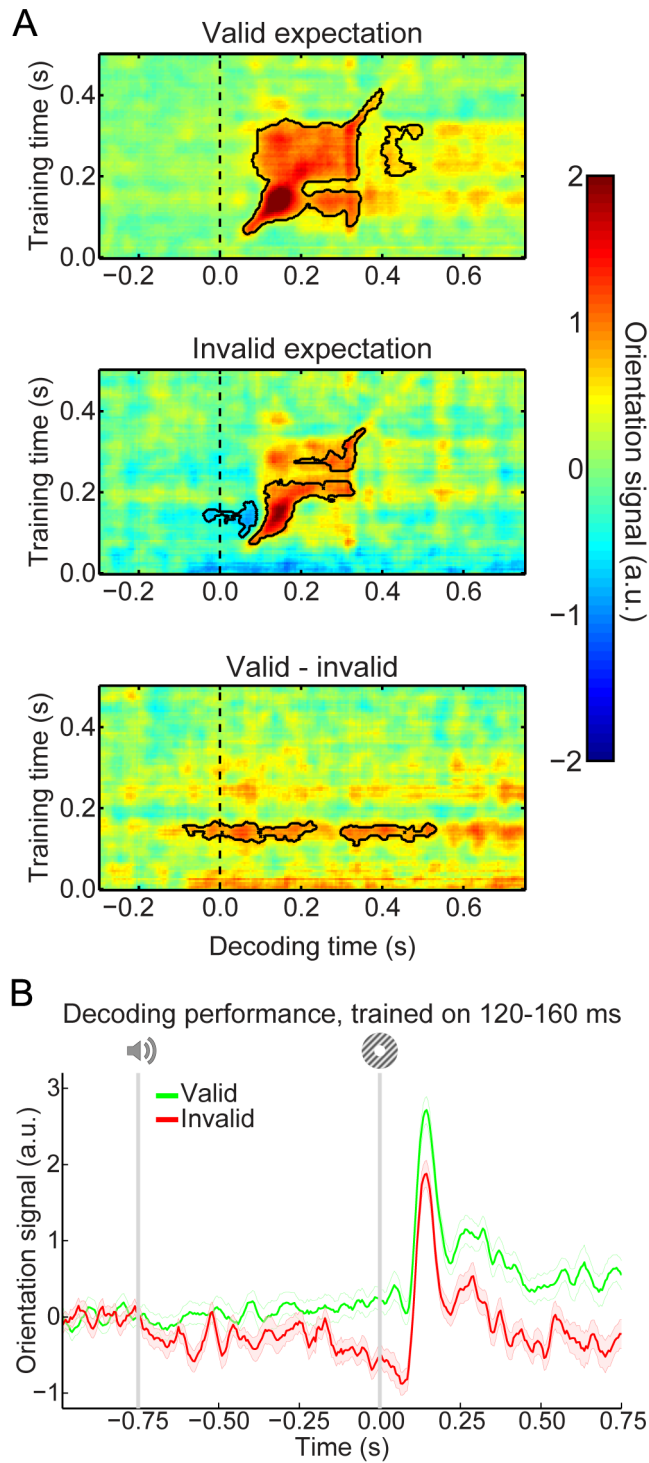
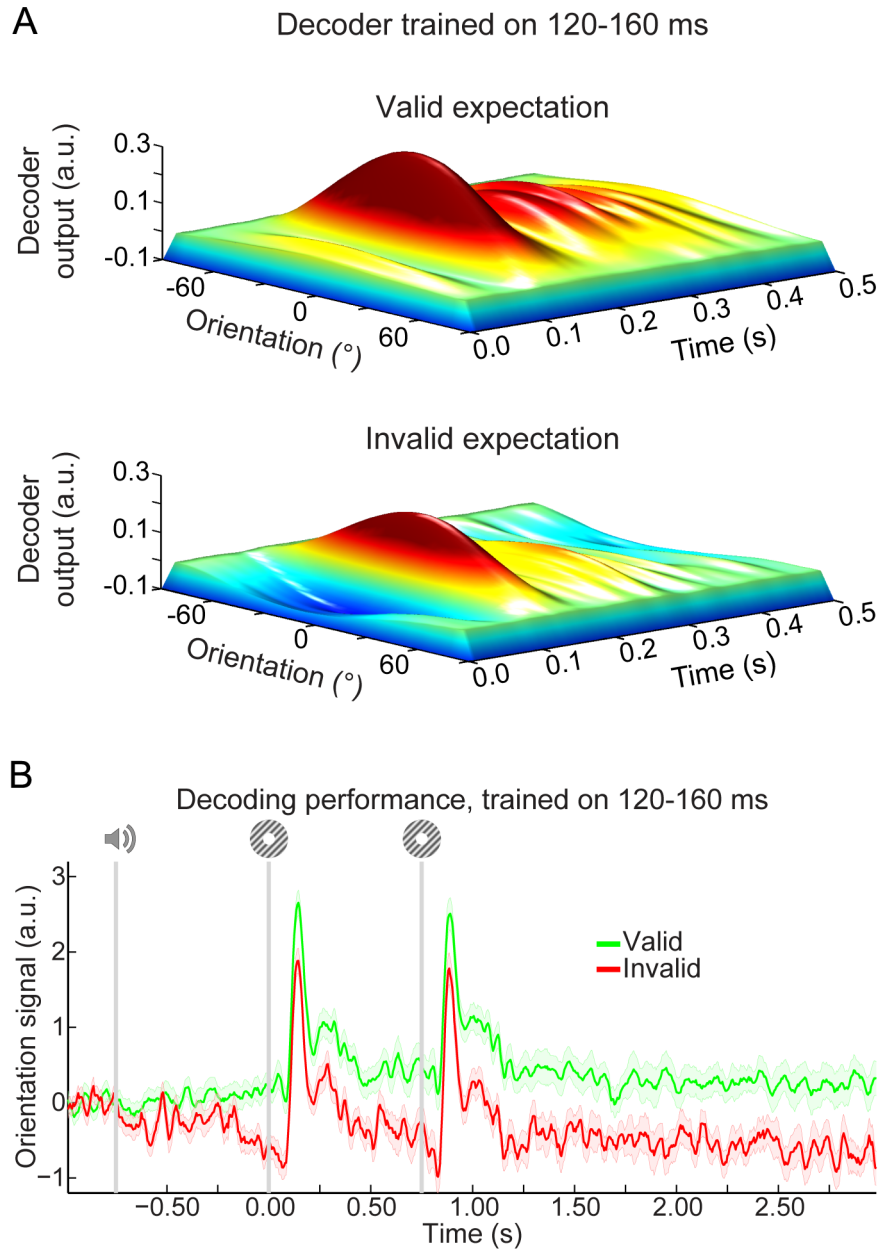


Figure 3 – figure supplement 1. Expectation effects in the absence of low-pass filtering. (A)

Temporal generalisation matrices of orientation decoding during the main experiment, without low-pass filtering the data. Otherwise, identical to Figure 3A. Solid black lines indicate significant clusters ($p < 0.05$). **(B)** Orientation decoding during the main task, averaged over training time 120 – 160 ms post-stimulus during the grating localiser. Shaded regions indicate SEM.



477

478 **Figure 3 – figure supplement 2. Expectation affects orientation decoding throughout the trial. (A)**

479 Output of the 32 orientation channels over time, during the main task, separately for gratings preceded

480 by a valid (top panel) and invalid (bottom panel) expectation cue. Here, decoders were trained on 120 –

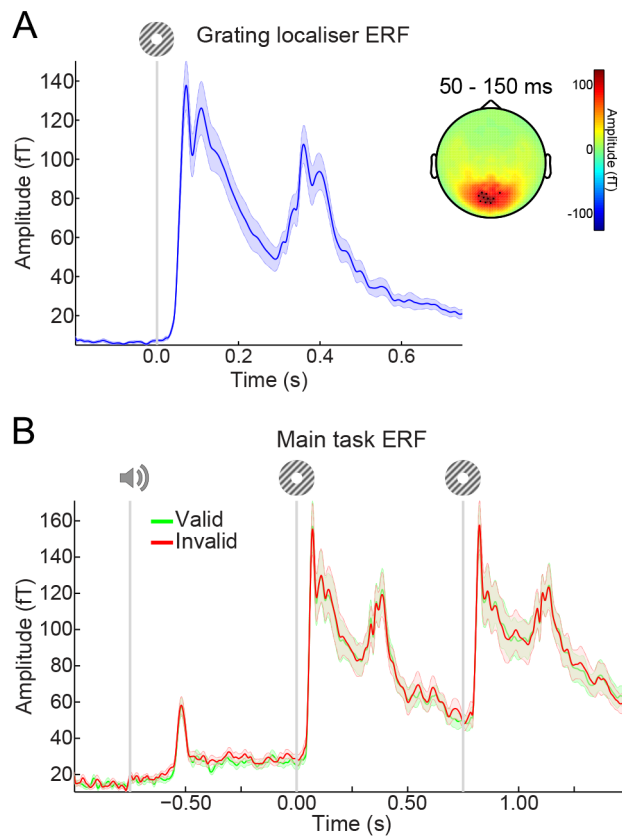
481 160 ms post-stimulus during the grating localiser, similar to the data in Figure 3B. That is, this figure

482 depicts similar results as in Figure 3B, but displaying the output of all 32 orientation channels, rather

483 than collapsing the channel responses into a decoding performance score (cf. Figure 2B). **(B)** Orientation

484 decoding during the main task, averaged over training time 120 – 160 ms post-stimulus during the

485 grating localiser. Same as in Figure 3 – figure supplement 1B, but with an extended x-axis, in order to
486 shown the sustained nature of the expectation templates. Shaded regions indicate SEM.
487
488
489
490



491
492 **Figure 3 – figure supplement 3. Event related fields. (A)** Event-related fields during the grating localiser,
493 in the 10 channels showing the largest response from 50 – 150 ms post-stimulus. **(B)** Event-related fields
494 during the main task, in the same 10 channels as in panel A, selected on the basis of the grating localiser
495 response. No significant differences between ERF for validly and invalidly predicted gratings. Shaded
496 regions indicate SEM.

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