

Open access • Posted Content • DOI:10.1101/2020.03.17.995704

Temporal context actively shapes EEG signatures of time perception — Source link ☑

Atser Damsma, Nadine Schlichting, Hedderik van Rijn Institutions: University of Groningen Published on: 18 Mar 2020 - bioRxiv (Cold Spring Harbor Laboratory) Topics: Time perception

Related papers:

- The Effect of a Regular Auditory Context on Perceived Interval Duration.
- Timing Rhythms: Perceived Duration Increases with a Predictable Temporal Structure of Short Interval Fillers.
- · Perception of time in articulated visual events
- · Modality-specific temporal constraints for state-dependent interval timing.
- · Alpha/beta power compresses time in sub-second temporal judgments



Running head: CONTEXT ACTIVELY SHAPES TIME PERCEPTION

Temporal context actively shapes EEG signatures of time perception

Atser Damsma^{1*}, Nadine Schlichting¹, Hedderik van Rijn¹

¹Department of Psychology, University of Groningen, Grote Kruisstraat 2/1, 9712 TS, Groningen, the

Netherlands

*Corresponding author: Atser Damsma (<u>a.damsma@rug.nl</u>)

CONTEXT ACTIVELY SHAPES TIME PERCEPTION

2

Abstract

2	Our subjective perception of time is optimized to temporal regularities in the environment. This is
3	illustrated by the central tendency effect: when estimating a range of intervals, short intervals are
4	overestimated whereas long intervals are underestimated to reduce the overall estimation error. Most
5	models of interval timing ascribe this effect to the weighting of the current interval with previous
6	memory traces after the interval has been perceived. Alternatively, the perception of the duration
7	could already be flexibly tuned to its temporal context. We investigated this hypothesis using an
8	interval reproduction task in which human participants (both sexes) reproduced a shorter and longer
9	interval range. As expected, reproductions were biased towards the subjective mean of each presented
10	range. EEG analyses showed that temporal context indeed affected neural dynamics during the
11	perception phase. Specifically, longer previous durations decreased CNV and P2 amplitude and
12	increased beta power. In addition, multivariate pattern analysis showed that it is possible to decode
13	context from the transient EEG signal quickly after both onset and offset of the perception phase.
14	Together, these results suggest that temporal context creates dynamic expectations which actively
15	affect the <i>perception</i> of duration.
16	Keywords: time perception; context; Bayesian perception; EEG

17

1

Significance Statement

18 The subjective sense of duration does not arise in isolation, but is informed by previous experiences. 19 This is demonstrated by abundant evidence showing that the production of duration estimates is 20 biased towards previously experienced time intervals. However, it is yet unknown whether this 21 temporal context actively affects perception or only asserts its influence in later, post-perceptual 22 stages as proposed by most current formal models of this task. Using an interval reproduction task, we 23 show that EEG signatures flexibly adapt to the temporal context during perceptual encoding. 24 Furthermore, interval history can be decoded from the transient EEG signal even when the current 25 duration was identical. Thus, our results demonstrate that context actively influences perception.

Introduction

CONTEXT ACTIVELY SHAPES TIME PERCEPTION

26

3

20	Introduction
27	The way humans experience time is not only driven by the current stimulus, but is also
28	influenced by previous experiences. According to Bayesian observer models, humans integrate noisy
29	sensory representations (the likelihood) with previously learned stimulus statistics (the prior
30	distribution). This is illustrated by the temporal context or central tendency effect: when presented
31	with a range of intervals, short intervals are overestimated and long intervals are underestimated
32	(Jazayeri & Shadlen, 2010). Furthermore, the prior distribution has been shown to be dynamically
33	updated, such that more recent intervals have a greater influence on the current estimate (Dyjas,
34	Bausenhart, & Ulrich, 2012; Taatgen & van Rijn, 2011; Wiener, Thompson, & Branch Coslett, 2014).
35	Although there is abundant behavioral evidence for Bayesian integration in human time perception
36	(Acerbi, Wolpert, & Vijayakumar, 2012; Cicchini, Arrighi, Cecchetti, Giusti, & Burr, 2012; Gu,
37	Jurkowski, Lake, Malapani, & Meck, 2015; Hallez, Damsma, Rhodes, van Rijn, & Droit-Volet, 2019;
38	Jazayeri & Shadlen, 2010; Maaß, Riemer, Wolbers, & van Rijn, 2019; Maaß, Schlichting, & van Rijn,
39	2019; Roach, McGraw, Whitaker, & Heron, 2017; Schlichting et al., 2018; Shi, Church, & Meck,
40	2013), its temporal locus and neural underpinnings are not yet understood.
41	Computational models of interval timing often (implicitly) assume that only after perception
42	has completed, the noisy interval percept is weighted with previous memory traces representing the
43	prior (e.g., Di Luca & Rhodes, 2016; Jazayeri & Shadlen, 2010; Taatgen & van Rijn, 2011).
44	Alternatively, however, prior experience might actively affect perception, as evidenced by recent
45	behavioral (Cicchini, Benedetto, & Burr, 2020; Cicchini, Mikellidou, & Burr, 2017; Zimmermann &
46	Cicchini, 2020), fMRI (St. John-Saaltink, Kok, Lau, & De Lange, 2016) and single neuron findings
47	(Sohn, Narain, Meirhaeghe, & Jazayeri, 2019). Specifically, Sohn et al. (2019) showed that neurons in
48	the prefrontal cortex of monkeys exhibited different firing rate patterns based on the prior during
49	interval estimation.
50	In humans, evidence is now emerging that electroencephalography (EEG) signatures in

In humans, evidence is now emerging that electroencephalography (EEG) signatures in
timing tasks are modulated by recently perceived durations. In a bisection task, longer prior durations
led to a larger amplitude of the contingent negative variation (CNV) and increased beta oscillations
power (Wiener, Parikh, Krakow, & Coslett, 2018; Wiener & Thompson, 2015). Crucially, however,

CONTEXT ACTIVELY SHAPES TIME PERCEPTION

4

54 these studies required an active comparison to the standard interval, in which EEG signatures have 55 been shown to reflect an adjustment of the decision threshold (Ng, Tobin, & Penney, 2011; see also Boehm, Van Maanen, Forstmann, & van Rijn, 2014). Any context-based changes in these signatures 56 might reflect updating of the comparison process. It is therefore still an open question what the 57 58 temporal locus of the context effect is: Does the prior exert its influence in post-perceptual stages or are purely perceptual processes already affected by previous experiences? 59 We tested the influence of temporal context in an interval reproduction task, which allowed us 60 to distill EEG signals during the perception phase in which no decision or motor response was 61 required that could yield fallacious conclusions regarding the effect of context effects during 62 63 perception. Participants reproduced two different interval ranges (the *short* and the *long context*). The ranges shared one interval (the overlapping interval), providing a condition in which the physical 64 65 stimulus was the same, but the temporal context was different. We show that temporal context affects 66 three EEG signatures that have previously been associated with time perception during the perception 67 phase: the CNV and beta oscillations, but also the offset P2, which has been shown to predict 68 subjective interval perception better than the CNV (Kononowicz & van Rijn, 2014; Kruijne, Olivers, 69 & van Rijn, 2021). A data-driven approach reveals that temporal context can be decoded from 70 transient neural dynamics during the perception phase using multivariate pattern analysis (MVPA). 71 Together, these results show that temporal context actively shapes the perception of duration, 72 falsifying most current formal theories of interval timing.

73

Materials and Methods

74 Participants

Twenty-seven healthy adults (22 females; age range 18 - 33 years, M = 21.33, SD = 3.78years) participated in the experiment for course credits in the University of Groningen Psychology program or monetary compensation (\in 14). Two participants were excluded from analysis during preprocessing due to excessive artifacts in the EEG data. The study was approved by the Psychology Ethical Committee of the University of Groningen (17141-S-NE). Written informed consent was

CONTEXT ACTIVELY SHAPES TIME PERCEPTION

5

80 obtained before the experiment. After the experiment, the participants were debriefed about the aim of

81 the study.

82 Stimuli and apparatus

Stimuli were presented using the Psychophysics Toolbox 3.0.12 (Brainard, 1997; Kleiner et
al., 2007) in Matlab 2014b. Intervals were presented as continuous 440 Hz sine wave tones. These
auditory stimuli were presented on Sennheiser HD 280 Pro stereo headphones at a comfortable sound
level. Visual stimuli were presented in the center of the screen in Helvetica size 25 in white on a dark
grey background using a 27-inch Iiyama ProLite G27773HS monitor with a 1920x1080 resolution at
100 Hz. The index-finger trigger buttons of a gamepad (SideWinder Plug & Play Game Pad,
Microsoft Corporation) were used to record responses.

90 **Procedure**

91 Participants performed an auditory interval reproduction task (Figure 1A). Every trial started 92 with a central fixation cross with a uniform random duration between 2 and 3 s. Then, an exclamation 93 mark was presented for 0.7 s, after which the auditory interval was presented (the *perception phase*) 94 while the exclamation mark remained on the screen. To signal the next phase, the exclamation mark 95 was replaced by a question mark which was presented for 1.5 s. Next, the continuous tone was 96 presented again, with the question mark remaining on the screen, which participants had to terminate 97 by pressing a button (the *reproduction phase*). Participants were instructed to match the duration of 98 this second tone to the duration of the first tone as accurately as possible.

99 The task involved two different interval ranges, the short context (0.625 s, 0.75 s, and 0.9 s)100 and the long context (0.9 s, 1.08 s, and 1.296 s) (Figure 1A). Crucially, there was an overlapping 101 interval that was presented in both contexts (0.9 s). The experiment consisted of four blocks, two of 102 which used intervals of the short context, and two of which used intervals of the long context. Block 103 order was counterbalanced across participants, with the constraint that the context would alternate 104 every block. Within a block, each duration of the short or the long context was presented 30 times. 105 amounting to a total of 90 trials per block and 360 trials over the whole experiment. The presentation order was random, with the constraint that every possible subsequent pair of intervals was presented 106 107 equally often (i.e., first-order counterbalancing). The hand needed for reproduction was switched after

CONTEXT ACTIVELY SHAPES TIME PERCEPTION

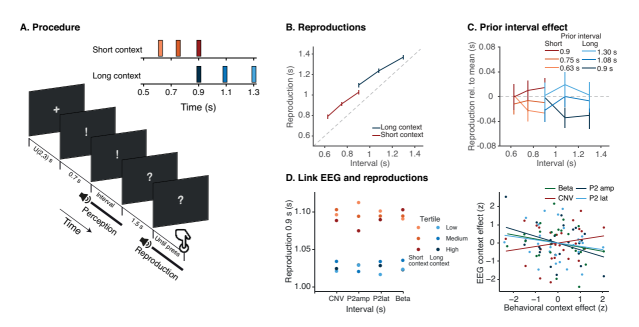


Figure 1. Task and behavioral results. A) Behavioral procedure of the experiment. Participants performed an interval reproduction task in which they heard a tone for a certain duration (*perception phase*). After an ISI of 1.5 s, they were asked to reproduce this duration by pressing a button to indicate the offset of the *reproduction phase*. In separate blocks, the perception phase consisted of three short or three relatively long durations (the *short* and the *long context*, respectively). One interval was presented in both contexts (the *overlapping interval* of 0.9 s). B) Average behavioral reproduction results. Error bars represent the standard error of the mean. C) Average reproduction of the overlapping interval (0.9 s) for the different intervals in the previous trial, relative to average reproduction in the context condition. Overall, reproductions were longer when the prior interval was longer. D) Link between the EEG signatures and reproductions. The left panel shows the reproduction of the overlapping interval for relatively low, medium, and high values (i.e., tertiles) of the CNV amplitude, P2 amplitude, P2 latency, and beta power. The right panel shows the correlation between participants' behavioral context effect and their context effect in the different EEG signatures (all values were *z*-scored). Dots represent individual participants, while the lines represent linear regression lines.

- 108 two blocks. Prior to each block, participants were instructed which hand (i.e., which gamepad button)
- 109 they would use to terminate the duration and which set of intervals would be presented (termed set A
- and set B for the short and long context, respectively; see also Maaß, Schlichting, et al., 2019), while
- 111 they were not informed about the relative durations or distributions associated with the sets (i.e., that
- the sets were associated with a short and long interval range). Participants could take self-timed
- 113 breaks between blocks. Prior to the experiment, participants performed two practice trials with
- 114 durations outside the range of both context conditions (0.4 s and 2 s). Experiment scripts are available
- 115 at <u>https://osf.io/sgbjz/.</u>

116 **EEG acquisition**

- 117 EEG signals were recorded from 62 Ag/AgCl electrodes, placed in accordance with the
- 118 international 10-20 system (WaveGuard EEG cap, eemagine Medical Imaging Solutions GmbH,

CONTEXT ACTIVELY SHAPES TIME PERCEPTION

7

119 Berlin, Germany). The ground electrode was placed onto the left side of the collarbone and the 120 mastoids served as location for the reference electrodes. The electrooculogram (EOG) was recorded 121 from the outer sides of both eyes and from the top and bottom of the left eye. Data was collected at a 122 sampling frequency of 512 Hz using a TMSi Refa 8-64 amplifier. Before the experiment, impedances 123 of all electrodes were reduced to below $5k\Omega$. Participants were instructed to blink only between trials 124 and not to move during the experiment.

125 **EEG pre-processing**

EEG pre-processing was performed using the FieldTrip toolbox (Oostenveld, Fries, Maris, & Schoffelen, 2011). EEG data was re-referenced to the averaged mastoids and filtered using a Butterworth IIR band-pass filter with a high-pass frequency of 0.01 Hz and a low-pass frequency of 80 Hz. Subsequently, trial epochs were created from -1 s until 6 s relative to the onset of the perception phase. Artifacts were corrected using independent component analysis (ICA). Epochs that exceeded an amplitude range of 120 μ V were removed from the dataset. On average, 10.72% (*SD* = 6.10) of the 360 trials were discarded.

133 Data Analysis

134 Behavioral analysis. Reproductions lower than 0.1 s and higher than 2 s were excluded from 135 analysis (0.2% of the data). To test whether reproductions were influenced by context, we fitted a 136 linear mixed model (LMM) using the *lme4* package (Bates, Mächler, Bolker, & Walker, 2015) in R 137 (R Core Team, 2016), including interval, context, their interaction and prior interval (i.e., the interval in the previous trial) as fixed factors. To facilitate interpretation of the results, interval and prior 138 139 interval were centered at 0.9 s and the factor context was recoded using effect coding (-0.5 for short 140 and 0.5 for long context). In addition to the random intercept of participant, we sequentially added 141 random slope terms and tested whether they improved the model with a likelihood ratio test. We will 142 here report the results of the best fitting model, which included random slopes for interval and prior 143 interval.

144 ERP analysis. All EEG analyses reported here focused on the perception phase. An overview
145 of the EEG results in the reproduction phase is available in the supplementary materials (section 1) at

CONTEXT ACTIVELY SHAPES TIME PERCEPTION

8

146 <u>https://osf.io/sgbjz/</u>. The CNV and beta signatures in the reproduction phase show trends that are

147 qualitatively similar to the perception phase, although they appear to be less strong.

148 **CNV.** The CNV analysis was performed on a fronto-central electrode cluster (electrodes Cz. C1, C2, FC2, FC1, FC2) (Kononowicz & van Rijn, 2014; Ng et al., 2011). A 10 Hz Butterworth low-149 150 pass filter was applied and the ERP was baselined to the average signal in the 0.1 s window before interval onset. To test the effect of global context during the perception phase, we compared the ERP 151 152 of the overlapping interval in the short and the long context using a cluster-based permutation test 153 (Maris & Oostenveld, 2007) in the window 0-1.2 s from interval onset. The permutation test assessed 154 whether the difference was different from zero by computing 100.000 permutations using the t-155 statistic, controlling for multiple comparisons with a cluster significance threshold of p < .05. To assess the influence of the prior interval on CNV, we calculated the average amplitude in the time 156 157 window that showed CNV differences in the previously mentioned permutation test (0.3-1.01 s), per 158 participant, context and prior interval for the overlapping interval. Next, we tested an LMM predicting 159 this amplitude, including context and prior interval as fixed factors, and participant as a random 160 intercept term.

161 Offset P2. The P2 analysis focused on the EEG signal averaged over the same fronto-central 162 electrode cluster as the CNV analysis, to which a 1-20 Hz Butterworth band-pass filter was applied to 163 minimize CNV-based contamination (cf., Kononowicz & van Rijn, 2014). The ERP was baselined to 164 the average signal in the 0.1 s window around interval offset (cf., Kononowicz & van Rijn, 2014). Similar to the CNV analysis, the ERPs of the overlapping interval in the short and the long context 165 were compared using a cluster-based permutation test in the window 0-0.5 s after interval offset. 166 Next, we calculated P2 amplitude was as the average amplitude between 0.14 and 0.3 s after interval 167 168 offset (this window was based on Kononowicz & van Rijn, 2014). We fitted an LMM predicting P2 169 amplitude, with interval, prior interval, and context as fixed factors, and participant as a random intercept term. The random slope of interval improved the fit and was added to the model. P2 latency 170 171 was calculated as the 50% area latency - the time point at which half of the area under the curve is 172 reached - within the same window (Liesefeld, 2018; Luck, 2005). P2 latency was analyzed using an 173 LMM with the same fixed factors as the P2 amplitude model.

CONTEXT ACTIVELY SHAPES TIME PERCEPTION

9

Because the 1 Hz high-pass filter might induce artifactual effects of opposite polarity before the actual peak (Tanner, Morgan-Short, & Luck, 2015), we also performed the P2 analysis on the data without additional filtering (that is, besides the band-pass filter between 0.01 Hz and 80 Hz applied during pre-processing). We found similar qualitative results, which are reported in the supplementary materials section 2.1 at https://osf.io/sgbjz/.

Multivariate pattern analysis. To investigate transient neural dynamics in more detail, we tested whether it is possible to decode global and local context through MVPA of the EEG signal. Following Wolff, Kandemir, Stokes, and Akyürek (2020), we used a sliding window approach in which the EEG fluctuations were pooled over electrodes and time. A window of 50 data points (98 ms) was moved across the signal in steps of 8 ms, separately for each channel. Within the window, the signal was down-sampled to 10 samples (by taking the average over 5 samples) and baselinecorrected by subtracting the mean within the window from all 10 individual samples.

186 To decode whether an overlapping-interval trial was presented in the short or the long 187 context, the 10 samples per electrode in each time window served as input for 5-fold cross-validation. 188 In each fold, we calculated the Mahalanobis distance (De Maesschalck, Jouan-Rimbaud, & Massart, 189 2000; Wolff, Jochim, Akyürek, & Stokes, 2017; Wolff et al., 2020) between the test trials and the 190 averaged signal of the short and long context, using the covariance matrix estimated from the training 191 trials with a shrinkage estimator (Ledoit & Wolf, 2004). To make the distance estimates more reliable, 192 the 5-fold cross-validation was repeated 50 times and results were averaged. The eventual decoding 193 distances were smoothed with a Gaussian smoothing kernel (SD = 16 ms). To test whether the 194 distance between conditions was significantly different from zero, a cluster-based permutation test 195 was performed.

A similar analysis was performed to decode the duration of the prior interval from the neural dynamics in the current trial. For the overlapping interval, the Mahalanobis distance between every test trial and the average of the prior interval conditions was calculated. This resulted in six difference time series for each condition (including the 0.9 s condition for each context separately and the difference with the trial's own condition). In this way, we aimed to determine whether the distance was higher when the difference between the prior interval condition of the test trial and the other

CONTEXT ACTIVELY SHAPES TIME PERCEPTION

10

possible prior interval conditions was larger. Next, for every time point, we performed a linear
regression on the Mahalanobis distance, using the absolute difference between prior interval
conditions (in seconds) and the difference between context (coded as 0 or 1) as predictors, allowing us
to disentangle the effect of sequential and global context on transient neural dynamics. A clusterbased permutation test was performed on the resulting slope values for prior interval and context, to
test whether they deviated from zero (using a one-sided *t*-test).
To investigate which electrodes are most informative in decoding the context of an

overlapping interval trial, we performed channel-wise decoding: the procedure to decode global context outlined above was performed separately for every electrode. Topographies were created to show the average decoding accuracy at the different electrodes during time windows in which the Mahalanobis distance resulting from the context decoding procedure outlined above (i.e., using all electrodes) was significantly higher than zero.

214 Because the context conditions were blocked in our experimental design, the decoding 215 accuracy might have been inflated by nonstationarities in the EEG signals, which lead to stochastic 216 dependence between trials (Lemm, Blankertz, Dickhaus, & Müller, 2011). Post-hoc, we controlled for 217 this notion by calculating the Mahalanobis distance between the different blocks, for each participant. 218 This allowed us to differentiate between the distances between blocks that were presented in the first 219 and second half of the experiment, and thereby, to test whether the original decoding results could be 220 due to within-block similarities beyond context. In this way, we compared the Mahalanobis distance 221 between the trials in a particular block and the 'same context' and 'different context' block in the 222 other half of the experiment. We found that the results were qualitatively similar to the original 223 analysis, with significant differences between the short and long context immediately after interval onset and after interval offset (analysis details and full results can be found in the supplementary 224 materials section 2.2. at https://osf.io/sgbjz/). 225

Time-frequency analysis. To assess oscillatory power during the perception phase we performed a time-frequency analysis using a single Hanning taper with an adaptive time window of 6 cycles per frequency in steps of 15 ms for frequencies from 4 to 40 Hz, with the amount of spectral smoothing set to 1. We calculated the absolute power from the baseline window of -0.2-0 s relative to

CONTEXT ACTIVELY SHAPES TIME PERCEPTION

11

230 interval onset. The analysis was again focused on fronto-central electrodes (Cz, C1, C2, FCz, FC1,

FC2). Similar to the CNV analysis, all time-frequency analyses were performed on the overlapping

interval to isolate the effect of context while keeping the actual stimulus constant.

233 Per participant, for every time-frequency point, we fitted a linear regression model including 234 prior interval (a continuous variable ranging from the shortest to the longest interval in seconds) and 235 context (short and long context coded as 0 and 1, respectively) as predictors (following an approach 236 similar to Wiener et al., 2018). For every time-frequency point, this resulted in two slope values, 237 expressing the relative influence of the global context and the previous interval. Next, a one-sample t-238 test against zero was performed for the two slope values at each time-frequency point, which was 239 corrected for multiple comparisons using cluster-based permutation (Maris & Oostenveld, 2007). The 240 statistical testing was performed on the frequency range of 8-30 Hz to include alpha power (8–14 Hz; 241 Kononowicz & van Rijn, 2015) and beta power (15–30 Hz; e.g., Haegens et al., 2011; Jenkinson & 242 Brown, 2011; Kononowicz & van Rijn, 2015) during the time window of 0-1.2 s after interval onset.

243 Linking EEG signatures and behavior. We tested in two ways whether EEG signatures during the perception phase predicted behavioral reproductions. First, we computed *single trial* values 244 245 of CNV amplitude, P2 amplitude, P2 latency and beta power. Following the methods described above, 246 for every trial, CNV amplitude was calculated as the average EEG signal in the window 0.3-1.01 s after interval onset, P2 amplitude as the average between 0.14 and 0.3 s after interval offset, P2 247 248 latency as the 50% area latency in the same window, and beta power was calculated as the average power in the time window 0.48-0.84 s after interval onset and the frequency range 23-30 Hz, which 249 250 was based on the permutation test. CNV, P2 amplitude, P2 latency, and beta power values that 251 deviated more than 4 SD from the average were excluded from analysis (0.06%, 0.01%, 0.00% and 0.46% of the trials, respectively). Similar to the behavioral analysis described above, reproductions 252 253 shorter than 0.1 s and longer than 2 s were also excluded from analysis. Next, we computed four LMMs with reproduction as the dependent factor, and CNV amplitude, P2 amplitude, P2 latency, and 254 beta power as fixed factors, respectively. Similar to the analyses described above, the CNV and beta 255 power analyses were focused on the overlapping interval trials. To control for the effect of context on 256 257 both EEG signatures and behavior, context and prior interval were also added as fixed factors to the

CONTEXT ACTIVELY SHAPES TIME PERCEPTION

12

models. The P2 analysis included all intervals, so here, interval was entered as an additional fixed
factor. In all models, participant was included as a random intercept term, and adding random slopes
did not improve the model fit in any of the models.

Second, in addition to the single trial analysis, we looked at individual differences: Do 261 262 participants who show a large context effect in the EEG signatures also show a large behavioral context effect? To this end, for the overlapping interval, we estimated the behavioral context effect 263 (i.e., the difference in reproduction between the long and short context) for each participant, and 264 265 compared it to the context effect of CNV amplitude, P2 amplitude and latency, and beta power, quantified as described in the previous paragraphs. To assess whether these measures were related for 266 each participant, we performed a one-tailed Pearson's correlation test between the individual 267 268 behavioral context effects and the EEG context effects.

269

Results

270 Behavioral results

271 Figure 1B shows the average reproductions for the different intervals. The results of the LMM showed that the reproductions increased with duration ($\beta = 0.77$, SE = 0.03, t = 24.33, p < .001). We 272 273 found a significant effect of global context, showing that reproductions were longer in the long compared to the short context ($\beta = 0.05$, SE = 0.01, t = 7.23, p < .001). In addition, the increase with 274 duration (i.e., the slope) was lower for the long compared to the short context ($\beta = -0.18$, SE = 0.03, t 275 = -7.01, p < .001). Besides the global context effect, reproductions were longer when the interval in 276 the previous trial was longer ($\beta = 0.08$, SE = 0.02, t = 3.41, p = .002). Figure 1C shows the 277 278 reproductions for the different previous intervals, relative to the average reproduction. 279 **ERPs**

280 **CNV.** Figure 2A and 2B show the average fronto-central ERP during the perception phase for 281 the different intervals in the short and the long context, respectively. In addition, Figure 2D shows a 282 direct comparison between the short and the long context of this ERP for the overlapping interval (0.9 283 s). The cluster-based permutation test showed that the CNV was more negative in the short than in the 284 long context in the time windows 0.30-0.65 s (p = .004) and 0.71-1.01 s (p = .003). Thus, while the

CONTEXT ACTIVELY SHAPES TIME PERCEPTION

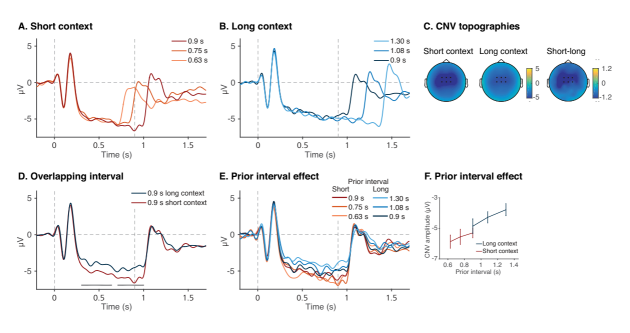


Figure 2. Average ERPs at the fronto-central cluster (Cz, C1, C2, FCz, FC1, FC2) relative to the onset of the perception phase for the different durations in the short (A) and long (B) context. In all panels, vertical grey dashed lines indicate interval onset and offset of the overlapping interval (0.9 s). C) Topographies of the overlapping interval (0.9 s), for the short context, long context, and their difference, during the window of significant difference as indicated by the cluster-based permutation test. D) Average ERP of the overlapping interval (0.9 s) in the short and the long context. Grey horizontal bars indicate significant differences according to the cluster-based permutation test. E) Average ERP of the overlapping interval, split up according to the interval in the previous trial. Red and blue lines show whether the overlapping interval appeared in the short or the long context, respectively. F) Average CNV amplitude for the middle interval, in the time window of significant difference between the short and long context, for the different previous intervals. Error bars represent the standard error of the mean.

- actual interval was the same, CNV amplitude during perception differed depending on the temporal
- context.
- 287 Figure 2E shows the average ERP for the overlapping interval, split for the different previous
- durations, and Figure 2F shows the average CNV in the 0.3-1.01 s window for the different prior
- 289 interval conditions. The LMM results showed that CNV amplitude at the overlapping interval became
- less negative with longer previous trials ($\beta = 2.50$, SE = 0.97, t = 2.57, p = .011). There was no
- evidence for an additional significant effect of context ($\beta = 0.43$, SE = 0.42, t = 1.03, p = .308),
- suggesting that the global context effect on CNV might be largely driven by the previous trial. Post-
- 293 hoc, we tested whether including the interaction between context and prior interval improved the
- model fit, but a likelihood ratio test showed that this was not the case ($\chi^2(1) = 0.10$, p = .750).

14

CONTEXT ACTIVELY SHAPES TIME PERCEPTION

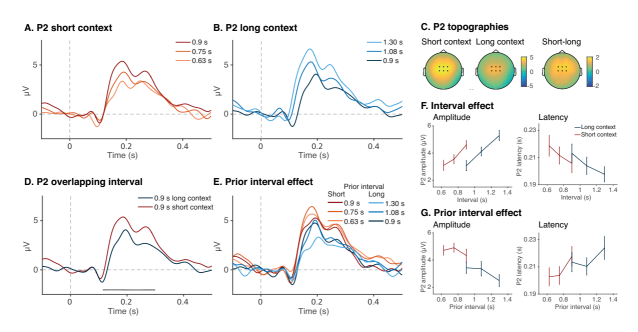


Figure 3. Amplitude and latency of the P2 at the fronto-central cluster (Cz, C1, C2, FCz, FC1, FC2) after the offset of the perception phase. A, B) Grand average ERPs baselined at the offset of the perception phase in the short and the long context, respectively. C) Topographies of P2 amplitude of the overlapping interval (0.9 s), for the short context, long context, and their difference, during the window of significant difference as indicated by the cluster-based permutation test. D) Average ERP of the overlapping interval (0.9 s) in the short and the long context. Grey horizontal bars indicate significant differences according to the cluster-based permutation test. F) Effect of interval on P2 amplitude and latency. The left panel shows P2 amplitude, calculated as the average amplitude in the window 0.14-0.3 s after interval offset for every participant and interval. The right panel shows P2 latency, calculated as the 50% area latency in the same window. G) Effect of the prior interval on P2 amplitude (left) and latency (right). In all figures, error bars represent the standard error of the mean.

Offset P2. Amplitude. Figure 3A and 3B shows the offset P2 for the different intervals in the 295 296 short and the long context, respectively. Figure 3D directly compares the P2 for the overlapping 297 interval in the short and long context. The cluster-based permutation test showed that the amplitude 298 was higher for the short compared to the long context in the window 0.11-0.3 s. Figure 3F shows the 299 average P2 amplitude as a function of interval and context. The LMM showed that the P2 increased 300 with duration ($\beta = 5.56$, SE = 0.62, t = 8.97, p < .001), but that the intercept was significantly lower 301 for the long compared to the short context ($\beta = -0.87$, SE = 0.28, t = -3.10, p = .002). Figure 3E and 302 3G (left panel) show the effect of the prior interval on P2 amplitude for the overlapping interval. In 303 line with the global context effect, the model showed that the P2 decreased with longer previous 304 intervals ($\beta = -1.71$, SE = 0.51, t = -3.35, p = .001). Together, these results show that P2 amplitude reflects the actual duration, as well as the global and local context in which the duration appeared. 305

CONTEXT ACTIVELY SHAPES TIME PERCEPTION

15

Latency. Figure 3F (right panel) shows that P2 latency decreased with the duration of the current interval, which was confirmed by the LMM predicting latency ($\beta = -0.04$, SE = 0.01, t = -3.66, p < .001). There was no evidence that P2 latency was affected by the context, as the fixed factors contex and prior interval did not reach significance (ps > .247). In summary, whereas P2 amplitude reflects the current duration and the general and sequential temporal context, P2 latency only decreases with longer current durations.

312 Multivariate pattern analysis

313 Figure 4A shows the decoding accuracy for the overlapping interval. The permutation test showed a positive cluster immediately after interval onset (0-0.17 s; p = .009) and after interval offset 314 315 (0.99-1.37 s; p < .001). Figure 4C shows the topographies of the channel-wise decoding results during 316 these two clusters, which reflects high parietal and left-lateralized decoding accuracy and high fronto-317 central and right-lateralized decoding accuracy, respectively. Figure 4B shows the slope value of prior 318 interval in the regression analysis predicting Mahalanobis distance. The permutation test showed that 319 there was no evidence for significant clusters for the slope of prior interval or context in the regression analysis (p = .999), showing that MVPA could not distinguish between prior interval conditions based 320 321 on the transient EEG signal.

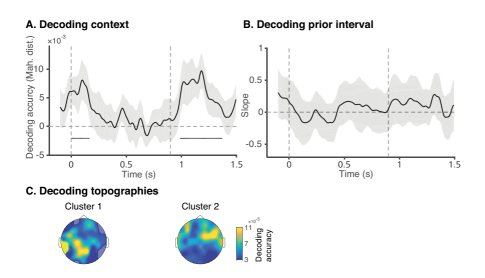


Figure 4. Decoding accuracy relative to the onset of the perception phase. A) Decoding accuracy of context for the overlapping interval as represented by the Mahalanobis distance. Grey horizontal bars indicate a significant difference from zero according to the cluster-based permutation test. Error shading represent 95% Cl of the mean. B) Decoding accuracy of prior interval in the overlapping interval, represented by the slope value of the regression of Mahalanobis distance with prior interval and context as predictors. C) Topographies of channel-wise context decoding accuracy for the overlapping interval, during the first significant cluster in panel A (left) and the second cluster (right). Colors represent the decoding accuracy in Mahalanobis distance.

16

CONTEXT ACTIVELY SHAPES TIME PERCEPTION

322 Time-frequency analysis

To assess oscillatory power during the perception phase, we calculated a linear regression of 323 frequency power at fronto-central electrodes with context (short vs long) and prior interval as 324 325 predictors for every time-frequency point during the overlapping interval. Figure 5A shows the slope 326 values representing the effect of context on the power of the different frequencies over time. We found a positive cluster in the window 0.48-0.84 s after interval onset in the 23-30 Hz frequency range 327 328 (p = .045), indicating increased beta power in the long context compared to the short context (see the 329 outlined area in Figure 5A). This beta effect is further illustrated in Figure 3C, which shows the average power in the 23-30 Hz over time, for the overlapping interval in the short and long context. 330 331 Figure 5B shows the slope values for prior interval, for which the permutation test indicated no 332 evidence for a cluster of slopes different from zero (ps > .051). In summary, these results suggest that 333 fronto-central beta power was higher in the long compared to the short context, while there was no 334 evidence for a similar influence of the previous trial.

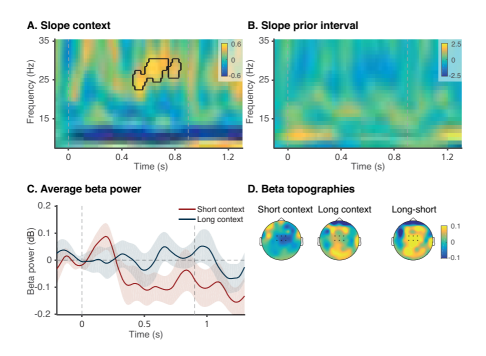


Figure 5. Slope values of regression on frequency power at fronto-central electrode cluster (electrodes Cz, C1, C2, FCz, FC1, FC2) relative to the onset of the perception phase. A) Slope values of the factor Context (short vs long) in the regression analysis at every time-frequency point. The outlined area marks a significant cluster according to the cluster-based permutation test performed in the time window 0-1.2 s and the frequency window 8-30 Hz. B) Slope values of the factor prior interval in the regression analysis predicting power. There was no evidence for significant clusters. In both panels, vertical dashed grey lines indicate the onset and offset of the perception phase. C) Average beta power in the time and frequency range of the significant cluster (23-30 Hz) for the short and long context for the overlapping interval. Error shading represents represent the standard error of the mean. D) Topographies of beta power for the overlapping interval, in the time and frequency range of the significant cluster (23-30 Hz) for the short context, the long context, and their difference.

CONTEXT ACTIVELY SHAPES TIME PERCEPTION

17

335 Linking EEG signatures and behavior

336 Figure 1C shows the effect of single-trial EEG signatures on reproductions of the overlapping interval. For illustration purposes, the single-trial EEG amplitudes and latencies were divided into 337 tertiles (low/short, medium, high/long) for each participant and context, and the average reproduction 338 339 was plotted for each tertile. The LMMs showed no evidence that single-trial CNV and beta power in the perception phase predicted reproductions in the reproduction phase ($\beta = 0.0003$, SE = 0.0004, t =340 0.85, p = .395 and $\beta = 0.0003, SE = 0.0019, t = -0.51, p = .614$, respectively). This was also the case 341 for P2 latency, with a trend towards shorter reproductions for later P2 peaks ($\beta = -0.08$, SE = 0.04, t =342 -1.80, p = .072). However, P2 amplitude after perception phase offset was predictive of that trial's 343 reproduction ($\beta = -0.0010$, SE = 0.0003, t = -3.45, p < .001). As the β -value indicates, higher P2 peaks 344 were followed by shorter reproductions. Given that context, interval and prior interval were also 345 346 included as fixed factors in the LMM, these results cannot be attributed to a mediating influence of 347 context, and therefore suggest that trial-by-trial variation in P2 amplitude might be a reliable predictor 348 of reproductions.

We additionally tested whether participants with a large behavioral context effect for the overlapping interval also showed a large context effect in the EEG signatures. This betweenparticipant relationship between these measures is depicted in Figure 1D. Analysis showed that the individual behavioral context effect was correlated with the P2 amplitude difference between contexts (r(23) = -.37, p = .033). We found no evidence for a similar relationship with P2 latency (r(23) = -.18, p = .196), CNV amplitude (r(23) = .19, p = .180) or beta power (r(23) = -.22, p = .861). Thus, in line with the single trial analysis, P2 amplitude differences predict reproduction outcomes.

356

Discussion

As the temporal locus of Bayesian computations in human time estimation is still unknown, we investigated whether temporal context actively influences neural signatures during the perception of time intervals. Behaviorally, we found that reproductions were biased towards the global temporal context as well as the duration in the previous trial. EEG results showed that CNV, P2 and beta power were modulated by previously perceived intervals, and that context could be decoded from transient

CONTEXT ACTIVELY SHAPES TIME PERCEPTION

18

362 brain dynamics at an early stage during perception. These results indicate that previously perceived 363 durations actively affect EEG signatures during interval estimation, showing that prior experiences act directly on perception. This observation goes against the (implicit) assumption of time perception 364 models that the likelihood is weighted with the prior only after perception. It is, however, in line with 365 366 recent behavioral evidence showing that context asserts its influence at early sensory stages (Cicchini et al., 2020; Zimmermann & Cicchini, 2020). Our findings suggest that experiences with the global 367 and recent temporal context actively calibrate cortical dynamics, in which the CNV and beta power 368 369 may reflect the anticipation of stimulus duration, and the P2 component the active evaluation of the 370 interval in the current context. Crucially, by focusing on the perception phase in a reproduction 371 paradigm, this is the first work demonstrating context effects that are not linked to explicit motor 372 preparation or response decisions.

373 Our findings argue against the idea that the CNV reflects the neural counterpart of the 374 absolute accumulator in pacemaker-accumulator models (Casini & Vidal, 2011; Macar & Besson, 375 1985; Macar & Vidal, 2004; Macar, Vidal, & Casini, 1999; Macar & Vitton, 1982; Pfeuty, Ragot, & 376 Pouthas, 2005), since no differences based on prior experience would be expected during the 377 perception of an interval. Instead, we found that the CNV during the perception of the overlapping 378 interval was more negative for the short compared to the long context, and for shorter previous 379 durations. This is consistent with anticipation and preparation accounts of the CNV (e.g., Boehm et 380 al., 2014; Elbert, 1993; Leuthold, Sommer, & Ulrich, 2004; Mento, 2013; Ng et al., 2011; Scheibe, 381 Schubert, Sommer, & Heekeren, 2009) and pacemaker-accumulator models that propose adaptive 382 spike rate accumulation (Simen, Balci, deSouza, Cohen, & Holmes, 2011): When interval offset is 383 expected quickly after onset, CNV amplitude increases more rapidly. This adaptation is in line with 384 studies showing a faster CNV development for relatively short foreperiods (Miniussi, Wilding, Coull, 385 & Nobre, 1999; Müller-Gethmann, Ulrich, & Rinkenauer, 2003; Trillenberg, Verleger, Wascher, Wauschkuhn, & Wessel, 2000; Van der Lubbe, Los, Jaśkowski, & Verleger, 2004), shorter standard 386 387 durations in an interval comparison task (Pfeuty et al., 2005), and after adaptation to a shorter interval (Li, Chen, Xiao, Liu, & Huang, 2017). The contextual adjustment of the speed with which the CNV 388 389 develops suggests that neural populations in the supplementary motor area (SMA), which is typically

CONTEXT ACTIVELY SHAPES TIME PERCEPTION

19

390 associated with the CNV (e.g., Coull, Vidal, & Burle, 2016), can perform flexible temporal scaling 391 based on the temporal context (Remington, Egger, Narain, Wang, & Jazayeri, 2018; Remington, Narain, Hosseini, & Jazaveri, 2018; Sohn et al., 2019), even in the absence of explicit motor 392 393 preparation. The prior might calibrate the speed of neural dynamics through different initial states at 394 the onset of the perception phase (Remington, Egger, et al., 2018; Sohn et al., 2019), as our 395 multivariate pattern analysis showed that global context can be decoded from EEG dynamics 396 immediately after the onset of the perception phase. Although the precise onset of significant 397 decoding should be interpreted with caution since the moving window approach and low-pass filtering 398 could smear out the accuracy over time (Grootswagers, Wardle, & Carlson, 2017), these results 399 suggest that temporal context affects the instantaneous neural response to to-be-timed stimuli. 400 The active anticipation based on context was also indexed by the P2 component. Specifically, 401 P2 amplitude increased with longer current durations, suggesting that it reflects hazard-based 402 expectancy: the probability that the interval offset will occur, given that is has not yet occurred 403 (Nobre, Correa, & Coull, 2007). This in line with previous studies showing that longer ISIs increase 404 P2 amplitude (e.g., Pereira et al., 2014; Röder et al., 2000). Importantly, however, P2 amplitude 405 decreased with longer previous durations, showing that the expectations are updated to the current 406 temporal context, even on a trial-by-trial basis. These results complement previous studies showing 407 that temporal expectancy modulates ERP amplitude (e.g., Kononowicz & van Rijn, 2014; Li et al., 408 2017; Todorovic & de Lange, 2012; Todorovic, van Ede, Maris, & de Lange, 2011; Wacongne et al., 409 2011). Interestingly, P2 amplitude at perception phase offset predicted interval reproductions, and 410 participants' behavioral context effect correlated with their context-based P2 effect. The lack of an equivalent CNV-effect highlights the predictive quality of the P2 (Kononowicz & van Rijn, 2014; 411 Kruijne et al., 2021), and indicates that the neural state at the end of the perception phase sets the 412 413 speed of cortical dynamics during reproduction (Sohn et al., 2019). Global context additionally 414 influenced beta power, such that beta power was higher in the long compared to the short context, in line with effects of beta power in single trial analyses (Kononowicz & van Rijn, 2015). Although beta 415 416 power has been proposed to reflect motor inhibition (Alegre et al., 2004; Hwang, Ghuman, Manoach, 417 Jones, & Luna, 2014; Kononowicz & van Rijn, 2015; Kühn et al., 2004), and most studies on the link

CONTEXT ACTIVELY SHAPES TIME PERCEPTION

20

418 between beta power and timing have a strong motor component, our results suggest that synchronized 419 beta oscillations also play a role during interval perception after which no immediate motor response is required. This finding complements recent studies showing that the accuracy and precision of time 420 estimates depend on beta (Wiener et al., 2018) and alpha-beta coupling (Kononowicz, Sander, van 421 422 Rijn, & Van Wassenhove, 2020). Additionally, the current global context effect on beta is in line with 423 Wiener et al.'s finding that longer previous durations increased beta power in the current trial. It has 424 to be noted, however, that we found no evidence for similar sequential effects on beta. 425 Besides the auditory stimuli which participants had to time, the current paradigm also 426 consisted of visual stimuli that indicated the phase of the trial (i.e., perception or reproduction). The 427 general overestimation we found in the behavioral results might potentially be explained by the 428 integration of these visual stimuli in temporal estimation (Shi & Burr, 2016). Future studies might 429 look further into potential modality differences in contextual calibration and their neural

430 underpinnings (Rhodes, Seth, & Roseboom, 2018; Roach et al., 2017; Zimmermann & Cicchini,

431 2020). Furthermore, we found no significant decoding corresponding to the windows of CNV

432 differences. This can be explained by the specific decoding method we employed, which focused on

transient dynamics, filtering out the stable CNV activity by baselining within a moving window. In

434 addition, decoding might be especially sensitive to stimulus onset and offset, with accuracy peaking

435 shortly afterwards and slowly dropping as the neural synchronization declines (e.g., Wolff et al.,

436 2017, 2020).

433

A comparison to Wiener and Thompson (2015), who found a larger CNV amplitude for 437 438 *longer* prior durations, suggests that contextual ERP effects might be dependent on the specific 439 experimental paradigm. In contrast to our reproduction experiment, their bisection task requires an 440 active decision during perception, and the CNV has been shown to reflect this decision process by 441 deflecting or plateauing after the standard interval in memory has been reached (Macar & Vidal, 2004; Ng et al., 2011; Pfeuty, Ragot, & Pouthas, 2003). A similar explanation could account for the 442 different nature of our offset P2 effect compared to Kononowicz and van Rijn (2014), who found a V-443 shaped P2 amplitude attenuation in a temporal comparison task (but see Kruijne et al., 2021). This 444 445 pattern reflects active comparison to the standard interval, which is not applicable to the current

CONTEXT ACTIVELY SHAPES TIME PERCEPTION

21

446	reproduction paradigm. In addition, the P2 measured in the current study shows similarities to the
447	positive offset peak named the late positive component of timing (LPCt) (Gontier et al., 2009; Paul et
448	al., 2011; Wiener & Thompson, 2015), although it has been argued that the P2 reflects perceptual
449	predictive processes while the LPCt indexes decision making (Kononowicz, van Rijn, & Meck, 2016).
450	The extent to which these components indeed reflect similar processes is still an open question, and
451	their occurrence seems to depend on the specific nature of the task. Future studies might directly
452	compare these neural differences in paradigms involving decision, motor or only perceptual timing
453	requirements.
454	In conclusion, our results show that previous durations actively influence flexible neural
455	dynamics during temporal encoding. These findings indicate that previous experiences in memory
456	create expectations that in turn calibrate our perception of the environment. The adaptive influence of
457	prior knowledge on perception could represent a more general Bayesian mechanism of magnitude
458	estimation (Petzschner, Glasauer, & Stephan, 2015), falsifying a class of models that assume discrete,
459	post-perceptual stages in which previous experiences exert their influence.

22

CONTEXT ACTIVELY SHAPES TIME PERCEPTION

460	Acknowledgements
461	This research has been partially supported by the EU Horizon 2020 FET Proactive grant
462	TIMESTORM - Mind and Time: Investigation of the Temporal Traits of Human-Machine
463	Convergence (Grant number 641100) and by the research programme Interval Timing in the Real
464	World financed by the Netherlands Organisation for Scientific Research (NWO, Grant number 453-
465	16-005, awarded to Hedderik van Rijn). The funding agencies had no involvement in the design of the
466	study, the analysis of the data, writing of the report, or in the decision to submit the article for
467	publication. We thank Sarah Maaß for fruitful discussions and programming the experiment, Ronja
468	Eike for her help in data collection and preprocessing, Emil Uffelmann for his help in data collection,
469	and Michael Wolff and Güven Kandemir for sharing their decoding knowledge with us.

470

CONTEXT ACTIVELY SHAPES TIME PERCEPTION

23

471

References

- Acerbi, L., Wolpert, D. M., & Vijayakumar, S. (2012). Internal Representations of Temporal Statistics
 and Feedback Calibrate Motor-Sensory Interval Timing. *PLoS Computational Biology*, 8(11).
 https://doi.org/10.1371/journal.pcbi.1002771
- Alegre, M., De Gurtubay, I. G., Labarga, A., Iriarte, J., Malanda, A., & Artieda, J. (2004). Alpha and
 beta oscillatory activity during a sequence of two movements. *Clinical Neurophysiology*, *115*(1),
 124–130. https://doi.org/10.1016/S1388-2457(03)00311-0
- Bates, D., Mächler, M., Bolker, B. M., & Walker, S. C. (2015). Fitting linear mixed-effects models
 using lme4. *Journal of Statistical Software*, 67(1). https://doi.org/10.18637/jss.v067.i01
- Boehm, U., Van Maanen, L., Forstmann, B., & van Rijn, H. (2014). Trial-by-trial fluctuations in CNV
 amplitude reflect anticipatory adjustment of response caution. *NeuroImage*, *96*, 95–105.
 https://doi.org/10.1016/j.neuroimage.2014.03.063
- 483 Brainard, D. H. (1997). The Psychophysics Toolbox. *Spatial Vision*, *10*(4), 433–436.
 484 https://doi.org/10.1163/156856897X00357
- Casini, L., & Vidal, F. (2011). The SMAs: Neural substrate of the temporal accumulator? *Frontiers in Integrative Neuroscience*, 5, 1–3. https://doi.org/10.3389/fnint.2011.00035
- 487 Cicchini, G. M., Arrighi, R., Cecchetti, L., Giusti, M., & Burr, D. C. (2012). Optimal encoding of
 488 interval timing in expert percussionists. *Journal of Neuroscience*, *32*(3), 1056–1060.
 489 https://doi.org/10.1523/JNEUROSCI.3411-11.2012
- 490 Cicchini, G. M., Benedetto, A., & Burr, D. C. (2020). Perceptual history propagates down to early
 491 levels of sensory analysis. *Current Biology*. https://doi.org/10.1016/j.cub.2020.12.004
- 492 Cicchini, G. M., Mikellidou, K., & Burr, D. (2017). Serial dependencies act directly on perception.
 493 *Journal of Vision*, 17(14). https://doi.org/10.1167/17.14.6
- 494 Coull, J. T., Vidal, F., & Burle, B. (2016). When to act, or not to act: That's the SMA's question.
 495 *Current Opinion in Behavioral Sciences*, Vol. 8, pp. 14–21.
 496 https://doi.org/10.1016/j.cobeha.2016.01.003
- 497 De Maesschalck, R., Jouan-Rimbaud, D., & Massart, D. L. (2000). The Mahalanobis distance.
 498 *Chemometrics and Intelligent Laboratory Systems*, 50(1), 1–18. https://doi.org/10.1016/S0169 499 7439(99)00047-7
- Di Luca, M., & Rhodes, D. (2016). Optimal Perceived Timing: Integrating Sensory Information with
 Dynamically Updated Expectations. *Scientific Reports*, 6, 1–15.
 https://doi.org/10.1038/srep28563
- 503 Dyjas, O., Bausenhart, K. M., & Ulrich, R. (2012). Trial-by-trial updating of an internal reference in
 504 discrimination tasks: Evidence from effects of stimulus order and trial sequence. *Attention*,
 505 *Perception, and Psychophysics*, 74(8), 1819–1841. https://doi.org/10.3758/s13414-012-0362-4
- Elbert, T. (1993). Slow Cortical Potentials Reflect the Regulation of Cortical Excitability. In *Slow Potential Changes in the Human Brain* (pp. 235–251). https://doi.org/10.1007/978-1-4899-1597 9_15
- Gontier, E., Paul, I., Le Dantec, C., Pouthas, V., Jean-Marie, G., Bernard, C., ... Rebaï, M. (2009).
 ERPs in Anterior and Posterior Regions Associated With Duration and Size Discriminations.
 Neuropsychology, 23(5), 668–678. https://doi.org/10.1037/a0015757
- Grootswagers, T., Wardle, S. G., & Carlson, T. A. (2017). Decoding dynamic brain patterns from
 evoked responses: A tutorial on multivariate pattern analysis applied to time series neuroimaging
 data. *Journal of Cognitive Neuroscience*, 29(4), 677–697. https://doi.org/10.1162/jocn a 01068
- 515 Gu, B. M., Jurkowski, A. J., Lake, J. I., Malapani, C., & Meck, W. H. (2015). Bayesian Models of

CONTEXT ACTIVELY SHAPES TIME PERCEPTION

24

- 516 Interval Timing and Distortions in Temporal Memory as a Function of Parkinson's Disease and 517 Dopamine-Related Error Processing. In *Time Distortions in Mind* (pp. 281–327).
- 518 https://doi.org/10.1163/9789004230699 012
- Haegens, S., Nácher, V., Hernández, A., Luna, R., Jensen, O., & Romo, R. (2011). Beta oscillations in
 the monkey sensorimotor network reflect somatosensory decision making. *Proceedings of the National Academy of Sciences of the United States of America*, 108(26), 10708–10713.
 https://doi.org/10.1073/pnas.1107297108
- Hallez, Q., Damsma, A., Rhodes, D., van Rijn, H., & Droit-Volet, S. (2019). The dynamic effect of
 context on interval timing in children and adults. *Acta Psychologica*, *192*, 87–93.
 https://doi.org/10.1016/j.actpsy.2018.10.004
- Hwang, K., Ghuman, A. S., Manoach, D. S., Jones, S. R., & Luna, B. (2014). Cortical neurodynamics
 of inhibitory control. *Journal of Neuroscience*, *34*(29), 9551–9561.
 https://doi.org/10.1523/JNEUROSCI.4889-13.2014
- Jazayeri, M., & Shadlen, M. N. (2010). Temporal context calibrates interval timing. *Nature Neuroscience*, 13(8), 1020–1026. https://doi.org/10.1038/nn.2590
- Jenkinson, N., & Brown, P. (2011). New insights into the relationship between dopamine, beta
 oscillations and motor function. *Trends in Neurosciences*, *34*(12), 611–618.
 https://doi.org/10.1016/j.tins.2011.09.003
- Kleiner, M., Brainard, D. H., Pelli, D. G., Broussard, C., Wolf, T., & Niehorster, D. (2007). What's
 new in Psychtoolbox-3? *Perception*, *36*, S14. https://doi.org/10.1068/v070821
- Kononowicz, T. W., Sander, T., Van Rijn, H., & Van Wassenhove, V. (2020). Precision timing with
 α-β oscillatory coupling: Stopwatch or motor control? *Journal of Cognitive Neuroscience*,
 32(9), 1624–1636. https://doi.org/10.1162/jocn_a_01570
- Kononowicz, T. W., & van Rijn, H. (2014). Decoupling interval timing and climbing neural activity:
 A dissociation between CNV and N1P2 amplitudes. *Journal of Neuroscience*, *34*(8), 2931–2939.
 https://doi.org/10.1523/JNEUROSCI.2523-13.2014
- Kononowicz, T. W., & van Rijn, H. (2015). Single trial beta oscillations index time estimation.
 Neuropsychologia, 75, 381–389. https://doi.org/10.1016/j.neuropsychologia.2015.06.014
- Kononowicz, T. W., van Rijn, H., & Meck, W. H. (2016). Timing and time perception: a critical
 review of neural timing signatures before, during, and after the To-Be-Timed Interval. In *Stevens Handbook of Experimental Psychology and Cognitive Neuroscience (4th ed.).*
- 547 Kruijne, W., Olivers, C. N. L., & van Rijn, H. (2021). Neural repetition suppression modulates time
 548 perception: Evidence from electrophysiology and pupillometry. *BioRxiv*, 2020.07.31.230508.
 549 https://doi.org/10.1101/2020.07.31.230508
- Kühn, A. A., Williams, D., Kupsch, A., Limousin, P., Hariz, M., Schneider, G. H., ... Brown, P.
 (2004). Event-related beta desynchronization in human subthalamic nucleus correlates with
 motor performance. *Brain*, *127*(4), 735–746. https://doi.org/10.1093/brain/awh106
- Ledoit, O., & Wolf, M. (2004). Honey, I shrunk the sample covariance matrix. *Journal of Portfolio Management*, 30(4), 110–119. https://doi.org/10.3905/jpm.2004.110
- Lemm, S., Blankertz, B., Dickhaus, T., & Müller, K. R. (2011). Introduction to machine learning for
 brain imaging. *NeuroImage*, 56(2), 387–399. https://doi.org/10.1016/j.neuroimage.2010.11.004
- Leuthold, H., Sommer, W., & Ulrich, R. (2004). Preparing for action: Inferences from CNV and LRP.
 Journal of Psychophysiology, 18(2–3), 77–88. https://doi.org/10.1027/0269-8803.18.23.77
- Li, B., Chen, Y., Xiao, L., Liu, P., & Huang, X. (2017). Duration adaptation modulates EEG
 correlates of subsequent temporal encoding. *NeuroImage*, 147(2), 143–151.
- 561 https://doi.org/10.1016/j.neuroimage.2016.12.015

CONTEXT ACTIVELY SHAPES TIME PERCEPTION

- Liesefeld, H. R. (2018). Estimating the Timing of Cognitive Operations With MEG/EEG Latency
 Measures: A Primer, a Brief Tutorial, and an Implementation of Various Methods. *Frontiers in Neuroscience*, *12*. https://doi.org/10.3389/fnins.2018.00765
- Luck, S. J. (2005). An Introduction to Event-Related Potentials and Their Neural Origins. An
 Introduction to the Event-Related Potential Technique.
- Maaß, S. C., Riemer, M., Wolbers, T., & van Rijn, H. (2019). Timing deficiencies in amnestic Mild
 Cognitive Impairment: Disentangling clock and memory processes. *Behavioural Brain Research*, 373, 112110. https://doi.org/10.1016/j.bbr.2019.112110
- Maaß, S. C., Schlichting, N., & van Rijn, H. (2019). Eliciting contextual temporal calibration: The
 effect of bottom-up and top-down information in reproduction tasks. *Acta Psychologica*, *199*,
 102898. https://doi.org/10.1016/j.actpsy.2019.102898
- Macar, F., & Besson, M. (1985). Contingent negative variation in processes of expectancy, motor
 preparation and time estimation. *Biological Psychology*, *21*(4), 293–307.
 https://doi.org/10.1016/0301-0511(85)90184-X
- Macar, F., & Vidal, F. (2004). Event-related potentials as indices of time processing: A review.
 Journal of Psychophysiology, Vol. 18, pp. 89–104. https://doi.org/10.1027/0269-8803.18.23.89
- Macar, F., Vidal, F., & Casini, L. (1999). The supplementary motor area in motor and sensory timing:
 Evidence from slow brain potential changes. *Experimental Brain Research*, *125*(3), 271–280.
 https://doi.org/10.1007/s002210050683
- Macar, F., & Vitton, N. (1982). An early resolution of contingent negative variation (CNV) in time
 discrimination. *Electroencephalography and Clinical Neurophysiology*, 54(4), 426–435.
 https://doi.org/10.1016/0013-4694(82)90206-1
- Maris, E., & Oostenveld, R. (2007). Nonparametric statistical testing of EEG- and MEG-data. *Journal of Neuroscience Methods*, *164*(1), 177–190. https://doi.org/10.1016/j.jneumeth.2007.03.024
- Mento, G. (2013). The passive CNV: Carving out the contribution of task-related processes to
 expectancy. *Frontiers in Human Neuroscience*, 7, 827.
 https://doi.org/10.3389/fnhum.2013.00827
- 589 Miniussi, C., Wilding, E. L., Coull, J. T., & Nobre, A. C. (1999). Orienting attention in time.
 590 Modulation of brain potentials. *Brain*, 122(8), 1507–1518.
 591 https://doi.org/10.1093/brain/122.8.1507
- Müller-Gethmann, H., Ulrich, R., & Rinkenauer, G. (2003). Locus of the effect of temporal
 preparation: Evidence from the lateralized readiness potential. *Psychophysiology*, 40(4), 597–
 611. https://doi.org/10.1111/1469-8986.00061
- Ng, K. K., Tobin, S., & Penney, T. B. (2011). Temporal accumulation and decision processes in the
 duration bisection task revealed by contingent negative variation. *Frontiers in Integrative Neuroscience*, 5, 77. https://doi.org/10.3389/fnint.2011.00077
- Nobre, A. C., Correa, A., & Coull, J. (2007). The hazards of time. *Current Opinion in Neurobiology*, 17(4), 465–470. https://doi.org/10.1016/j.conb.2007.07.006
- Oostenveld, R., Fries, P., Maris, E., & Schoffelen, J. M. (2011). FieldTrip: Open source software for
 advanced analysis of MEG, EEG, and invasive electrophysiological data. *Computational Intelligence and Neuroscience*, 2011. https://doi.org/10.1155/2011/156869
- Paul, I., Wearden, J., Bannier, D., Gontier, E., Le Dantec, C., & Rebaï, M. (2011). Making decisions
 about time: Event-related potentials and judgements about the equality of durations. *Biological Psychology*, 88(1), 94–103. https://doi.org/10.1016/j.biopsycho.2011.06.013
- Pereira, D. R., Cardoso, S., Ferreira-Santos, F., Fernandes, C., Cunha-Reis, C., Paiva, T. O., ...
 Marques-Teixeira, J. (2014). Effects of inter-stimulus interval (ISI) duration on the N1 and P2

CONTEXT ACTIVELY SHAPES TIME PERCEPTION

26

- 608 components of the auditory event-related potential. *International Journal of Psychophysiology*,
 609 94(3), 311–318. https://doi.org/10.1016/j.ijpsycho.2014.09.012
- Petzschner, F. H., Glasauer, S., & Stephan, K. E. (2015). A Bayesian perspective on magnitude
 estimation. *Trends in Cognitive Sciences*, 19(5), 285–293.
 https://doi.org/10.1016/j.tics.2015.03.002
- Pfeuty, M., Ragot, R., & Pouthas, V. (2003). When time is up: CNV time course differentiates the
 roles of the hemispheres in the discrimination of short tone durations. *Experimental Brain Research*, 151(3), 372–379. https://doi.org/10.1007/s00221-003-1505-6
- Pfeuty, M., Ragot, R., & Pouthas, V. (2005). Relationship between CNV and timing of an upcoming
 event. *Neuroscience Letters*, 382(1–2), 106–111. https://doi.org/10.1016/j.neulet.2005.02.067
- R Core Development Team. (2013). A language and environment for statistical computing. *R Foundation for Statistical Computing*, Vol. 1. Retrieved from http://www.r-project.org/
- Remington, E. D., Egger, S. W., Narain, D., Wang, J., & Jazayeri, M. (2018). A Dynamical Systems
 Perspective on Flexible Motor Timing. *Trends in Cognitive Sciences*, 22(10), 938–952.
 https://doi.org/10.1016/j.tics.2018.07.010
- Remington, E. D., Narain, D., Hosseini, E. A., & Jazayeri, M. (2018). Flexible Sensorimotor
 Computations through Rapid Reconfiguration of Cortical Dynamics. *Neuron*, 98(5), 1005 1019.e5. https://doi.org/10.1016/j.neuron.2018.05.020
- Rhodes, D., Seth, A. K., & Roseboom, W. (2018). Multiple Duration Priors Within and Across the
 Senses. *BioRxiv*, 467027. https://doi.org/10.1101/467027
- Roach, N. W., McGraw, P. V., Whitaker, D. J., & Heron, J. (2017). Generalization of prior
 information for rapid Bayesian time estimation. *Proceedings of the National Academy of Sciences of the United States of America*, 114(2), 412–417.
 https://doi.org/10.1073/pnas.1610706114
- Röder, B., Rösler, F., & Neville, H. J. (2000). Event-related potentials during auditory language
 processing in congenitally blind and sighted people. *Neuropsychologia*, *38*(11), 1482–1502.
 https://doi.org/10.1016/S0028-3932(00)00057-9
- Scheibe, C., Schubert, R., Sommer, W., & Heekeren, H. R. (2009). Electrophysiological evidence for
 the effect of prior probability on response preparation. *Psychophysiology*, *46*(4), 758–770.
 https://doi.org/10.1111/j.1469-8986.2009.00825.x
- Schlichting, N., Damsma, A., Aksoy, E. E., Wächter, M., Asfour, T., & van Rijn, H. (2018). Temporal
 Context Influences the Perceived Duration of Everyday Actions: Assessing the Ecological
 Validity of Lab-Based Timing Phenomena. *Journal of Cognition*, 2(1).
 https://doi.org/10.5334/joc.4
- Shi, Z., & Burr, D. (2016). Predictive coding of multisensory timing. *Current Opinion in Behavioral Sciences*, 8, 200–206. https://doi.org/10.1016/j.cobeha.2016.02.014
- Shi, Z., Church, R. M., & Meck, W. H. (2013). Bayesian optimization of time perception. *Trends in Cognitive Sciences*, 17(11), 556–564. https://doi.org/10.1016/j.tics.2013.09.009
- 646 Simen, P., Balci, F., deSouza, L., Cohen, J. D., & Holmes, P. (2011). A model of interval timing by
 647 neural integration. *Journal of Neuroscience*, *31*(25), 9238–9253.
 648 https://doi.org/10.1523/JNEUROSCI.3121-10.2011
- Sohn, H., Narain, D., Meirhaeghe, N., & Jazayeri, M. (2019). Bayesian Computation through Cortical
 Latent Dynamics. *Neuron*, 103(5), 934-947.e5. https://doi.org/10.1016/j.neuron.2019.06.012
- St. John-Saaltink, E., Kok, P., Lau, H. C., & De Lange, F. P. (2016). Serial dependence in perceptual
 decisions is reflected in activity patterns in primary visual cortex. *Journal of Neuroscience*, *36*(23), 6186–6192. https://doi.org/10.1523/JNEUROSCI.4390-15.2016

CONTEXT ACTIVELY SHAPES TIME PERCEPTION

- Taatgen, N. A., & van Rijn, H. (2011). Traces of times past: Representations of temporal intervals in memory. *Memory and Cognition*, 39(8), 1546–1560. https://doi.org/10.3758/s13421-011-0113-0
- Tanner, D., Morgan-Short, K., & Luck, S. J. (2015). How inappropriate high-pass filters can produce
 artifactual effects and incorrect conclusions in ERP studies of language and cognition.
 Psychophysiology, 52(8), 997–1009. https://doi.org/10.1111/psyp.12437
- Todorovic, A., & de Lange, F. P. (2012). Repetition suppression and expectation suppression are
 dissociable in time in early auditory evoked fields. *Journal of Neuroscience*, *32*(39), 13389–
 13395. https://doi.org/10.1523/JNEUROSCI.2227-12.2012
- Todorovic, A., van Ede, F., Maris, E., & de Lange, F. P. (2011). Prior expectation mediates neural
 adaptation to repeated sounds in the auditory cortex: An MEG study. *Journal of Neuroscience*, *31*(25), 9118–9123. https://doi.org/10.1523/JNEUROSCI.1425-11.2011
- Trillenberg, P., Verleger, R., Wascher, E., Wauschkuhn, B., & Wessel, K. (2000). CNV and temporal
 uncertainty with "ageing" and "non-ageing" S1-S2 intervals. *Clinical Neurophysiology*, *111*(7),
 1216–1226. https://doi.org/10.1016/S1388-2457(00)00274-1
- Van der Lubbe, R. H. J., Los, S. A., Jaśkowski, P., & Verleger, R. (2004). Being prepared on time: On
 the importance of the previous foreperiod to current preparation, as reflected in speed, force and
 preparation-related brain potentials. *Acta Psychologica*, *116*(3), 245–262.
 https://doi.org/10.1016/j.actpsy.2004.03.003
- Wacongne, C., Labyt, E., Van Wassenhove, V., Bekinschtein, T., Naccache, L., & Dehaene, S.
 (2011). Evidence for a hierarchy of predictions and prediction errors in human cortex. *Proceedings of the National Academy of Sciences of the United States of America*, 108(51),
 20754–20759. https://doi.org/10.1073/pnas.1117807108
- Wiener, M., Parikh, A., Krakow, A., & Coslett, H. B. (2018). An intrinsic role of beta oscillations in memory for time estimation. *Scientific Reports*, 8(1), 1–17. https://doi.org/10.1038/s41598-018-26385-6
- Wiener, M., & Thompson, J. C. (2015). Repetition enhancement and memory effects for duration.
 NeuroImage, *113*, 268–278. https://doi.org/10.1016/j.neuroimage.2015.03.054
- Wiener, M., Thompson, J. C., & Branch Coslett, H. (2014). Continuous carryover of temporal context
 dissociates response bias from perceptual influence for duration. *PLoS ONE*, *9*(6).
 https://doi.org/10.1371/journal.pone.0100803
- Wolff, M. J., Jochim, J., Akyürek, E. G., & Stokes, M. G. (2017). Dynamic hidden states underlying
 working-memory-guided behavior. *Nature Neuroscience*, 20(6), 864–871.
 https://doi.org/10.1038/nn.4546
- Wolff, M. J., Kandemir, G., Stokes, M. G., & Akyürek, E. G. (2020). Unimodal and Bimodal Access
 to Sensory Working Memories by Auditory and Visual Impulses. *The Journal of Neuroscience : The Official Journal of the Society for Neuroscience*, 40(3), 671–681.
 https://doi.org/10.1523/JNEUROSCI.1194-19.2019
- Zimmermann, E., & Cicchini, G. M. (2020). Temporal Context affects interval timing at the
 perceptual level. *Scientific Reports*. https://doi.org/10.1038/s41598-020-65609-6
- 693