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Article type : Special Issue Article

Reviewer 1: Marzia Del Zotto (University Hospitals of Geneva, Switzerland) Reviewer 2: Ryszard Aukstulewicz (Humboldt-Universität zu Berlin, Germany)

**Title:**

**Timing of repetition suppression of event-related potentials to unattended objects**

**Running title:**

**Time-course of repetition suppression of ERPs**

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**Keywords**

Event-related potential (ERP); general linear modelling (GLM); object recognition; predictive coding (PC); repetition enhancement (RE)

This article has been accepted for publication and undergone full peer review but has not been through the copyediting, typesetting, pagination and proofreading process, which may lead to differences between this version and the Version of Record. Please cite this article as doi: 10.1111/ejn.13972

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## Abstract

Current theories of object perception emphasize the automatic nature of perceptual inference. Repetition suppression (RS), the successive decrease of brain responses to repeated stimuli, is thought to reflect the optimization of perceptual inference through neural plasticity. While functional imaging studies revealed brain regions that show suppressed responses to the repeated presentation of an object, little is known about the intra-trial time course of repetition effects to everyday objects. Here we used event-related potentials (ERP) to task-irrelevant line-drawn objects, while participants engaged in a distractor task. We quantified changes in ERPs over repetitions using three general linear models (GLM) that modelled RS by an exponential, linear, or categorical “change detection” function in each subject. Our aim was to select the model with highest evidence and determine the within-trial time-course and scalp distribution of repetition effects using that model. Model comparison revealed the superiority of the exponential model indicating that repetition effects are observable for trials beyond the first repetition. Model parameter estimates revealed a sequence of RS effects in three time windows (86-140ms, 322-360ms, and 400-446ms) and with occipital, temporo-parietal, and fronto-temporal distribution, respectively. An interval of repetition enhancement (RE) was also observed (320-340ms) over occipito-temporal sensors. Our results show that automatic processing of task-irrelevant objects involves multiple intervals of RS with distinct scalp topographies. These sequential intervals of RS and RE might reflect the short-term plasticity required for optimization of perceptual inference and the associated changes in prediction errors (PE) and predictions, respectively, over stimulus repetitions during automatic object processing.

## Introduction

Stimulus repetition-related phenomena are ubiquitous in psychophysics, psychology, and neuroscience (Henson, 2003; Ibbotson, 2005; Grill-Spector et al., 2006; Krekelberg et al., 2006; Clifford et al., 2007; Kohn, 2007; Auksztulewicz & Friston, 2016; Barron et al., 2016). Functional neuroimaging studies have observed both repetition suppression (RS) and repetition enhancement (RE) effects (reviewed in (Segaert et al., 2013)). RS, also referred to as stimulus specific adaptation (SSA), is thought to reflect a rapid form of experience-dependent plasticity affecting perception and response properties of neurons. It has also been linked to optimization of the brain’s predictions about the sensory environment (Solomon & Kohn, 2014) by discounting expected properties (Clifford et al., 2007; Summerfield et al., 2008; Webster, 2011; Vogels, 2016).

Repetition-related phenomena in the visual system are complex and likely represent a compound of distinct neural processes, the mechanisms of which are not fully understood yet (Ibbotson, 2005; Grill-Spector et al., 2006; Solomon & Kohn, 2014). At least three mechanisms contribute to SSA, including i) somatic afterhyperpolarization, ii) synaptic depression due to depletion of presynaptic vesicles, and iii) circuit-level mechanisms (Kohn, 2007; von der Behrens et al., 2009). Adaptation is thought to modify neural population coordination and its effects cascade through the processing stages, possibly affecting multiple networks (Solomon & Kohn, 2014).

Predictive coding (PC) offers a neurobiologically plausible, mechanistic model for stimulus repetition related phenomena, and accommodates observations in psychophysics, electrophysiology and functional neuroimaging (Friston, 2005; Auksztulewicz & Friston, 2016). PC suggests that the brain

maintains and updates an internal model of the environment to infer the most likely causes of sensory inputs and actively generates predictions (Clark, 2015). In this framework, sensory systems are hierarchically organized where each level receives inputs from the level below that signals a mismatch between predicted and observed events, a prediction error (PE; Hinton et al., 1995; Rao & Ballard, 1999; Friston, 2005; Hohwy, 2013). In turn, each level sends its input to the level below conveying predictions that are thought to explain away PEs at the lower level. Perceptual inference, i.e. the process of determining the most likely cause of sensory inputs, thus rests on message passing across the hierarchy. From the perspective of PC, adaptation or RS can be understood as a neural correlate of perceptual learning, i.e. the optimized process of perceptual inference where PEs to repeated stimuli are explained away more efficiently due to synaptic plasticity in sensory circuits (Baldeweg, 2007; Garrido et al., 2009). Conversely, RE could represent increased neural activity during the “sharpening” of predictions corresponding to increasing precision (implemented by increased postsynaptic gain) over repetitions. Encoding precision is important since veridical perception not only rests on the content of sensory signals, but also on the confidence or precision of the signals that drive inference. It has been suggested that the brain might employ mechanisms to encode precision by relying on modulatory neurotransmitters that regulate the gain or excitability of populations (Friston et al., 2016). Specifically, cholinergic mechanisms might affect the encoding of sensory precision by modulating postsynaptic gain (Feldman and Friston, 2010; Moran et al., 2013). Alternatively, cholinergic mechanisms can also affect the function of N-methyl-D-aspartate (NMDA) receptors (Aramakis et al., 1997; Chen et al., 2008) which, in turn, are thought to be involved in signaling both prediction errors and predictions in cortical hierarchies (Friston 2005; Corlett et al. 2011; Stephan et al. 2016). In sum, PC offers potential explanations for the phenomena observed during repeated stimulus presentation: larger responses to unpredicted events, and attenuation as well as enhancement to repeated events (Egner et al., 2010; de Gardelle et al., 2013; Recasens et al., 2015).

While studies on repetition of faces and words are relatively abundant (for a recent review, see Schweinberger & Neumann, 2016), there are few electroencephalography (EEG) studies where time-course and topographic distribution of repetition effects to objects were investigated at the whole-scalp level. Typically, most studies focused the analysis on a preselected set of electrodes and confined amplitude measurements to time windows either based on visual inspection of the data or on previous reports in the literature. Furthermore, previous studies often investigated brain responses only to the first repetition relative to the initial presentation, thus ignoring the dynamics of brain responses to further repetitions (Schendan & Kutas, 2003; Henson et al., 2004; Eddy et al., 2006; Gruber et al., 2006; Gruber & Muller, 2006; Guillaume et al., 2009; Gilbert et al., 2010; Kim et al., 2012; Andrade et al., 2015; Gosling et al., 2016). The aim of the current study was to determine the time course and scalp distribution of repetition effects without prior hypotheses about the dynamics (RS vs RE), time course, or scalp distribution of repetition effects. To this end, we analysed the spatio-temporal dynamics of ERPs to black and white line drawings of common objects over six consecutive presentations. We used statistical parametric mapping (SPM; Friston, 2007) to analyse ERP amplitudes at each and every sensor in the poststimulus 50-500ms time window using a mass-univariate approach. Given that most studies focused on the analysis of ERPs to the first repetition only, little is known about the time course of the decay of scalp-recorded ERPs to object stimuli that are repeated multiple times. Therefore we set up three GLMs with parametric regressors incorporating three hypotheses about the time-course of repetition effects. We used an exponential,

a linear, and a “change detection” model to identify ERP components that showed a reliable repetition effect and performed Bayesian model comparison (Stephan et al., 2009) in order to identify the model that best explained the observed data.

## Methods

### Participants and ethics statement

Seventeen students (mean age 21.06 ys, SD 1.56 ys, 8 female, 15 right-handed) volunteered to participate in the study. The experimental protocols were approved by the Institutional Review Board of the Institute for Psychology, Hungarian Academy of Sciences. All participants gave their written informed consent after the nature of the experiment had been fully explained. They received a monetary compensation for their participation in the study. All participants had normal or corrected-to-normal vision. The experiments were conducted in compliance with the Declaration of Helsinki.

### Stimuli and procedure

In each of the four blocks, we recorded event-related potentials (ERP) to 60 black and white line drawings of common objects taken from the picture inventory by Szekely et al. (2004). The 60 object pictures were selected from the following semantic categories: small artifacts (n=39, e.g., book, flag); large artifacts (n=1, bed); objects found in nature (n=2, flower, leaf); things to wear (n=5, e.g., coat, hat); body parts (n=2, feather, heart); foods (n=11, e.g., apple, mushroom). Stimuli were organized into microsequences of 6-10 presentations of an object followed by a close-up or wider-angle view of the same object, and an additional repetition of the same object with its original viewing angle another two times (Figure 1A). ERPs to changes of viewing angle were not analysed here. They were included to study boundary extension effects (e.g., Czigler et al., 2013) and will be published elsewhere. Thus the length of microsequences varied pseudo-randomly between 9-13 presentations. To study repetition effects here we analysed ERPs to the first six stimuli, exclusively. In each block we used 60 individual black line-drawn objects on white background. Stimulus duration was 250ms with 320ms inter-stimulus interval (ISI). Each picture subtended 11.4° visual angle and was presented on a dark grey background. A black fixation cross was presented in the centre of the screen laid over a grey disk subtending 1.17° visual angle. To minimize eye-movements subjects were instructed to fixate at the cross throughout the experiment. Similar to our prior studies (Stefanics et al., 2011; Stefanics & Czigler, 2012; Csukly et al., 2013; Kovacs-Balint et al., 2014; Farkas et al., 2015) we employed a behavioural task to minimize the variation of attentional effects on the processing of object stimuli across participants by engaging the participants' attention. Pseudo-randomly every 3-6 seconds, the fixation cross became wider or longer (Fig. 1A). The participants' task was a speeded button-press to the changes of the cross and reaction time (RT) was recorded. Trials occurring within an 800 ms interval after a change in the fixation cross were excluded from the analysis. RTs and hit rates were compared between experimental blocks with analyses of variance (ANOVA). Stimuli were presented using Cogent 2000 and Cogent Graphics developed at the Wellcome Department of Imaging Neuroscience (<http://www.vislab.ucl.ac.uk/Cogent/index.html>). Participants sat in a comfortable chair in a sound-attenuated, dimly lit room during EEG recording.

## EEG recordings and preprocessing

EEG was recorded from 61 Ag/AgCl electrodes according to a modified international 10–20 system (AF7, Fp1, Fpz, Fp2, AF8, AF3, AFz, AF4, F7, F5, F3, F1, Fz, F2, F4, F6, F8, FT7, FC5, FC3, FC1, FCz, FC2, FC4, FC6, FT8, T7, C5, C3, C1, Cz, C2, C4, C6, T8, TP7, CP5, CP3, CP1, CPz, CP2, CP4, CP6, TP8, P7, P5, P3, P1, Pz, P2, P4, P6, P8, PO7, PO3, POz, PO4, PO8, O1, Oz, O2). An electrode attached to the tip of the nose was used as reference. The ground electrode was placed on the forehead. EEG was recorded from DC with a low-pass filter at 100 Hz (Neuroscan Synamp, Victoria, Australia). Eye movements were monitored by two horizontal and two vertical bipolar EOG electrodes. Data were digitized at 32 bit resolution and a sampling rate of 500 Hz. EEG was filtered off-line between 1 and 30 Hz (24 dB/octave) and re-referenced to the common average.

Epochs extending -100 ms before to 550 ms after stimulus onset were extracted from the continuous EEG for each object for the first six presentations. Epochs were baseline corrected to the pre-stimulus 100 ms period. To avoid other potential artefacts, epochs with values exceeding  $\pm 75 \mu\text{V}$  on any EEG or EOG channel were rejected from the analysis using the open source software EEGLAB (RRID: SCR\_007292, (Delorme and Makeig, 2004)) in the Matlab development environment (MathWorks, Natick, USA). After artifact rejection, the total number of trials (summed over the four blocks) used for calculating the mean ERPs that entered the GLM was 197 (sd=11), 184 (sd=13), 183 (sd=11), 182 (sd=12), 187 (sd=11), 178 (sd=11), for 1<sup>st</sup>, 2<sup>nd</sup>, 3<sup>rd</sup>, 4<sup>th</sup>, 5<sup>th</sup>, and 6<sup>th</sup> presentation of stimuli, respectively.

## Space $\times$ time SPM analysis

Mean ERP data were converted to scalp  $\times$  time images for statistical analysis using the open source software SPM12 (v6470, RRID: SCR\_007037; Litvak et al., 2011), following similar preprocessing and statistical procedures as in previous work (e.g., Henson et al., 2008; Garrido et al., 2013; Auztulewicz & Friston, 2015; Stefanics et al., 2018). The data were interpolated to create a 32 $\times$ 32 pixel scalp map for each time-point in the poststimulus 50-500 ms interval. Given a sampling frequency of 500 Hz, the time dimension consisted of 226 samples in each averaged ERP. Images were stacked to create a 3D space-time image volume that was smoothed with a Gaussian kernel (full-width at half-maximum (FWHM)=[16mm 16mm 16ms]) in accordance with the assumptions of Random Field Theory (Worsley & Friston, 1995; Kiebel & Friston, 2004).

A GLM with four main regressors corresponding to the four experimental blocks, and a generic decay/rise function as a parametric modulator for each stimulus presentation in each block, was estimated for each participant. The design matrix for a single subject is depicted in Fig. 1B. The decay/rise function was used to quantify changes in ERP amplitude over repetitions within microsequences as unpredicted firstly-presented objects became more predictable over consecutive repetitions. We chose an exponential function as parametric modulator since several electrophysiological and neuroimaging findings indicating that response attenuation over repeated stimulus presentations typically follows a non-linear decay that is well approximated by an exponential function (e.g., Puce et al., 1999; Sanchez-Vives et al., 2000; Baldeweg, 2006; Boehnke et al., 2011; de Gardelle et al., 2013; Kaliukhovich & Vogels, 2014). This allowed us to estimate modulation of the repetition sequence with an exponential time course. Importantly this design is flexible enough to capture both decay (attenuation or suppression) and rise (facilitation or enhancement) processes. For example, a decrease of a positive ERP component would show a

positive correlation with our regressor across repetitions, whereas an increase of a positive component would show a negative correlation with our regressor (conversely for negative components). The estimated regression coefficients (beta parameter estimates) of the decay function for each scalp time-point for each participant were analysed at the group level for all three models (exponential, linear, “change detection”), using a standard two-stage summary statistics approach (Friston et al. 2005; Mumford & Nichols 2009). These parameter estimates represent the relationship (similarity) between the dynamics of ERPs over repeated stimulus presentations and the parametric decay regressors.

On the group level, we used F-tests to find scalp time-points where mean ERPs were significantly modulated by repeated stimulus presentations. The resulting statistical parametric maps (SPM) were family-wise error (FWE) corrected for multiple comparisons at the voxel level ( $p < 0.05$  (FWE)) using Random Field Theory (Flandin & Friston, 2017).

#### Model comparison

Beside using an exponential function as parametric modulator we explored two alternative models previously considered for studying repetition effects (Noppeney and Penny, 2006; Lieder et al., 2013) and compared them with the exponential model. In particular, we considered repetition effects as 1) categorical decay (i.e., object 1st presentation > presentations 2–6) and 2) linear decrease. The categorical decay corresponds to a “change detection” model, i.e., it represents the hypothesis that whenever a new object is presented in a sequence, it elicits a phasic response which disappears completely under the following repetitions. While the linear model is not realistic (given that repetition suppression effects must become consecutively smaller in physiological systems with decay mechanisms), we included this regressor as a “null” model, similar to Noppeney and Penny (2006). Thus, we set up another two GLMs, incorporating these two hypotheses. The design matrices for a single subject are depicted in Fig. 1C and D, respectively.

In order to compare the models formally, we used the Bayesian Information Criterion (BIC) (Schwarz, 1978) approximation to the log model evidence (LME). Under Gaussian noise (as assumed by the GLM), this leads to an approximation of LME that is a function of the residual sum of squares (RSS):

$$LME \approx -\frac{1}{2}n \ln\left(\frac{RSS}{n}\right) - \frac{1}{2}k \ln(n) \quad (1)$$

where  $n$  is the number of data points and  $k$  is the number of parameters estimated by the model.

We first computed the LME for each voxel in individual participants. In order to perform model comparison at the group level, we computed the sum of  $\Delta LME$  (between models) across subjects for each voxel. This is equal to the logarithm of the group Bayes factor (GBF; Stephan et al., 2007) and corresponds to a fixed effects group-level Bayesian model selection (BMS; Stephan et al., 2009) procedure. Group model comparison was done both within a functionally defined mask (of voxels showing repetition effects under all models) as well as on all voxels in the 3D space-time image volume (to perform an unconstrained comparison). The mask comprised all voxels from the SPM analyses where all three models had yielded a significant whole-brain corrected effect (logical “AND”



conjunction). We then used a non-parametric Wilcoxon signed rank test to assess the null hypothesis of zero median for  $\Delta$ LME across all voxels.

## Results

### Reaction time and hit rate

We compared reaction times and hit rates for the occasional changes in the fixation cross between experimental blocks. An ANOVA of reaction times across the four blocks yielded no significant effect ( $F(3,48)=1.12$ ,  $p=0.35$ ) indicating lack of evidence for a change in reaction speed across the blocks. The mean reaction time was 452ms ( $SD=87$ ms). An ANOVA of hit rate across the four blocks yielded no significant effect ( $F(3,48)=0.396$ ,  $p=0.76$ ) indicating a high detection performance without signs of a change in hit rate during the experiment. The mean hit rate was 95.26% ( $SD=6.7\%$ ).

### Model comparison

We assessed the three models by performing model comparison at the group level as described above. The functionally defined mask was a conjoint mask of voxels showing significant repetition effects under any of the three models (logical “AND” conjunction). Fixed-effects Bayesian model comparison revealed that the Exponential model was clearly superior compared to the other two models, and that the Change detection model performed better than the Linear model, both for voxels within the functional mask (Fig. 2A), as well as for voxels within the whole volume (Fig. 2B).

To characterize the distribution of  $\Delta$ LME values more formally, we performed null hypothesis testing, as described above. The results are summarized in Table 1.

### ERP results

Given that model comparison showed that the GLM with the exponential decay function explains the data best, we used the space x time clusters with significant results for the winning model to illustrate repetition effects. We start from scalp topographies and then show using time-windowed data and conventional ERP plots of the effects that lead to significant results in SPM. Scalp topographies of the SPMs are shown in Figure 3A for the cluster maxima. Model estimates to line drawings showed a sequence of RS effects. Over repeated presentations of our visual stimuli, brain responses showed a sequence of RS and RE effects. An early RS effect was observed at bilateral occipital areas in the 86-140ms interval followed by RE at midline occipital sites (320-340ms). Temporo-parietal electrodes over the right hemisphere showed RS in the 322-360ms time window followed by a further interval of RS in the 400-446ms time window at right fronto-temporal sites. Details of test statistics are given in Table 2. A video animation of the time-course of repetition effects over the whole scalp is available in the Supporting information of the online version of this paper.

For visual comparison, ERP amplitudes for each subject are shown in Figure 3B for each of the six stimulus presentations at selected electrodes in the intervals where significant repetition effects were observed. The early RS effect over occipital areas suggests an abrupt suppression of PE responses from the first to the second stimulus presentation, with constant response amplitude for further repetitions. In turn, the subsequent RS effects over temporal and frontal areas show a more smoothly decaying response profile. The occipital RE effect also shows a smooth time course of incrementally increasing response amplitude to repeated stimulus presentations.

For visualisation, grand-average ERP waveforms to the first and sixth stimulus presentations are shown in Figure 2C at electrode sites where SPM analysis had shown significant repetition effects. Our visual stimuli (line drawings of objects) evoked the canonical posterior P1 and P2 components, as well as prominent negative-going peaks at parieto-temporal and frontal electrodes. Note that ERPs are shown for illustration purpose only and no further null hypothesis tests are conducted; all statistical tests were carried out as part of the SPM analysis covering the entire space-time volume, comprising data from all electrodes in the poststimulus 50-500 ms interval (see Methods for details).

## Discussion

In this study we used Statistical Parametric Mapping, a comprehensive analysis framework to investigate effects of stimulus repetition on ERPs to line drawings of everyday objects. For each subject we estimated a GLM including a decay function to quantify changes in brain potentials over the whole scalp in the poststimulus 50-500ms time window. We observed three consecutive intervals of RS in the 86-140ms, 322-360ms, and 400-446ms time window with occipital, temporo-parietal, and fronto-temporal distribution (Figure 3A), respectively. Furthermore, we found an interval of RE in the 320-340ms window with occipito-temporal distribution.

Importantly, similar to previous studies (e.g., Stefanics et al., 2011, 2018; Stefanics & Czigler, 2012; Csukly et al., 2013; Kovacs-Balint et al., 2014; Farkas et al., 2015), our experimental design implemented a primary task to control attentional effects that might modulate repetition-related neural activity (Vuilleumier et al., 2005; Egner et al., 2010; Chennu et al., 2013, 2016; Aukstulewicz & Friston, 2016; Gosling et al., 2016). The constantly high hit-rate and fast RT over the experiment indicated that participants complied with the task and attended the fixation cross. This suggests that repetition effects observed in our study likely reflect automatic perceptual inference operating outside the focus of visual attention. Theories of perception as unconscious inference originate from Helmholtz's classical idea that perceptual experience is the "conclusion" of unconscious inductive inference from sensory input (Hatfield, 2002; Kiefer, 2017). In current theories of cortical information processing such as predictive coding (Friston, 2005), RS is viewed as the result of a process during which the brain minimizes the prediction error (the difference between the predicted and the actual input) with increasing efficacy (due to updating of predictions and the associated synaptic plasticity of cortical connections) during repeated presentations of the same stimulus type. This decrease of prediction errors during processing of a given stimulus, and the change in efficacy of stimulus processing via plasticity in underlying neural circuits corresponds to perceptual inference and learning, respectively (Baldeweg, 2007).



Our findings represent an important advance in the understanding of the time course and scalp distribution of repetition effects in ERP correlates of object perception as they are based on an unbiased mass-univariate statistical approach that covers the entire space×time volume of EEG signals. Instead of relying on visual inspection and preselection of a subset of channels for amplitude tests, this approach included data from all electrodes and all time points in the post-stimulus 50-500ms interval. The analysis takes into account the smoothness of scalp potentials (correlations between neighbouring channels and temporal smoothness of the EEG signal) and used a stringent method for correcting for multiple comparisons (FWE) using Random Field Theory.

RS has been widely used in fMRI studies that examine how specific representations may be encoded by neuronal population activity (e.g., Barron et al., 2016). Furthermore, models of network mechanisms have been put forward to explain repetition suppression at the level of population dynamics (Grill-Spector et al., 2006). In electrophysiology, RS effects have been studied frequently over trials; by contrast, their temporal evolution within trials has received little attention. A predictive coding view suggests that RS reflects the minimisation of prediction error by updating predictions about the content and precision of sensory inputs (Auksztulewicz & Friston, 2015). Importantly, predictive coding also suggests that for non-trivial stimuli, which are processed at several (spatial, temporal, or semantic) scales, multiple significant within-trial intervals of RS should occur. This is because model updating and the “explaining away” of PEs during perceptual inference occurs at several levels of the cortical hierarchy, with temporal delays inherent to the recurrent message passing between areas, and increasing synaptic efficacy due to short-term plasticity is found on all levels (Friston, 2005; Baldeweg, 2006; Baldeweg, 2007; Garrido et al., 2009). Our findings suggest that, for the particular stimuli used here (line drawings of everyday objects) this implicit perceptual process might take place in neural circuitry comprising occipital, temporal, and frontal areas.

Attention is known to counteract repetition suppression, i.e., it increases neural response to the repeated stimuli (e.g., Kok et al., 2012; Auksztulewicz & Friston, 2015). However, in our experiment the object stimuli were task-irrelevant and our protocol employed a primary task to engage the participants’ attention. Therefore attentional effects are unlikely to account for the enhanced responses we observed in the 320-340ms window with occipito-temporal distribution. Repetition enhancement has received less consideration than RS thus far (Segaert et al., 2013). RE effects in studies with implicit tasks have been suggested to reflect perceptual identification (Schnyer et al., 2002) and involvement of explicit memory processes (Segaert et al., 2013). The predictive coding theory assumes that activity of a subset of neural elements that participate in perceptual inference represent predictions about the content and precision of sensory inputs. During stimulus repetition, neural activity underlying both kinds of predictions might increase, and manifest as enhanced neural responses. A prior study using fMRI reported that RS and RE co-occurred in a single cortical region during stimulus repetition (de Gardelle et al., 2013). While our present analyses do not link our scalp ERP results to the source level, our findings of an early interval of RS in the 86-140ms time window followed by an interval of RE in the 320-340ms window with posterior distributions suggest occipito-temporal generator sources. Repetition paradigms with masked object (Eddy et al., 2006) and face (Henson et al., 2008) stimuli, as well an auditory study by Recasens et al. (2015), have also reported distinct intervals of RS and RE effects, indicating that not only unattended but even subliminal processing of faces and objects, as well as pure tones, are associated with both increase and

decrease of ERPs. It is notable that the same pattern of results have been obtained by Henson et al. (2008), Recasens et al. (2015) and our current study, namely that the RE effects occur later than RS effects.

A limitation of our study is the relatively short stimulus onset asynchrony (SOA) of 570ms. Due to the short SOA, we cannot fully exclude that the pre-stimulus 100ms baseline period contained ERP components from the previous trial. However, it is unlikely that baseline correction (which consists of subtracting a constant) could have significantly contributed to the observed sequence of multiple short intervals of repetition effects.

Scalp-recorded ERPs result from the linear summation of electric fields generated in the brain, therefore it is non-trivial to determine whether changes in a certain ERP component relative to a baseline period were caused by an increase/decrease of a negative/positive potential. Based on our current analysis we cannot unequivocally disambiguate whether our results are due to RS of negative components or RE of positive components; this ambiguity is inherent to all ERP studies. Nevertheless, several canonical ERP peaks have known timing and scalp distribution. Here, we interpreted amplitude changes during the period of these peaks in the most parsimonious way. For example, a period of decreased ERP overlapping with the early P1 peak is interpreted as repetition suppression of a dipolar source activity in the extrastriate cortex projecting its positive field over posterior electrodes (Di Russo et al., 2002; Murphy et al., 2012). While our current results provide objective phenomenological descriptions of repetition effects of scalp-recorded ERPs, future steps will involve studying repetition effects with more mechanistically interpretable computational and biophysically plausible network models (e.g., Garrido et al., 2009).

### Figure captions

Figure 1. Paradigm and 1<sup>st</sup>-level design matrix. A) We used a simple stimulus repetition paradigm where line drawings of everyday objects were repeated 6 to 10 times. Between the 6<sup>th</sup>-10<sup>th</sup> presentations a change in the viewing angle was introduced, after which the original picture was repeated two times. Note that our analysis focused on the first six presentations where stimuli did not change over repetitions. B) Covariates plotted over the 1<sup>st</sup>-level design matrix. Image number corresponds to images for mean ERPs to the 1-6 presentations in four blocks (x axis). ERPs were modelled with a parametric modulator and a main regressor for each block (y axis right). The exponential function (mean centered) used for modelling repetition effects for the first block is plotted in blue over the design matrix (y axis left). C and D) Covariates plotted over the 1<sup>st</sup>-level design matrix for the Change detection and the Linear models, respectively.

Figure 2. Histograms of  $\Delta$ LME. A) Histograms over the voxels within a mask defined by the “logical AND” conjunction of significant voxels under any of the three models, and B) over all voxels in the whole 3D space-time volume.

Figure 3. Statistical parametric maps (SPM) and time course of repetition effects. A) Main effect of stimulus repetition (pooled across the four experimental blocks), F-values thresholded at  $p < 0.05$  FWE (whole-scalp corrected), overlaid on the contrast images. Asterisks mark scalp locations for electrodes shown in B and C. Numbers show activations as indicated in Table 2. Panels from top to bottom show observed intervals of RS in the 86-140ms, RE in the 320-340ms, and RS in the 322-

360ms and 400-446ms time windows, with occipital, occipito-temporal, temporo-parietal, and fronto-temporal distribution, respectively. Note that activation #4 is not plotted separately as it showed a similar dynamics and temporal topography to that of activation #2 peaking at 346ms. B) Box plots of ERP amplitudes for each stimulus presentation. Red lines represent the mean; points (subjects) shown together with the 95% confidence interval of the mean (1.96 SEM) in red and a 1 SD interval in blue. Time windows of significant repetition effects are indicated in each subplot at the electrode sites closest to corresponding cluster maxima. C) Grand mean ERP waveforms elicited by the 1st and 6th stimulus presentation with 95% confidence interval are shown for illustration purposes. Note that statistics were carried out on 3D scalp space-time parameter estimates which were based on ERPs for the 1st, 2nd, 3rd, 4th, 5th, and 6th stimulus presentation. Black horizontal bars mark intervals of significant repetition effects.

### Table captions

Table 1. Summary of model comparison statistics. The results show that the distribution of  $\Delta$ LME values are not Gaussian and median  $\Delta$ LME values are significantly different from zero for all comparisons. The absolute value of median  $\Delta$ LME in all comparisons was  $>12$ . Notably, a difference in LME  $>5$  is considered as very strong evidence in favour of the superior model (Kass and Raftery, 1995).

Table 2. Test statistics for repetition effects. Significant activations are arranged and numbered according to size. P-values and statistics are given for up to three peaks within each activation.

### Conflict of interest

We have no conflict of interest or competing interests to disclose.

### Author contributions

GS and IC designed the study. GS, JH, and KES analysed the data. GS, JH, IC and KES interpreted the results. All authors contributed to writing the manuscript.

### Acknowledgements

The authors acknowledge support by the René and Susanne Braginsky Foundation (KES) and the University of Zurich (KES). This research was supported by the NKFIH research grant 119587 (IC). We are grateful to Zsuzsanna D'Albini and Judit Roschéné Farkas for collecting data.

## Data accessibility

Data are available from the corresponding author upon request.

## Abbreviations

EEG, Electroencephalography; ERP, event-related potential; SPM, statistical parametric mapping; GLM, general linear model; RS, repetition suppression; RE, repetition enhancement; PE, prediction error; SSA, stimulus specific adaptation; PC, predictive coding; FWHM, full-width at half-maximum; FWE, family-wise error;

## References

- Andrade GN, Butler JS, Mercier MR, Molholm S, Foxe JJ (2015) Spatio-temporal dynamics of adaptation in the human visual system: a high-density electrical mapping study. *Eur J Neurosci* 41:925-939.
- Aramakis V, Bandrowski A, Ashe J (1997) Activation of muscarinic receptors modulates NMDA receptor-mediated responses in auditory cortex. *Exp Brain Res*. 113:484-496.
- Auksztulewicz R, Friston K (2015) Attentional Enhancement of Auditory Mismatch Responses: a DCM/MEG Study. *Cereb Cortex* 25:4273-4283.
- Auksztulewicz R, Friston K (2016) Repetition suppression and its contextual determinants in predictive coding. *Cortex* 80:125-140.
- Baldeweg T (2006) Repetition effects to sounds: evidence for predictive coding in the auditory system. *Trends Cogn Sci* 10:93-94.
- Baldeweg T (2007) ERP Repetition Effects and Mismatch Negativity Generation. *Journal of Psychophysiology* 21:204-213.
- Barron HC, Garvert MM, Behrens TE (2016) Repetition suppression: a means to index neural representations using BOLD? *Philos Trans R Soc Lond B Biol Sci* 371.
- Boehnke SE, Berg DJ, Marino RA, Baldi PF, Itti L, Munoz DP (2011) Visual adaptation and novelty responses in the superior colliculus. *Eur J Neurosci* 34:766-779.
- Bogacz R (2017) A tutorial on the free-energy framework for modelling perception and learning. *J Math Psychol* 76:198-211.
- Chen G, Chen P, Tan H, Ma D, Dou F, Feng J, Yan Z (2008) Regulation of the NMDA receptor-mediated synaptic response by acetylcholinesterase inhibitors and its impairment in an animal model of Alzheimer's disease. *Neurobiol Aging*. 29:1795-1804.
- Chennu S, Noreika V, Gueorguiev D, Blenkmann A, Kochen S, Ibanez A, Owen AM, Bekinschtein TA (2013) Expectation and attention in hierarchical auditory prediction. *J Neurosci* 33:11194-11205.
- Chennu S, Noreika V, Gueorguiev D, Shtyrov Y, Bekinschtein TA, Henson R (2016) Silent Expectations: Dynamic Causal Modeling of Cortical Prediction and Attention to Sounds That Weren't. *J Neurosci* 36:8305-8316.
- Clark A (2015) *Surfing Uncertainty: Prediction, Action, and the Embodied Mind*. In. Oxford: Oxford University Press.
- Clifford CW, Webster MA, Stanley GB, Stocker AA, Kohn A, Sharpee TO, Schwartz O (2007) Visual adaptation: neural, psychological and computational aspects. *Vision Res* 47:3125-3131.
- Corlett PR, Honey GD, Krystal JH, Fletcher PC. (2011) Glutamatergic model psychoses: prediction error, learning, and inference. *Neuropsychopharmacology*. 36:294-315.

- Csukly G, Stefanics G, Komlosi S, Czigler I, Czobor P (2013) Emotion-related visual mismatch responses in schizophrenia: impairments and correlations with emotion recognition. *PLoS One* 8:e75444.
- Czigler I, Intraub H, Stefanics G (2013) Prediction beyond the borders: ERP indices of boundary extension-related error. *PLoS One* 8:e74245.
- Dayan P, Hinton GE, Neal RM, Zemel RS (1995) The Helmholtz machine. *Neural Comput* 7:889-904.
- de Gardelle V, Waszczuk M, Egner T, Summerfield C (2013) Concurrent repetition enhancement and suppression responses in extrastriate visual cortex. *Cereb Cortex* 23:2235-2244.
- Delorme A, Makeig S (2004) EEGLAB: an open source toolbox for analysis of single-trial EEG dynamics including independent component analysis. *J Neurosci Methods* 134:9-21.
- Di Russo F, Martinez A, Sereno MI, Pitzalis S, Hillyard SA (2002) Cortical sources of the early components of the visual evoked potential. *Hum Brain Mapp* 15:95-111.
- Eddy M, Schmid A, Holcomb PJ (2006) Masked repetition priming and event-related brain potentials: a new approach for tracking the time-course of object perception. *Psychophysiology* 43:564-568.
- Egner T, Monti JM, Summerfield C (2010) Expectation and surprise determine neural population responses in the ventral visual stream. *J Neurosci* 30:16601-16608.
- Farkas K, Stefanics G, Marosi C, Csukly G (2015) Elementary sensory deficits in schizophrenia indexed by impaired visual mismatch negativity. *Schizophr Res* 166:164-170.
- Feldman H, Friston KJ (2010) Attention, uncertainty, and free-energy. *Front Hum Neurosci.* 4:215.
- Flandin G, Friston KJ (2017) Analysis of family-wise error rates in statistical parametric mapping using random field theory. *Hum Brain Mapp.* doi: 10.1002/hbm.23839.
- Friston K (2005) A theory of cortical responses. *Philos Trans R Soc Lond B Biol Sci* 360:815-836.
- Friston KJ (2007) Statistical parametric mapping. In Friston, K.J., Ashburner, J., Kiebel, S.J., Nichols, T.E., Penny, W.D. (eds) *Statistical Parametric Mapping: The Analysis of Functional Brain Images*. Academic Press, London. pp. 10-31.
- Friston KJ, Brown HR, Siemerkus J, Stephan KE (2016) The dysconnection hypothesis (2016). *Schizophr Res.* 176:83-94.
- Friston KJ, Stephan KE, Lund TE, Morcom A, Kiebel S. (2005) Mixed-effects and fMRI studies. *Neuroimage.* 24:244-252.
- Garrido MI, Kilner JM, Kiebel SJ, Stephan KE, Baldeweg T, Friston KJ (2009) Repetition suppression and plasticity in the human brain. *Neuroimage* 48:269-279.
- Garrido MI, Sahani M, Dolan RJ (2013) Outlier responses reflect sensitivity to statistical structure in the human brain. *PLoS Comput Biol* 9:e1002999.
- Gilbert JR, Gotts SJ, Carver FW, Martin A (2010) Object repetition leads to local increases in the temporal coordination of neural responses. *Front Hum Neurosci* 4:30.
- Gosling A, Thoma V, de Fockert JW, Richardson-Klavehn A (2016) Event-Related Potential Effects of Object Repetition Depend on Attention and Part-Whole Configuration. *Front Hum Neurosci* 10:478.
- Grill-Spector K, Henson R, Martin A (2006) Repetition and the brain: neural models of stimulus-specific effects. *Trends Cogn Sci* 10:14-23.
- Gruber T, Giabbiconi CM, Trujillo-Barreto NJ, Muller MM (2006) Repetition suppression of induced gamma band responses is eliminated by task switching. *Eur J Neurosci* 24:2654-2660.
- Gruber T, Muller MM (2006) Oscillatory brain activity in the human EEG during indirect and direct memory tasks. *Brain Res* 1097:194-204.
- Guillaume C, Guillery-Girard B, Chaby L, Lebreton K, Hugueville L, Eustache F, Fiori N (2009) The time course of repetition effects for familiar faces and objects: an ERP study. *Brain Res* 1248:149-161.
- Hatfield G (2002) Perception as Unconscious Inference. In: *Perception and the Physical World: Psychological and Philosophical Issue in Perception* (Heyer D, Mausfeld R, eds), pp 115-143: John Wiley & Sons.

- Henson RN (2003) Neuroimaging studies of priming. *Prog Neurobiol* 70:53-81.
- Henson RN, Mouchlianitis E, Matthews WJ, Kouider S (2008) Electrophysiological correlates of masked face priming. *Neuroimage* 40:884-895.
- Henson RN, Rylands A, Ross E, Vuilleumier P, Rugg MD (2004) The effect of repetition lag on electrophysiological and haemodynamic correlates of visual object priming. *Neuroimage* 21:1674-1689.
- Hinton GE, Dayan P, Frey BJ, Neal RM (1995) The "wake-sleep" algorithm for unsupervised neural networks. *Science* 268:1158-1161.
- Hohwy J (2013) *The predictive mind*. Oxford: Oxford University Press.
- Ibbotson MR (2005) Physiological Mechanisms of Adaptation in the Visual System. In: *Fitting the Mind to the World: Adaptation and After-Effects in High-Level Vision* (Clifford CWG RG, ed), pp 17-46. Oxford: Oxford Univ. Press.
- Kaliukhovich DA, Vogels R (2014) Neurons in macaque inferior temporal cortex show no surprise response to deviants in visual oddball sequences. *J Neurosci* 34:12801-12815.
- Kass RE, Raftery AE (1995) Bayes Factors. *JASA* 90:773-795.
- Kiebel SJ, Friston KJ (2004) Statistical parametric mapping for event-related potentials: I. Generic considerations. *Neuroimage* 22:492-502.
- Kiefer A (2017) Literal Perceptual Inference. In: *Philosophy and Predictive Processing: 17* (Metzinger T, Wiese W, eds). Frankfurt am Main: MIND Group.
- Kim MS, Jang KM, Che H, Kim DW, Im CH (2012) Electrophysiological correlates of object-repetition effects: sLORETA imaging with 64-channel EEG and individual MRI. *BMC Neurosci* 13:124.
- Kohn A (2007) Visual adaptation: physiology, mechanisms, and functional benefits. *J Neurophysiol* 97:3155-3164.
- Kok P, Rahnev D, Jehee JF, Lau HC, de Lange FP (2012) Attention reverses the effect of prediction in silencing sensory signals. *Cereb Cortex* 22:2197-2206.
- Kovacs-Balint Z, Stefanics G, Trunk A, Hernadi I (2014) Automatic detection of trustworthiness of the face: a visual mismatch negativity study. *Acta Biol Hung* 65:1-12.
- Krekelberg B, Boynton GM, van Wezel RJ (2006) Adaptation: from single cells to BOLD signals. *Trends Neurosci* 29:250-256.
- Lieder F, Daunizeau J, Garrido MI, Friston KJ, Stephan KE (2013) Modelling Trial-by-Trial Changes in the Mismatch Negativity. *PLoS Comput Biol* 9(2): e1002911.
- Litvak V, Mattout J, Kiebel S, Phillips C, Henson R, Kilner J, Barnes G, Oostenveld R, Daunizeau J, Flandin G, Penny W, Friston K (2011) EEG and MEG data analysis in SPM8. *Comput Intell Neurosci* 2011:852961.
- Moran RJ, Campo P, Symmonds M, Stephan KE, Dolan RJ, Friston KJ (2013) Free energy, precision and learning: the role of cholinergic neuromodulation. *J Neurosci*. 33:8227-8236.
- Mumford JA, Nichols T. (2009) Simple group fMRI modeling and inference. *Neuroimage*. 47:1469-1475.
- Murphy JW, Kelly SP, Foxe JJ, Lalor EC (2012) Isolating early cortical generators of visual-evoked activity: a systems identification approach. *Exp Brain Res* 220:191-199.
- Noppeney U, Penny WD (2006) Two approaches to repetition suppression. *Hum Brain Mapp* 27:411-416.
- Puce A, Allison T, McCarthy G (1999) Electrophysiological studies of human face perception. III: Effects of top-down processing on face-specific potentials. *Cereb Cortex* 9:445-458.
- Rao RP, Ballard DH (1999) Predictive coding in the visual cortex: a functional interpretation of some extra-classical receptive-field effects. *Nat Neurosci* 2:79-87.
- Recasens M, Leung S, Grimm S, Nowak R, Escera C (2015) Repetition suppression and repetition enhancement underlie auditory memory-trace formation in the human brain: an MEG study. *Neuroimage* 108:75-86.
- Sanchez-Vives MV, Nowak LG, McCormick DA (2000) Cellular mechanisms of long-lasting adaptation in visual cortical neurons in vitro. *J Neurosci* 20:4286-4299.



- Schendan HE, Kutas M (2003) Time course of processes and representations supporting visual object identification and memory. *J Cogn Neurosci* 15:111-135.
- Schnyer DM, Ryan L, Trouard T, Forster K (2002) Masked word repetition results in increased fMRI signal: a framework for understanding signal changes in priming. *Neuroreport* 13:281-284.
- Schwarz GE (1978) Estimating the dimension of a model, *Annals of Statistics* 6:461-464.
- Schweinberger SR, Neumann MF (2016) Repetition effects in human ERPs to faces. *Cortex* 80:141-153.
- Segaert K, Weber K, de Lange FP, Petersson KM, Hagoort P (2013) The suppression of repetition enhancement: a review of fMRI studies. *Neuropsychologia* 51:59-66.
- Solomon SG, Kohn A (2014) Moving sensory adaptation beyond suppressive effects in single neurons. *Curr Biol* 24:R1012-1022.
- Stefanics G, Czigler I (2012) Automatic prediction error responses to hands with unexpected laterality: an electrophysiological study. *Neuroimage* 63:253-261.
- Stefanics G, Heinzle J, Horváth AA, Stephan KE (2018) Visual mismatch and predictive coding: A computational single-trial ERP study. *J Neurosci* (in press) DOI: 10.1523/JNEUROSCI.3365-17.2018
- Stefanics G, Kimura M, Czigler I (2011) Visual mismatch negativity reveals automatic detection of sequential regularity violation. *Front Hum Neurosci* 5:46.
- Stephan KE, Diaconescu AO, Iglesias S. (2016) Bayesian inference, dysconnectivity and neuromodulation in schizophrenia. *Brain*. 139(Pt 7):1874-1876.
- Stephan KE, Penny WD, Daunizeau J, Moran RJ, Friston KJ (2009) Bayesian model selection for group studies. *Neuroimage* 46:1004-1017.
- Stephan KE, Weiskopf N, Drysdale PM, Robinson PA, Friston KJ (2007) Comparing hemodynamic models with DCM. *Neuroimage* 38:387-401.
- Summerfield C, Trittschuh EH, Monti JM, Mesulam MM, Eger T (2008) Neural repetition suppression reflects fulfilled perceptual expectations. *Nat Neurosci* 11:1004-1006.
- Szekely A et al. (2004) A new on-line resource for psycholinguistic studies. *Journal of Memory and Language* 51:247-250.
- Vogels R (2016) Sources of adaptation of inferior temporal cortical responses. *Cortex* 80:185-195.
- von der Behrens W, Bauerle P, Kossl M, Gaese BH (2009) Correlating stimulus-specific adaptation of cortical neurons and local field potentials in the awake rat. *J Neurosci* 29:13837-13849.
- Vuilleumier P, Schwartz S, Duhoux S, Dolan RJ, Driver J (2005) Selective attention modulates neural substrates of repetition priming and "implicit" visual memory: suppressions and enhancements revealed by FMRI. *J Cogn Neurosci* 17:1245-1260.
- Webster MA (2011) Adaptation and visual coding. *J Vis* 11.
- Worsley KJ, Friston KJ (1995) Analysis of Fmri Time-Series Revisited - Again. *Neuroimage* 2:173-181.

$\Delta$ LME	Median	Mean	Std	% of voxels >0	Kolmogorov-Smirnov test (D)	Kolmogorov-Smirnov test (p)	Wilcoxon signed rank test (Z)	Wilcoxon signed rank test (p)
EXP vs LIN (within mask)	38.77	43.59	14.24	100	1	<0.00001	-53.58	<0.00001
EXP vs CDT (within mask)	16.73	21.48	11.42	100	1	<0.00001	-53.58	<0.00001
CDT vs LIN (within mask)	-21.76	-22.11	9.63	0.57	0.9839	<0.00001	-53.57	<0.00001
EXP vs LIN (all voxels)	25.03	29.15	18.58	98.36	0.9704	<0.00001	-375.75	<0.00001
EXP vs CDT (all voxels)	13.78	13.73	11.03	89.28	0.8439	<0.00001	-348.01	<0.00001
CDT vs LIN (all voxels)	-12.77	-15.43	19.98	21.43	0.7286	<0.00001	-283.30	<0.00001

Table 1.

Activation number	Activation size (# voxels )	Latency (ms)	p-value (FWE-corr.)	F-statistic	Equivalent Z-statistic
1	7795	422	0.00180	61.2678	4.81524
		440	0.00404	52.1248	4.60680
		380	0.03835	31.7057	3.95946
2	4156	346	0.00363	53.2627	4.63474
		346	0.00737	46.0016	4.44467
3	570	102	0.00474	50.4278	4.56392
4	52	298	0.03762	31.8552	3.96560
5	8	112	0.04366	30.7134	3.91801
6	2	108	0.04893	29.8551	3.88109

Table 2.





