Encoding of task regularities links grid-like signals to human timing behavior

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Abstract

Grid cells have been proposed to encode task regularities that allow predicting future states. Entorhinal grid-like signals might therefore mirror behavioral biases associated with relying on task regularities, like regression-to-the-mean biases in time estimation. Here, we tested this proposal using functional magnetic resonance imaging and a rapid timing task in humans. Indeed, trial-wise entorhinal activity reflected task accuracy and the degree to which interval estimates regressed towards the mean of all tested intervals. Grid-like signals were observed exclusively for the interval closest to the mean, which was explained by differences in temporal stability across intervals. Finally, both behavioral and entorhinal results were explained by a Bayesian observer model that assumes the integration of current-trial sensory evidence with prior expectations. Together, we find that entorhinal activity and grid-like signals reflect behavioral performance in a timing task, supporting the proposed role of grid cells in encoding task structure for predictive coordination of behavior.

1 Introduction

² The ability to recognize and utilize statistical regularities governed by the co-occurrence of stimuli,

actions, and events is crucial for any successful interaction with the environment (Vetter, Wolpert,

4 2000; Clark, 2013; Friston, Buzsáki, 2016). Learning about such regularities allows us to predict

5 future states of the world, for instance to track moving objects during occlusion, which is an es-

⁶ sential ability underlying flexible and robust behavior broadly (Fiser et al., 2010; Heald et al., 2023;

⁷ Schapiro, Turk-Browne, 2015; Schapiro et al., 2016). When catching an approaching ball, for ex-

⁸ ample, we anticipate the moment it will reach us not solely based on our estimates of its current

⁹ speed and distance, but also based on our knowledge of how previous balls have behaved in simi-

¹⁰ lar situations. Through experience, we have learned about the probability associated with certain

speeds and arrival times, which now guides when and how we act. How does the brain encode such statistical regularities and how do they afford predictive inference in the service of behavior?

¹³ A prominent metaphor of how the brain encodes statistical regularities can be found in the con-¹⁴ cept of cognitive maps, referring to relational map-like representations of places and events that

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support mnemonic and predictive processes (O'Keefe, Nadel, 1978; Moser et al., 2014; Stachenfeld
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¹⁷ erature on cognitive maps and predictions pointed to the hippocampal formation as a key neural

component involved in both, fueling efforts to unify theories of its contribution to a range of tasks
 (Whittington et al., 2020; Stachenfeld et al., 2017; Ambrogioni, Ólafsdóttir, 2023; Behrens et al.,

2018: Bellmund et al., 2018). An emerging view is that the hippocampal formation plays an impor-

tant role in encoding regularities that afford generalization across tasks or contexts (e.g., different

environments; Whittington et al. (2020); Bousquet et al. (1998); Fuhs, Touretzky (2007); Penny et al.

²³ (2013); Friston, Buzsáki (2016); Pezzulo et al. (2017)), thus greatly accelerating learning and reduc-

²⁴ ing behavioral errors in novel or noisy situations (Lisman, Redish, 2009; Stachenfeld et al., 2017).

²⁵ In particular, the hippocampus has been suggested to support the encoding of task regularities in

real time as a task is performed, as its activity reflects feedback and behavioral improvements even

²⁷ in fast-paced timing tasks (Polti et al., 2022).

Like the hippocampus, the entorhinal cortex is widely considered critical for cognitive mapping; not 28 only is it a major anatomical gateway for hippocampal-cortical interactions (Witter, Amaral, 2004), 29 but it also harbors grid cells (Hafting et al., 2005; Moser et al., 2014) proposed to provide a map-like 30 coordinate system useful for representing relationships between places and events in the world 31 (e.g., different spatial positions during navigation, but also non-spatial features (Constantinescu 32 et al., 2016; Aronov et al., 2017; Behrens et al., 2018; Bellmund et al., 2018; Bao et al., 2019; Theves 33 et al., 2019, 2020; Viganò, Piazza, 2020; Park et al., 2021). This grid-like coordinate system may be 34 predictive in nature, meaning that it likely anticipates future states of the agent (e.g., future po-35 sitions during navigation, Stachenfeld et al. (2017)) and affords efficient vector computations for 36 spatial planning (Banino et al., 2018; Bush et al., 2015; Bicanski, Burgess, 2020). At the population 37 level, grid cell activity is believed to exhibit a hexadirectional (i.e., six-fold rotationally symmetric) 38 modulation as a function of virtual running or gaze direction (Killian et al., 2012), which can be ob-39 served in the human entorhinal cortex using functional magnetic resonance imaging (fMRI) (Doeller 40 et al., 2010; Julian et al., 2018; Nau et al., 2018b). Importantly, while grid-like signals have been ob-41 served in a range of tasks and species (Kunz et al., 2019), most studies left open whether or not 42

43 grid-like signals were indeed relevant for task performance.

⁴⁴ Here, we set out to understand the contributions of the entorhinal cortex and grid-like signals in ⁴⁵ particular to the encoding of task regularities that afford predictive coordination of actions relative

to sensory events. In addition, we investigated the relationship between grid-like signals and task 46 performance - a relationship that is often assumed but remains to be demonstrated empirically. 47 For this purpose, we monitored human brain activity using fMRI while participants engaged in a 48 visual tracking task previously shown to engage grid-like signals in the human entorhinal cortex 49 (Nau et al., 2018b) as well as a time-to-contact (TTC) estimation task shown to engage the adjacent 50 hippocampus in a rapid and behavior-dependent manner (Polti et al., 2022). We specifically tested 51 whether and how entorhinal activity reflects behavioral biases previously linked to the encoding 52 of timing-task regularities (Jazayeri, Shadlen, 2010), focusing on the posteromedial portion of the 53 entorhinal cortex (pmEC); the presumed human homologue of the rodent medial entorhinal cortex 54 that harbours grid cells (Navarro Schröder et al., 2015; Maass et al., 2015; Syversen et al., 2021). 55

56 **Results**

In the following, we will present our experiment and results in 5 steps. First, we introduce our task 57 design and empirical measures in detail. Second, we show that task performance depended not 58 only on the interval that was tested in each trial, but also on the intervals tested in previous trials, 59 with behavioral responses showing a systematic bias towards the average interval. This regression-60 to-the-mean bias suggests that participants relied on prior knowledge that accumulated across tri-61 als. Third, we report that trial-wise pmEC activity mirrored this behavioral bias in real time, similar 62 to previous reports for the hippocampus (Polti et al., 2022). Fourth, we show that human pmEC 63 grid-like signals co-varied with the tested intervals across trials. This effect was explained by the 64 temporal stability of the grid-like signal (i.e. replicability across data partitions), and its amplitude 65 was correlated with behavioral performance in all participants. Finally, by showing that a Bayesian 66 observer model provides a parsimonious account of the data, we illustrate a potential computa-67 tional explanation of our results, namely that entorhinal grid-like signals may reflect the mismatch 68 between prior knowledge and sensory evidence obtained in each trial. Collectively, these results 69 suggest that non-spatial task factors (such as tested intervals) shape spatial representations in the 70 entorhinal cortex in service of timing behavior, and that entorhinal activity, and grid-like signals in 71 particular, reflect the rapid encoding of task regularities in service of predictive inference. More-72 over, our results also provide evidence for a link between grid-like signals and task performance. 73

74 Time-to-contact (TTC) estimation task

Our task consisted of two components; a visual tracking task that engages grid-like signals in the 75 human EC (Nau et al., 2018b), as well as a predictive timing task that engages other regions includ-76 ing the adjacent hippocampus (Polti et al., 2022). Over the course of 768 trials, participants tracked 77 a moving fixation target with their eyes until it was occluded, which then prompted them to predict 78 when the target would hit a visual boundary. In each trial, the fixation target moved 10 degrees of 79 visual angle (dva) into one of 24 directions (Fig. 1A, "Gaze trajectory") at one of 4 possible speeds, 80 yielding 4 different intervals to be estimated (t_{TTC}): 0.55, 0.65, 0.86, and 1.2 s (See *Methods*). Speed 81 and direction were held constant within each trial. After the target stopped moving, participants 82 estimated the time when it would have hit the visual boundary 5 dva apart, which they indicated via 83 button click (Fig. 1A, "TTC estimation"). Participants then received feedback reflecting the accuracy 84 of their estimated TTC relative to the ground-truth TTC. The next trial then started after a jittered 85 inter-trial interval (ITI). See the methods section for details. 86



A) Visual tracking & time-to-contact estimation task



Figure 1: A) Visual tracking and Time-To-Contact (TTC) estimation task. In each trial during fMRI scanning, participants fixated on a target (1), which started moving at one of 4 possible speeds and in one of 24 possible directions for 10 dva (2). After the target stopped moving, participants kept fixating and estimated when the fixation target would have hit a boundary 5 dva apart (3). After pressing a button at the estimated TTC, participants received feedback (4) based on their performance. B) Task performance. Root-mean-square error (RMSE) differences across TTCs show a quadratic pattern, i.e. target TTCs closer to the mean of the sampled TTCs (vertical dashed line) have a lower RMSE. We plot the mean and SEM (black dot and line). C) Regression effect. Participants responses regressed towards the mean of the sampled TTCs (0.82, horizontal dashed line), away from the identity line (diagonal dashed line). Regression line (black) and standard error (gray shade). BC) Single-participant data plotted as dots. Target TTCs are color coded.

87 Behavioral results

To examine whether participants performed the task well, we first compared their estimated TTC 88 to the target TTC (ground truth) using a mixed-effects model (MEM). Indeed, we found that the 89 estimated and target TTCs were tightly correlated (Fig. 1C; F(1) = 976.44, $p = 2.2x10^{-16}$, $\epsilon^2 = 0.91$, CI: 90 [0.88, 1]). However, TTC estimates were further systematically biased towards the mean of the 91 tested intervals (0.82 s, Fig. 1C, horizontal dashed line) in line with previous reports using inter-92 val timing tasks (Miyazaki et al., 2005; Jazayeri, Shadlen, 2010; Acerbi et al., 2012; Cicchini et al., 93 2012; Chang, Jazayeri, 2018). We quantified this regression-to-the-mean bias by fitting a line to the 94 TTC estimates on group-level, which resulted in a slope value of 0.53 (Fig. 1C, MEM fit, diagonal 95 solid line). For comparison, perfect task performance would lead to a slope of 1, whereas total 96 regression to the mean would result in a slope of 0. 97

To quantify task performance in more detail, we then calculated the precision and accuracy in TTC 98 estimation for each participant and time interval. Precision describes how similar the estimates 99 were across trials, whereas accuracy describes how similar they were to the ground-truth target 100 TTC (see methods for details). Together, these two measures combine into the root-mean-square 101 error (RMSE), which we computed as our final performance measure. Note that a given RMSE can 102 be the result of different precision-accuracy trade-offs (Fig. S1A). We found that the RMSE showed 103 a quadratic relationship to the t_{TTC}, meaning that lower RMSE's were observed for t_{TTC} closer to 104 the mean of the tested intervals (Fig. 1B; MEM, F(2) = 39.18, $p = 1.5x10^{-9}$, $\epsilon^2 = 0.68$, CI : [0.52, 1]). This 105 quadratic trend explained the data better than assuming a linear trend (Chi-square test, $\chi^2(3) =$ 106 66.15, $p = 2.85 \times 10^{-14}$). 107

Taken together, our behavioral results showed that participants performed the task well (Fig. 1B, C),
 and that their TTC estimates exhibited systematic regression-to-the-mean biases (Fig. 1C). These
 biases are well documented in the literature and suggest that participants relied on prior knowl-

edge beyond the current trial to estimate time (Miyazaki et al., 2005; Jazayeri, Shadlen, 2010; Acerbi et al., 2012; Cicchini et al., 2012; Chang, Jazayeri, 2018; Meirhaeghe et al., 2021; Polti et al., 2022).

113 Entorhinal cortex activity reflects trial-wise accuracy and biases in TTC estimation

Previous work on the present data showed that activity in the hippocampus reflected the mag-114 nitude of the behavioral regression-to-the-mean effect across trials (Polti et al., 2022). Here, we 115 therefore first tested whether a similar effect can be observed for the pmEC. Using mass-univariate 116 general linear models (GLM), we modeled the activity in each trial parametrically either as a func-117 tion of accuracy (i.e., the absolute difference between estimated and target TTC; Fig. 2A, top) or as a 118 function of the magnitude of the regression effect (i.e., the absolute difference between estimated 119 TTC and the mean of the tested intervals; Fig. 2A, bottom). To avoid effects of potential colinearity 120 on the final parameter estimates, these two predictors were fit in two independent GLMs, which in-121 cluded additional nuisance predictors (e.g., for head-motion, see methods). We found that pmEC 122 activity was higher in trials in which TTC estimates were more accurate (Fig. 2B, left; two-tailed 123 one-sample Wilcoxon signed-rank test; $V = 89, p = 1.8 \times 10^{-4}, r = -0.70, CI : [-0.85, -0.45]$), but also 124 that it linearly increased with stronger regression-to-the-mean biases (Fig. 2B, right; two-tailed one-125 sample Wilcoxon signed-rank test; V = 157, p = 0.015, r = -0.47, CI : [-0.72, -0.13]), resembling the 126 effects previously reported for the hippocampus (Polti et al., 2022). 127



Figure 2: Posteromedial entorhinal cortex (pmEC) activity predicts trial-wise TTC behavior. Schematic description of the parametric regressor (PR) used in each separate GLM. The Accuracy PR (Top) contained the absolute difference between estimated TTC and the identity line for each trial (petrol diagonal dashed line), whereas the Regression effect PR (Bottom) contained the absolute difference between estimated TTC and mean of the tested TTCs (0.82, magenta horizontal dashed line). B) Independent regions-ofinterest analysis for pmEC. We plot the beta estimates obtained for each participant for each of the two regressors. Negative values indicate higher pmEC activity with either higher accuracy (left) or higher magnitude of the regression effect (right). Depicted are the mean and SEM across participants (black dot and line) overlaid on single participant data (colored dots). Statistics reflect p<0.05 (*) obtained using a group-level two-tailed one-sample Wilcoxon signed-rank test against zero.

128 Entorhinal grid-like signals predict behavioral performance in time estimation

The results reported above indicate that trial-wise pmEC activity was associated with the accuracy 129 and the bias in TTC estimation. However, these analyses do not address whether or not grid-like 130 signals show an association to our behavioral measures as well. Therefore, we next examined grid-131 like signals in our data separately for each t_{TTC} using an established quadrature filter approach 132 (Doeller et al., 2010). The aim of this analysis was to examine whether pmEC voxels exhibited 133 visual grid-like signals in our task (i.e., six-fold rotationally symmetric modulations as a function of 134 gaze direction (Nau et al., 2018b; Julian et al., 2018; Staudigl et al., 2018)), and if so, whether these 135 signals show a relationship to the regression-to-the-mean bias as well. 136

We split the time series of each pmEC voxel into two halves for later cross-validation, and then 137 modeled each of the halves separately using a voxel-wise GLM. The GLM included two parametric 138 main predictors modeling the sine and cosine of gaze direction modulo 60° (Fig. 3A) in addition to 139 nuisance regressors (see methods). The ratio between the beta estimates obtained for these two 140 predictors then allowed us to infer the putative grid orientation (i.e., the phase of the hexadirec-141 tional modulation) for each voxel, which were then averaged across voxels and across scanning 142 runs within each data partition. This procedure resulted in one putative grid-orientation for each 143 t_{TTC} and each data partition. Using a second GLM, we then tested the reliability of these grid-like 144 signals in the respective held-out data by modeling the activity of each voxel as a function of gaze di-145 rection aligned with the putative grid orientation modulo 60°. In other words, the predictor tested 146 whether MRI signals observed for gaze directions aligned with the putative grid orientation were 147 stronger than those observed for directions misaligned to it. 148

We found that pmEC indeed exhibited a reliable grid-like modulation in our task as indicated by 149 a regions-of-interest (ROI) analysis. Critically, however, we observed a main effect of t_{TTC} on the 150 amplitude of this grid-like modulation (Fig. 3B; MEM, F(3) = 3.08, p = 0.029, $\epsilon^2 = 0.04$, CI : [0, 1]), 151 with only one of the four tested intervals showing the effect ($TTC_{0.86}$, Fig. 3B; Table S1). Trials in 152 which $TTC_{0.86}$ was tested yielded a stronger grid-like signal than all other TTCs (Table S1). Note 153 that $TTC_{0.86}$ was the interval that was closest to the mean of all intervals. A whole-brain voxel-wise 154 analysis later confirmed that this grid-like modulation for $TTC_{0.86}$ occurred in both hemispheres in 155 the entorhinal cortex (Fig. 3C, left and middle panels) as well as in a few other regions that shared 156 the putative grid orientation with the pmEC (e.g., the pre-supplementary motor area (preSMA), Fig. 157 3C, right panel; see Table S2 for post-hoc ROI-analysis). No effect was observed in pmEC when the 158 same cross-validation analysis was repeated for other directional periodicities such as 90° (Fig. 3D. 159 Left; Table S3A) and 45° (Fig. 3D, Center; Table S3B). Furthermore, as expected based on previous 160 work Nau et al. (2018b), the early visual cortex did not exhibit a grid-like signal (Fig. 3D, Right; Table 161 S3C). 162

Having established that pmEC activity was modulated in a grid-like fashion for one of the TTCs, we 163 next sought to understand the underlying differences across TTCs in more detail. Previous studies 164 using navigation paradigms (Kunz et al., 2015; Stangl et al., 2018) suggested that these differences 165 in grid-like signals may be due to (i) differences in spatial stability (i.e., grid orientations may differ 166 across voxels and therefore average out), or (ii) differences in temporal stability (i.e., grid orienta-167 tion may change over the course of the experiment). To test whether any one of these factors could 168 explain the pattern of results in our data, we estimated the spatial and temporal stability of the 169 pmEC grid-like signal for each of the 4 TTCs separately. We found that grid orientations clustered 170 across pmEC voxels for all t_{TTC} in a similar way (Fig. S3B; Table S4A), ruling out differences in spatial 171 stability between t_{TTC} (Fig. S3B; MEM, F(3) = 0.36, p = 0.78, $\epsilon^2 = -0.02$, CI : [0, 1]). However, tempo-172 ral stability predicted the amplitude of grid-like signals, which means that the more pmEC voxels 173 showed a stable grid orientation over data partitions, the stronger the resulting cross-validated 174 signal amplitude turned out to be (Fig. 3E; Spearman's rho = 0.57, $p = 5x10^{-4}$). Furthermore, as ex-175 pected based on the results for grid-like signal amplitude, $TTC_{0.86}$ trials showed the highest tempo-176 ral stability among all tested t_{TTC} (Fig. S3A; Table S4B, MEM, F(3) = 3.49, p = 0.018, $\epsilon^2 = 0.05$, CI : [0, 1]). 177

These findings provide evidence that pmEC grid-like signal amplitude differed across tested intervals, and that this difference was largely explained by how stable the putative grid orientation was over time. Importantly, we observed the grid-like signal only for the interval closest to the mean of all intervals. This result dovetails with our prior observation that trial-wise pmEC activity correlated with how strongly behavioral responses were biased towards the mean interval (Fig. 2),

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and it suggests that grid-like signals may also show a relationship to behavioral performance in our task, which was indeed the case. In all of our participants, the grid-like signal amplitude was negatively and linearly related to timing performance, meaning that TTCs with lower RMSE showed the strongest grid signals (Fig. 3F; MEM, $F(3) = 10.1, p = 0.004, \epsilon^2 = 0.28, CI : [0.07, 1]$). Again, this result was not observed for early visual cortex (V1, Fig. S3D; MEM, $F(3) = 2.6, p = 0.125, \epsilon^2 = 0.13, CI : [0.0, 1]$).



Figure 3: A) Visual grid-like analysis method. The hexadirectional signal is cross-validated across data partitions. Putative grid-orientation was estimated using half of the data and then used contrast orientation-aligned vs. to orientation-misaligned gaze movements in the other half (odd vs. even runs). B) Independent regions-of-interest (ROI) analysis for 6-fold symmetry in pmEC. We plot the amplitude of the hexadirectional signal in held-out data expressed as beta estimate. We found reliable cross-validated hexadirectional modulation at the group level only for TTC_{0.86} trials. There were consistent differences in pmEC fMRI activity for aligned vs. misaligned directions across target TTCs. Statistics reflect p<0.05 at FDR-corrected levels (*). C) Voxel-wise analysis results exhibiting activity modulation by gaze movement direction with 60° periodicity aligned to the pmEC grid orientation for TTC_{0.86} trials. We plot thresholded t-test results at 2mm resolution at p < 0.001 uncorrected levels overlaid on a structural template brain. Insert zooming in on EC and MNI coordinates added. D) Control symmetries and regions. Left: 4-fold symmetry in pmEC. Center: 8-fold symmetry in pmEC. Right: 6-fold symmetry in V1. E) TTC $_{0.86}$ pmEC temporal stability predicts corresponding grid-like modulation across participants. Each dot represents a single participant. Regression line (black) and standard error (gray shade). F) Within-subject pmEC grid-like modulation predicts TTC estimation error (RMSE). Stronger pmEC grid-like modulation elicited lower RMSE. Separate regression lines are plotted for each participant. B,D) Depicted are the mean and SEM across participants (black dot and line) overlaid on single participant data (colored dots). Target TTCs are color coded. EF) Participants are color coded.

8

A Bayesian observer model explains performance differences across target TTCs

The behavioral and physiological results presented above suggest that participants' interval esti-190 mates depended on both the current trial and previous trials. This pattern of results is reminiscent 191 of prior work on Bayesian integration (Körding, Wolpert, 2004; Petzschner et al., 2015), which has 192 indeed been argued to underlie contextual effects in interval timing (Jazaveri, Shadlen, 2010; Acerbi 193 et al., 2012; Cicchini et al., 2012). According to the Bayesian framework, the regression-to-the-mean 194 effects we observed at the behavioral level (Fig. 1C) may be a natural consequence of integrating 195 the sensory evidence in each trial (i.e., the inferred probability of a certain TTC being tested) with 196 an expectation informed by the statistical regularities governing prior trials (i.e., the inferred prior 197 distribution of all tested intervals), leading to the characteristic behavioral biases towards the mean 198 of the encoded interval distribution. A Bayesian observer model may therefore provide a parsimo-199 nious computational explanation of both the behavioral regression-to-the-mean effect (Fig 1C) and 200 the observed difference in grid-like activity across TTCs (Fig. 3B). 201

We tested this possibility post-hoc using a Bayesian observer model that was successfully used 202 to model timing behavior in previous work (Jazaveri, Shadlen, 2010, 2015; Remington et al., 2018; 203 Chang, Jazayeri, 2018). Briefly, the model predicts the optimal TTC estimate for each trial by com-204 bining two sources of information: (i) the probability that a specific TTC was tested in a trial given 205 the sensory evidence obtained during the visual tracking phase (Likelihood, Fig. 4A Left), and (ii) a 206 Gaussian prior centered on the mean of the sampled intervals (Prior; Fig. 4A Left). For each par-207 ticipant, a separate model was trained and tested using cross-validation, which built on fitting the 208 model on one half of the data and predicting TTC estimates in the remaining half (see Methods for 209 details). 210

In line with previous work (Jazayeri, Shadlen, 2010, 2015; Remington et al., 2018; Chang, Jazayeri, 211 2018), we found that the model accurately predicted participants' interval estimates across trials 212 (Fig. 4B), with a group average Mean Absolute Error (MAE) of 0.02. The model further recapitulated 213 each participant's task-performance measures (Fig. 1B), with model-derived RMSE values following 214 a quadratic pattern across t_{TTC} similar to the data (Fig. 4C; MEM, F(2) = 74.95, $p = 3.4x10^{-13}$, $\epsilon^2 =$ 215 0.8, CI : [0.7, 1]). Moreover, by relating the model estimates to the neuroimaging results obtained 216 for pmEC, we found that model-derived RMSE values were indeed correlated with the amplitude 217 of grid-like signals observed in all participants (Fig. 4D; MEM, F(1) = 8.15, p = 0.008, $\epsilon^2 = 0.22$, CI: 218 [0.03, 1]). Together, these modeling results suggest that the differences in participants' TTC esti-219 mates and in grid-like signals across TTCs can indeed be well explained using a Bayesian observer 220 model that assumes that predictions informed by temporal regularities in previous trials bias tim-221 ing behavior in the current trial. 222

One notable assumption of the model was that the prior distribution of intervals was centered on 223 the mean of the sampled intervals. To test whether this assumption was valid, we ran the model 224 four times, each time centering the prior close to one of the four tested intervals (Fig. S4). Centering 225 the prior on a value far from the mean interval could in theory result in an even better fitting model 226 and a closer match to our behavioral data. However, we found both the model fit (Fig. S4C, Table 227 S5) and the match between model-derived and real task performance (Fig. S4D) to be optimal 228 when the prior was assumed to be centered on the mean interval. Consequently, only the model 229 with a prior centered on the mean interval recapitulated the quadratic RMSE pattern across t_{TTC} 230 we observed in the data (Fig. S4D). 231



Figure 4: A) Schematic illustration of Bayesian TTC estimation. Left: A Bayesian observer integrates sensory Likelihood and Prior information sources to minimize their mismatch (yellow lines). One source of information comes from a prior expectation (Prior, magenta Gaussian distribution), centered at the mean of the sampled durations (magenta horizontal dashed line). Another source of information comes from noisy sensory inputs (Likelihood, Gaussian distributions of different widths and turquoise tones). On each trial, by combining these two information sources participants can produce a statistically optimal Estimated TTC. Center: An optimal Bayesian estimator that minimizes the mismatch between Prior and Likelihood would be expected to show a Root-Mean-Square-Error (RMSE) pattern across target TTCs best explained by a quadratic function (solid green line). The RMSE increases quadratically as a function of the Target TTC's distance to the mean of the prior (magenta vertical dashed line, 0.82 s). Right: As a result of the integration of these two sources of information, TTC estimates are biased 'away' from the identity line (petrol blue diagonal dashed line) and 'towards' the mean of the sampled TTCs (magenta horizontal dashed line). B) Bayesian observer model performance. We plot the cross-validated model prediction vs. participants' data. The model successfully captures participants' behavior, with a Mean Absolute Error (MAE) of 0.02. Colored lines represent participants' linear model fits across Target TTCs. For most participants the linear model fits are well aligned with the identity line (diagonal grey dashed line). C) Model RMSE per target TTC. Predicted participants' behavior using the crossvalidated Bayesian observer model. RMSE differences across TTCs show a quadratic pattern, replicating real participants' behavior (Fig. 1B). D) As expected, participants' pmEC grid-like signals predict model's RMSE, replicating our experimental observations (Fig. 3F). BC) Target TTCs are color coded. Single-participant data plotted as dots. BD) Separate regression lines are plotted for each participant.

Discussion 232

Prior

Predicted RMSE

The present study examined whether and how the human pmEC contributes to the encoding of 233 task regularities that guide timing behavior. We used fMRI to record human brain activity while par-234 ticipants performed a rapid time-to-contact estimation task, which allowed us to analyze trial-wise 235 pmEC activity as a function of time-estimation performance across a range of sampled intervals. 236 Moreover, the task included periods in which participants followed a moving fixation target with 237 their eyes, allowing us to estimate visual grid-like signals in the pmEC: the amplitude and stabil-238

(Beta estimate)

ity of a six-fold rotationally symmetric MRI-signal modulation as a function of gaze direction. We 239 found that pmEC activity closely tracked task performance across trials as well as behavioral re-240 sponse biases towards the mean interval, and we observed a bias towards that mean interval on 241 the level of grid-like signals. Traditionally, such regression-to-the-mean biases have been taken as 242 evidence for Bayesian integration in the brain (Petzschner et al., 2015), since they are well explained 243 by models that assume the integration between sensory evidence obtained in a trial with a prior 244 expectation derived from previous trials. Indeed, a Bayesian observer model previously used to 245 model timing behavior in humans and macaques (Jazayeri, Shadlen, 2010, 2015; Remington et al., 246 2018; Chang, Jazayeri, 2018; De Kock et al., 2021) provided a parsimonious account for both our 247 behavioral and pmEC results. In the following, we will discuss these results in light of previous liter-248 ature on timing behavior and relate them to prior work on spatial and non-spatial coding principles 249 in the hippocampal formation. 250

251 Task regularities bias TTC estimation in future trials

While participants were not explicitly told about the true range of intervals that were tested, their 252 estimates were nevertheless biased towards the mean interval (Fig. 1B, C). This regression-to-the-253 mean effect in time estimation is well documented and has been proposed to reflect participants' 254 reliance on the temporal regularities learned from previous trials (Miyazaki et al., 2005: Jazaveri, 255 Shadlen, 2010; Acerbi et al., 2012; Cicchini et al., 2012; Petzschner et al., 2015; Chang, Jazayeri, 256 2018; Polti et al., 2022). Specifically, integrating a prior expectation with the sensory evidence ob-257 tained in the current trial may help participants to anticipate the trajectory of moving objects during 258 occlusion (Fig. 4A). This prior expectation likely takes the form of a Gaussian distribution of time 259 intervals centered on the mean (Fig. S4D). Relying on this prior to make an interval judgement 260 will therefore most often be biased towards that mean, depending on how strong the sensory evi-261 dence is in a given trial about the true TTC that was tested. With increased mismatch between prior 262 expectation and sensory evidence, participants' estimates may be biased more towards the mean 263 (Fig. 1B, C), potentially reflecting an increase in uncertainty about their estimate (Jazayeri, Shadlen, 264 2010; Petzschner et al., 2015). Overall, this strategy may lead to large errors in some trials, but it 265 may nevertheless be adaptive, since the mean is still a "good guess" for the large majority of trials. 266 This is especially true for those trials in which the evidence derived from the senses is sparse or 267 noisy. Interestingly, we tested only four intervals drawn from a uniform distribution, meaning that 268 the mean interval does not actually account for a large proportion of the trials. However, previous 269 work has shown that interval estimates tend to be encoded using a Gaussian distribution even 270 when the intervals were sampled uniformly (Acerbi et al., 2012), in line with our observation that 271 our data was well described using a Bayesian model that assumed a Gaussian distribution centered 272 on the mean interval (Fig. 4). 273

274 Entorhinal cortex encodes task regularities that afford time estimation

Entorhinal activity reflected the behavioral response biases towards the mean across trials, as well 275 as overall task performance. In our view, this result is striking in our view as the functions of the 276 entorhinal cortex are mostly studied in the context of spatial navigation (Epstein et al., 2017; Kunz 277 et al., 2019) and long-term memory formation (Fernández et al., 1999; Hargreaves et al., 2012; 278 Staresina et al., 2013; Schiller et al., 2015), not in the context of rapid timing tasks. Notably, similar 279 tasks have been used successfully for studying predictive processes in rodents (Henke et al., 2021), 280 humans (Jazaveri, Shadlen, 2010; Acerbi et al., 2012; Cicchini et al., 2012; Chang, Jazaveri, 2018; 281 Polti et al., 2022), and non-human primates (Jazayeri, Shadlen, 2015; Wang et al., 2018; Sohn et al., 282

2019), but not in the context of entorhinal functions. Our results suggest that computations in the 283 human entorhinal cortex contribute to timing task performance in real time as the task is being 284 performed. In fact, we found trial-wise pmEC activity to follow a pattern that closely resembled the 285 one previously reported for the adjacent hippocampus, as well as for other regions prominently 286 including the striatum (Polti et al., 2022; Rolando et al., 2024). The contributions of the entorhinal 287 cortex to task performance may therefore closely depend on other regions shown to preferentially 288 encode distinct behaviorally relevant factors, such as task details versus task structure (Doeller 289 et al., 2008; Geerts et al., 2020). Specifically, the hippocampal-entorhinal region has been shown 290 to signal the encoding of task structure (e.g., graphs reflecting transition probabilities between se-291 quentially presented stimuli (Garvert et al., 2017)) and abstract task spaces (Constantinescu et al., 292 2016; Theves et al., 2019, 2020; Viganò, Piazza, 2020; Park et al., 2021), which may reflect the learn-293 ing of generalizable principles that guide behavior across tasks. Our results are in line with these 294 ideas and support recently proposed computational accounts of entorhinal function that center 295 on structured, factorized representations allowing inference and generalisation (Whittington et al., 296 2020). 297

Our results further dovetail with work on temporal-context encoding (Schapiro et al., 2012; Hsieh 298 et al., 2014) and sequence memory (Fortin et al., 2002; Montchal et al., 2019; Bellmund et al., 2020a, 299 2022) in the hippocampal-entorhinal region. For example, animal studies have shown that damage 300 to the entorhinal cortex impairs memory for relations (Buckmaster et al., 2004), and inactivation 301 of the rodent MEC prevents context-dependent learning of intervals (Bigus et al., 2023). In general, 302 recent years have seen a growth in the literature on the links between MEC and timing behavior in 303 rodents (Heys, Dombeck, 2018; Heys et al., 2020; Vo et al., 2021; Dias et al., 2021; Bigus et al., 2023). 304 Our results extend these reports to the human pmEC, revealing a direct relationship between pmEC 305 activity and behavioral performance in a timing task. Specifically, its activity mirrored a behavioral 306 bias towards the mean of the tested intervals; a phenomenon that occurs for any type of magni-307 tude estimation (Petzschner et al., 2015; Petzschner, Glasauer, 2011). Investigating the relationship 308 between these mean biases and neural activity across a range of tasks therefore provides fertile 309 ground for investigations of the domain-general functions of the hippocampal-entorhinal region 310 (Behrens et al., 2018; Bellmund et al., 2018; Stachenfeld et al., 2017). In line with this idea, hip-311 pocampal activity has been linked to regression-to-the-mean biases during the estimation of spa-312 tial distances (Wiener et al., 2016) and intervals (Polti et al., 2022), and entorhinal activity has been 313 shown to be modulated by spatial context during virtual navigation (Julian, Doeller, 2021). 314

315 Non-spatial task factors shape entorhinal grid-like signals

In addition to co-variations in pmEC activity and task-performance measures across trials, we found 316 that pmEC grid-like signals in particular reflected participants' performance across intervals (Fig. 3F). 317 Like behavioral performance, also the grid-like signal seemed to be biased towards the mean inter-318 val, with cross-validation revealing a robust grid-like modulation exclusively for the interval closest 319 to the mean. This effect was driven by the temporal stability, not spatial stability, of the grid-like 320 signal (Figs. S3A, B), in line with previous reports (Kunz et al., 2015; Stangl et al., 2018). Furthermore, 321 our results also replicate previous work on visual grid-like signals in humans (Nau et al., 2018b; lu-322 lian et al., 2018; Staudigl et al., 2018), while going beyond these studies by reporting a relationship 323 between these signals and behavioral performance. Eve movements may engage similar processes 324 in the entorhinal cortex as navigation (Nau et al., 2018a), while offering higher experimental con-325 trol and study-design efficiency. Since behavioral performance was well explained by a Bayesian 326 observer model, we tested whether this model also predicted the grid-like signal differences across 327

intervals, which was indeed the case (Fig. 4C). Cross-validated model predictions recapitulated the
 link between task performance and the grid-like signal differences across intervals, potentially of
 fering a parsimonious and normative explanation of our findings. The entorhinal cortex, and grid
 cells in particular, may support the encoding of task regularities that ultimately manifest in both
 neural activity and behavior as regression-to-the-mean effects.

Why would a spatial grid-like signal be modulated by the range of intervals that was tested? Previ-333 ous work has shown that task-dependent factors such as environmental features, goal locations, 334 and rewards can distort the grid-cell pattern in rodents (Barry et al., 2007, 2012; Krupic et al., 2015; 335 Hardcastle et al., 2015; Keinath et al., 2018; Boccara et al., 2019; Butler et al., 2019), as well as 336 grid-like (Viganò et al., 2023) and behavioral (Bellmund et al., 2020b; Chen et al., 2015) response 337 patterns in humans. It therefore seems plausible that also human grid-like signals can be modu-338 lated by behaviorally-relevant features of the task. In our task, the most relevant feature was the 339 intervals that were tested, possibly explaining why the timing error was lowest (Fig. 1C) and the 340 temporal stability of the grid-like signal was highest (Fig. S3A) for the interval closest to the mean. 341 Previous work suggested that the distortions of the grid-cell patterns could be the consequence 342 of conflicting sources of information (Barry et al., 2007; Hardcastle et al., 2015; Krupic et al., 2015), 343 which would be broadly in line with our Bayesian modeling results (Fig. 4). In our case, the con-344 flicting sources of information may be the prior expectation derived from previous trials, which 345 is inherently biased towards the mean interval, and the current-trial evidence derived from the 346 senses. Therefore, we speculate that the amplitude and orientation of grid-like signals depend 347 on the agreement between these two sources of information. This alignment becomes higher the 348 closer the tested interval is to the mean interval. 349

³⁵⁰ Predictive processing as a domain-general principle of entorhinal function?

While many of the above considerations remain to be tested, it has long been recognized that learn-351 ing task regularities ultimately requires the encoding of relationships between stimuli, actions, and 352 events (Körding, Wolpert, 2004; Petzschner, Glasauer, 2011; Petzschner et al., 2015), which has 353 been proposed to build on a relational coding scheme that has often been discussed for the me-354 dial temporal lobe (Manns, Eichenbaum, 2006; Whittington et al., 2020) even for non-spatial feature 355 spaces (e.g., Aronov et al. (2017); Bellmund et al. (2018); Behrens et al. (2018); Constantinescu et al. 356 (2016); Bao et al. (2019); Theves et al. (2019, 2020); Viganò et al. (2021); Park et al. (2021); Wagner 357 et al. (2023); Nitsch et al. (2023)). Our task explicitly required participants to make temporal pre-358 dictions, but the central ideas and observations presented in this article may therefore very well 359 translate also to other tasks and less explicit situations. Taking a predictive-processing perspective 360 may more generally help to understand the functional contributions of the entorhinal cortex across 361 behavioral domains and species (e.g., Cothi de et al. (2022)), which includes, but is not limited to, the 362 Bayesian framework. In fact, a growing number of studies support the idea that Bayesian models 363 can provide a normative explanation for a range of observations in the spatial navigation literature, 364 including the integration of visual cues and landmarks during path integration (Cheng et al., 2007; 365 Petzschner, Glasauer, 2011; Kessler et al., 2022). Intriguingly, the same Bayesian model (Kang et al., 366 2023) that explains distortions in spatial memory in humans (Hartley et al., 2004; Chen et al., 2015; 367 Bellmund et al., 2020b; Keinath et al., 2021) can also explain distortions of the grid-cell pattern in 368 rodents (e.g., Krupic et al. (2015)), pointing towards a unified computational account for behavior 369 and entorhinal activity across species. 370

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371 Predictions for future work

Our results suggest that non-spatial task factors shape human entorhinal activity in real time as the 372 task is performed, a phenomenon that future studies should investigate across a range of tasks. 373 It is important to note that we modeled our results using a Bayesian observer model due to its 374 successful application to similar problems in the past (Jazayeri, Shadlen, 2010; Acerbi et al., 2012; 375 Remington et al., 2018; Chang, Jazayeri, 2018), but other computational frameworks may be able 376 to explain our data equally well. Approaches that model the mismatch between prior expectations 377 and sensory evidence more explicitly seem especially promising in this context (e.g., prediction er-378 rors in reinforcement learning (Momenneiad et al., 2017; Niv, 2009; Dayan, Daw, 2008; Stachenfeld 379 et al., 2017)). Second, our results are in line with the idea that participants encoded the tested 380 intervals using a Gaussian distribution centered on the mean intervals, but a more fine-grained 381 sampling of intervals would be necessary to convincingly show that this was actually the case. The 382 ideal scenario would be to sample many more time intervals, not from one but from multiple distri-383 butions, each centered on a different value. In this case, one would predict that the grid-like signal 384 is stable at the center of each of these distributions. Finally, to investigate the range of non-spatial 385 factors that affect grid-like signals, and therefore to understand potential domain-general contri-386 butions of the pmEC to human cognition, future work should test a range of tasks beyond time 387 estimation and spatial navigation. 388

389 Conclusion

Taken together, using fMRI and a rapid time-to-contact estimation paradigm in humans, we showed 390 that time estimates are biased towards the mean of the tested intervals, and that this mean bias is 391 reflected in entorhinal activity across trials (similar to the hippocampus, Polti et al. (2022)). More-392 over, we report a novel relationship between grid-like signals and behavioral performance, as the 393 amplitude of the grid-like signal correlated with participants' time-estimation error, and the puta-394 tive grid orientation was stable exclusively for the interval closest to the mean. Finally, both our 395 behavioral and neuroimaging results were well explained (post hoc) by a Bayesian observer model 396 that assumes the integration of prior expectations and sensory evidence in each trial. These re-397 sults point to an involvement of the pmEC in interval timing, likely building on the encoding of task 398 regularities that afford predictive inference in the service of goal-directed behavior. 399

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410 Author Contributions

411 MN, IP and CFD conceived the study. IP and MN designed the experimental paradigm, visualized 412 the results, and embedded them in the literature with help from RK, VW and CFD. IP implemented

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- the experimental code and acquired and analyzed the data with close supervision from MN. MN
- and IP wrote the manuscript with critical feedback from RK, VW, and CFD. CFD provided general
- ⁴¹⁵ supervision of the project and secured funding. IP and MN are shared-first authors.

416 **Declaration of interest**

⁴¹⁷ The authors declare no conflicts of interest.

418 Data and code availability

Source data and analysis code are available on Open Science Framework (https://osf.io/cs8d6/), along side pre-processed eye-tracker data (https://osf.io/mrhk9/. Raw fMRI data is available at the following
 G-Node Infrastructure repository: https://gin.g-node.org/ipolti/TTC_HPCF.git. The original Bayesian Ob server Model has been shared by Jazayeri and colleagues and can be found here: https://jazlab.org/
 resources/.

424 Methods

425 Participants

The data used in this study were used in a previous report (Polti et al., 2022). Thirty-nine participants 426 (16 women, 23 men, 19-35 years old) were recruited for this study. Five participants were excluded: 427 one was excluded because the eye-tracker calibration failed, one was excluded because they did 428 not follow the task instructions, and three participants were excluded because of technical issues 429 during scanning. A total of 34 participants entered the analysis. The study was approved by the 430 regional committee for medical and health research ethics (project number 2017/969) in Norway 431 and participants gave their written consent prior to scanning in accordance with the Declaration of 432 Helsinki (World Medical Association, 2013). 433

434 Task

We used a Time-to-contact (TTC) task that required participants to estimate the time when a moving 435 dot reached a target location. Each trial began with the smooth visual pursuit of a dot moving in 436 1 of 24 predefined linear trajectories with 1 of 4 possible speeds: 4.2°/s, 5.8°/s, 7.5°/s and 9.1°/s. 437 All trajectories had two segments, one where the dot was visible ("Gaze trajectory", Fig. 1A; length 438 of 10 dva) and one where the dot was occluded (length of 5 dva). Because all trajectories had the 439 same length, the 4 speeds led to 4 target TTC (t_{TTC}) durations: 1.2s, 0.88s, 0.67s, and 0.55s. A t_{TTC} 440 duration was defined as the time it takes the dot to traverse the occluded segment. When the 441 dot reached the end of the visible segment, a fixation cross remained in place until participants 442 had performed a TTC estimation judgment. The latter consisted of clicking a button when the 443 dot presumably reached the end of the occluded segment ("Boundary", Fig. 1A). After giving a 444 response, participants received visual feedback for 1 s reflecting their TTC estimation accuracy. 445 When accuracy was not high, a visual cue signaled the TTC error direction: either responding too 446 early (underestimation) or too late (overestimation). At the feedback offset, the dot became visible 447 again and remained static for a variable ITI sampled randomly from a uniform distribution ranging 448 from 0.5 to 1.5 s. Then, a new trial began with the dot moving in a different direction. Over the 449 course of 768 trials, we sampled eye movement directions with 15° resolution. Participants were 450 never explicitly informed about the full visual trajectory or the range of t_{TTC} . See previous work of 451 Polti et al. (2022) for more details on the task. 452

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453 Behavioral analysis

Participants indicated the estimated TTC in each trial via button press. To test if participants discriminated the four target TTCs we used a linear mixed-effect model with estimated TTC (e_{TTC}) as the dependent variable, target TTC (t_{TTC}) as the predictor and separate intercepts and slopes per participant.

We partitioned participants' behavioral performance for each t_{TTC} i using two parameters, Accuracy and Precision (Fig. S1A). Accuracy_i was computed as the absolute difference between the average e_{TTC} (e_{TTC}) and t_{TTC} , divided by e_{TTC} :

$$Accuracy_{i} = \frac{|e_{\overline{TTC}} - t_{TTC}|}{e_{\overline{TTC}}}$$
(1)

Precision_i was computed as the coefficient of variation (CV), given by the standard deviation of the e_{TTC} divided by the e_{TTC} :

$$Precision_{i} = CV_{i} = \frac{\sqrt{\sum (e_{TTC} - e_{TTC})^{2}/N}}{e_{TTC}}$$
(2)

where *N* is the number of e_{TTC} values. These two parameters can be integrated in a single measurement that reflects the precision-accuracy trade-off when written as a Pythagorean sum, i.e. the root-mean-square error (RMSE):

$$RMSE_{i} = \sqrt{Precision_{i}^{2} + Accuracy_{i}^{2}}$$
(3)

In order to analyze the pattern of RMSE across target TTCs, we used a mixed-effect model with RMSE as the dependent variable, target TTC as a quadratic predictor, and separate slopes and intercepts per participant (Fig. 1B). We also ran the same model again while using a linear instead of quadratic trend. We then tested which model best explained the changes in RMSE across target TTCs using a chi-square test.

471 Imaging data acquisition & preprocessing

Imaging data were acquired on a Siemens 3T MAGNETOM Skyra located at the St. Olavs Hospi-472 tal in Trondheim, Norway. A T1-weighted structural scan was acquired with 1mm isotropic voxel 473 size. Following EPI-parameters were used: voxel size=2mm isotropic, TR=1020ms, TE=34.6ms, flip 474 angle=55°, multiband factor=6. Participants performed a total of four scanning runs of 16-18 min-475 utes each including a short break in the middle of each run. Functional images were corrected for 476 head motion and co-registered to each individual's structural scan using SPM12 (www.fil.ion.ucl.ac. 477 uk/spm/). We used the FSL topup function to correct field distortions based on one image acquired 478 with inverted phase-encoding direction (https://fsl.fmrib.ox.ac.uk/fsl/fslwiki/topup). Functional images 479 were then spatially normalized to the Montreal Neurological Institute (MNI) brain template and 480 smoothed with a Gaussian kernel with full-width-at-half-maximum of 4 mm for regions-of-interest 481 analysis or with 8 mm for whole-brain analysis. Time series were high-pass filtered with a 128 s 482 cut-off period. 483

484 Region-of-interest (ROI) definition and analysis

Rodent studies have consistently reported grid cells in the medial entorhinal cortex (Heys et al.,
2014; Moser et al., 2014), which likely corresponds to the posteromedial portion of the entorhinal
cortex in humans (Maass et al., 2015; Navarro Schröder et al., 2015). We therefore used a (bilateral) posteromedial entorhinal cortex (pmEC) mask from Navarro Schröder et al. (2015) in all our

fMRI analyses (Fig. S2). As a control region, we chose the early visual cortex (V1) for which, to our 489 knowledge, no hexadirectional signals have been reported. V1 masks were generated for each indi-490 vidual participant based on the automatic parcellation derived from FreeSurfer's structural recon-491 struction (https://surfer.nmr.mgh.harvard.edu/). A pre-supplementary motor area (preSMA) mask was 492 obtained from the JuBrain SPM anatomy toolbox (https://www.fz-juelich.de/inm/inm-1/EN/Forschung/ 493 docs/SPMAnatomyToolbox/SPMAnatomyToolbox node.html) in order to post-hoc confirm whether activ-494 ity observed in voxel-wise analyses corresponded to preSMA. All masks were spatially normalized 495 to the MNI brain template space and re-sliced to the functional imaging resolution using SPM12. 496

All ROI analyses described in the following were conducted using the following procedure. We extracted beta weights estimated for the respective regressors of interest for all voxels within a region in both hemispheres, averaged them across voxels within that region and performed a onesample Wilcoxon test on group level against zero as implemented in the software R (https://www. R-project.org).

502 EC activity as a function of accuracy and as a function of the regression effect

To examine the relationship between behavioral biases and brain activity, we used mass-univariate 503 general linear models (GLM) to model the trial-wise activity of the pmEC voxels as a function of 504 accuracy (i.e. the absolute difference between estimated and target TTC in each trial) and as a 505 function of the regression effect (i.e. the absolute difference between the estimated TTC and the 506 mean of the tested intervals, which was 0.82 s) in the TTC task. To avoid effects of potential co-507 linearity between these regressors, we estimated model weights using two independent GLMs, 508 which modeled the time course of each trial with either one of the two regressors. The GLMs 509 also included one regressor modeling ITIs, one for button presses and one for periods of rest, 510 which were all convolved with the canonical hemodynamic response function in SPM12. In addition, 511 the models included the six realignment parameters obtained during pre-processing as well as a 512 constant term modeling the mean of the time series. After fitting each model, we used the weights 513 estimated for the two regressors to perform ROI analyses for the EC using a two-tailed one-sample 514 Wilcoxon test (Fig. 2). 515

516 Hexadirectional analysis of visual grid-like representations

Prior work showed that the MRI signal in the human entorhinal cortex exhibits a six-fold rotationally symmetric modulation as a function of gaze direction, which is thought to reflect grid-cell population activity (Nau et al., 2018b; Julian et al., 2018). Here, we tested whether such grid-like signals were also detectable in our data. The analysis builds on cross-validation to first estimate the putative grid orientation relative to the screen, and then testing in held-out data whether gaze directions aligned to this putative grid orientation are associated with stronger MRI signals than directions misaligned to it.

To estimate the putative grid orientation, we first modeled the activity in each voxel in half of the 524 data using a GLM (odd vs even runs) that incorporated two main regressors of interest. These 525 regressors modeled the sine and cosine of the movement direction of the fixation target θ with a 526 periodicity of 60° , i.e. $\sin(6\theta)$ and $\cos(6\theta)$. For each trial, θ included the tracking and TTC estimation 527 task phases. In addition, nuisance regressors modeled ITIs, the feedback phase, button presses 528 and periods of rest, and two parametric regressors modeled the effect of feedback on the activity 529 during the feedback phase and another one modeling the effect of TTC bias on button presses. 530 All regressors were convolved with the HRF. Because different fixation-target speeds also led to 531

different TTCs and thus trial durations, and because the amplitude of the HRF-convolved signal
 scaled with duration, we re-scaled the resulting main regressors to obtain a balanced regressor
 amplitude for all speeds. Moreover, we added the realignment parameters and a constant term to
 the model. Each speed level was modeled separately. Weights for all regressors were estimated
 using SPM12.

We used the beta weights corresponding to the sine ($\beta \sin$) and cosine ($\beta \cos$) of each target TTC level 537 to estimate the putative grid orientation relative to the screen ϕ for voxels within the entorhinal 538 cortex using the following formula: $\phi = \arctan(\beta \sin /\beta \cos)$. The estimated orientations were then cir-539 cularly averaged across voxels and across runs within each data partition, resulting in a single grid 540 orientation for each target TTC and data partition. We then used the estimated grid orientation 541 in a second GLM to estimate the amplitude of the grid-like signal in its independent data counter-542 part (Fig. 3A). To do so, we again modeled nuisance variance as described before, this time adding 543 one main regressor per target TTC modeling the cosine of each fixation-target movement direc-544 tion modulo 60° aligned to the previously estimated mean orientation using the following formula: 545 $\cos(6(\theta - \phi))$. Again, all regressors but the realignment parameters and the constant term were con-546 volved with the HRF, and all main regressors were rescaled to match their amplitude across TTCs. 547 We then again estimated weights for all regressors using SPM12 and averaged them across the 548 pmEC ROI. For each target TTC, we tested its corresponding estimated weight against zero using a 549 one-tailed one-sample Wilcoxon test on group-level (Fig. 3B; Table S1). In order to test for a main 550 effect of target TTC on grid-like signal amplitude, we ran a mixed-effects model with target TTC 551 as predictor, the estimated weights as the dependent variable and participants as the error term. 552 We used two-tailed paired Wilcoxon signed-rank tests to compare differences in grid-like signal 553 between target TTCs (Table S1). 554

To test whether the grid-like signal found in the pmEC exhibited a specifically 6-fold symmetric periodicity (60°), not other periodicities, we repeated the cross-validation analysis for 4-fold (i.e. 90°, Fig. 3D, Left) and 8-fold (i.e. 45°, Fig. 3D, Center) symmetries. In addition, we tested for 6-fold symmetry in a control region (early visual cortex (V1), Fig. 3D, Right).

In order to exhibit a reliable grid-like signal, a voxel's grid orientation should remain stable over time (i.e. temporal stability). For each participant and target TTC, we therefore computed the percentage of pmEC voxels that maintained an orientation difference of less than 15° between training and test data partitions. We then tested if differences in temporal stability could explain individual differences in the pmEC grid-like modulation for TTC_{0.86} using a Spearman's rank-order correlation (Fig. 3E).

Finally, to asses the behavioral relevance of the pmEC grid-like signal, we ran a linear mixed effect model with the RMSE for all four target TTCs as the dependent variable and their corresponding pmEC grid-like signal as a predictor. The model included separate intercepts and slopes for each participant. We expected that a stronger visual grid-like modulation would predict lower RMSE values (i.e. better performance in the TTC task, Fig. 3F).

570 Additional analyses of pmEC visual grid-like signals

Besides having high temporal stability, a robust grid-like signal is also expected to have high spatial stability (i.e., pmEC voxel-wise grid orientations should cluster in order to provide a robust mean grid orientation). We used Rayleigh's *z*-value as a measure of spatial stability: higher *z* values correspond to higher voxel-wise grid orientation clustering. For each participant and target TTC (t_{TTC}), we computed Rayleigh's test for non-uniformity of circular data on the pmEC voxel-wise grid orientations from the training data partition. We tested the Rayleigh's *z*-values separately for each t_{TTC} against zero using one-tailed one-sample Wilcoxon tests on group-level. In order to compare spatial stability between t_{TTC} , we again used a mixed-effects model incorporating spatial stability as a dependent variable, t_{TTC} as a predictor, and participants as the error term (Fig. S3B). We further used a Spearman's rank-order correlation to test if individual differences in spatial stability could predict the cross-validated grid-like signal amplitude estimated for TTC_{0.86} (Fig. S3C; Spearman's rho = 0.49, p = 0.003).

Differences in grid-like signal amplitude across t_{TTC} could also be explained by differences in temporal stability. We thus used a mixed-effects model with temporal stability as a dependent variable, t_{TTC} as a predictor and participants as the error term to test this possibility (Fig. S3A). This analysis was conducted both for the pmEC and the early visual cortex control (Fig. S3D).

587 Bayesian observer model

The Bayesian observer model, developed originally by Jazayeri and colleagues (Jazayeri, Shadlen,
 2010), has been successfully adapted and applied several times to model timing behavior (Jazayeri,
 Shadlen, 2015; Remington et al., 2018; Chang, Jazayeri, 2018; De Kock et al., 2021). We adapted the
 original code provided at https://jazlab.org/resources/ in order to model participants' behavior in our
 TTC estimation task.

The Bayesian observer model is composed of three stages. In the first stage, the observer makes a noisy duration measurement (m_{VI}) of the visual tracking period (VT, Fig. 1A), from the movement onset of the fixation disc until it becomes occluded. Measurement noise was modeled as a zeromean scalar Gaussian likelihood distribution (Gibbon, 1977). Specifically, the standard deviation of measurement noise was scaled as a function of the VT duration with constant of proportionality w_m (Weber fraction for VT duration measurement).

⁵⁹⁹ During the second stage, the Bayesian observer integrates two sources of information to minimize ⁶⁰⁰ their mismatch: the likelihood of m_{vt} and a prediction based on prior knowledge (a statistical rep-⁶⁰¹ resentation of the temporal context, i.e., the distribution of sampled VT durations). The prior was ⁶⁰² modeled as a Gaussian distribution centered on the mean of the range of sampled VT durations, ⁶⁰³ with standard deviation equal to the standard deviation of the sampled VT durations. As a result of ⁶⁰⁴ Bayesian integration, the posterior distribution is then computed over the sampled VT durations.

In the third stage, the Bayesian observer selects the optimal VT duration estimate corresponding 605 to the mean of the posterior based on a guadratic loss function. This value is further scaled by 0.5, 606 a gain factor (Gf) equal to the ratio of distances between the occluded and visible segments of the 607 path (5 / 10 dva; Fig. 1A). The scaled value corresponds to the optimal estimated TTC (e_{TTC}), which 608 is then augmented by TTC production noise to account for motor variability. TTC production noise 609 was modeled as a scalar Gaussian distribution centered at the mean of the posterior. The standard 610 deviation of TTC production noise scaled as a function of e_{TTC} with constant of proportionality w_p 611 (Weber fraction for TTC production). To account for idiosyncratic response biases (e.g. consistently 612 producing TTC responses earlier or later than expected), the model includes an offset parameter 613 o_b (Remington et al., 2018). 614

In order to fit the Bayesian observer model to behavioral data, we used a cross-validation procedure. We first split the data of each participant into two independent halves (384 trials each) with an equal number of trials for each t_{TTC} . The trials were randomly sampled without replacement throughout the experimental session. The model parameters (w_m , w_p , o_b) were fitted separately on each data partition by maximizing the log-likelihood of participants' responses given the sampled

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TTCs using the *fminsearch* MATLAB function. Parameter searches were repeated 10 times using different initialization values, showing a high degree of consistency between iterations and data partitions (Fig. S4A). Given such consistency, we used as final model parameters the median value across all the fits to the data (Fig. S4B). These model parameters were further used for model simulations, where we generated the same number of trials and t_{TTC} combinations as participants had in each data partition.

To evaluate the model predictions we computed the RMSE of each *t*_{TTC} as a summary statistic and 626 calculated the Mean Absolute Error (MAE) between the observed and predicted RMSE (Fig. 4B). 627 To corroborate that the model replicated participants' quadratic RMSE pattern across t_{TTC} , we ran 628 a mixed-effects model with the model RMSE as a dependent variable, target TTC as a quadratic 629 predictor and separate intercepts and slopes per participant. Furthermore, to test the model suit-630 ability to replicate the observed relationship between pmEC grid-like signals and RMSE, we used 631 a mixed-effects model with model RMSE as the dependent variable, participants' pmEC grid-like 632 signal as a predictor, and separate intercepts and slopes per participant. 633

In our model, we assumed a prior Gaussian distribution centered on the mean of the sampled 634 VT durations (1.64 s), and Gaussian likelihood functions centered on the respective value of each 635 sampled VT duration (1.1 s, 1.34 s, 1.72 s and 2.4 s; Fig. 4A, Left). With these settings, the smallest 636 mismatch between prior and likelihood occurs for the sampled VT duration 1.72 s (Fig. 4A, Left, 637 yellow lines). Consequently, the lowest RMSE values are observed at the corresponding t_{TTC} of 638 this VT duration (0.86 s; Fig. 4A, Center). We explored alternative configurations by running model 639 simulations where the prior was centered close to a different VT duration each time. The different 640 prior values were obtained by multiplying each VT duration with the ratio between the average 641 VT duration 1.64 s and the VT duration 1.72 s. When scaled by Gf the prior means correspond to 642 0.52 s, 0.64 s, 0.82 s and 1.14 s, and the sampled VT durations correspond to the t_{TTC} 0.55 s, 0.67 643 s, 0.86 s and 1.2 s (Fig. S4D). For each participant and model, we simulated the same number of 644 trials as were originally tested and calculated the MAE between the observed and simulated RMSE 645 (Fig. S4C). To examine potential differences in MAE across model simulations with different prior 646 means, we ran a mixed-effects model with MAE as the dependent variable, prior mean value as 647 predictor and participants as the error term. We found a main effect of prior mean on the MAE (; 648 $F(3) = 44.23, p = 2.2x10^{-16}, \epsilon^2 = 0.55, CI : [0.44, 1]$). We expected the model with the prior centered on 649 the mean interval to show the lowest MAE. To test this prediction, we conducted one-tailed paired 650 Wilcoxon signed-rank tests between each model pair (Table S5). 651

652 Eye tracking

We used an MR-compatible infrared eye tracker with long-range optics (Eyelink 1000) to monitor 653 gaze position at a rate of 500 hz during the experiment. After blink removal, the eve tracking data 654 was linearly detrended, median centered, downsampled to the screen refresh rate of 120 hz and 655 smoothed with a running-average kernel of 100 ms. Fixation error was computed separately for 656 each participant and trial as the euclidean distance between the fixation target and the measured 657 gaze position. In order to test for systematic imbalances or biases in viewing behavior, we used sep-658 arate mixed-effects models (one per t_{TTC}) with fixation error as the dependent variable, direction 659 as predictor, and participants as the error term. There were no significant differences in fixation 660 error across directions (Fig. S1B; Table S6). 661

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⁸⁹² Supplementary Material



Figure S1: A) Precision (Coefficient of Variation) as a function of Accuracy (the absolute difference between average estimated TTC and target TTC, normalized by the average estimated TTC) separately for each target TTC. The distance from the origin is the TTC performance, i.e. the Root-Mean-Square-Error (RMSE). Solid grey quarter circle lines represent the locus of the mean RMSE for each target TTC. The same RMSE can arise from low accuracy and high precision, and from high accuracy and low precision. Mean RMSE for each target TTC represented by a colored dot with a black edge. Target TTCs are color coded. B) Euclidean distance between fixation target and gaze (fixation error). There were no significant differences in fixation error across all 24 gaze directions. Fixation quality does not affect the gaze-dependent hexadirectional modulation in EC presented in this study. Each dot per direction represents a single participant. Target TTCs are color coded. Group-level mean (black line) and SEM (gray shade). Fixation error displayed as degree of visual angle (radial axis). AB) Single-participant data plotted as dots.

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Figure S2: Regions of interest (ROIs). Red) posterior medial entorhinal cortex (pmEC) ROI representing the human homolog of rodent medial entorhinal cortex. This mask was obtained from Navarro Schröder et al. (2015). Green) pre-supplementary motor area (preSMA) obtained from the JuBrain SPM anatomy toolbox. Blue) early visual cortex (V1) anatomically defined for each participant using FreeSurfer's cortical parcellation. ROIs superimposed onto a 2mm resolution skull-stripped structural template brain. MNI coordinates added.

Figure S3: Additional pmEC grid-like signal analyses. A) Significant differences in temporal stability of grid-like signals in pmEC voxels across target TTCs. $TTC_{0.86}$ showed the highest percent of pmEC temporally stable voxels. B) No significant difference in spatial stability of grid-like signals in pmEC voxels across target TTCs. C) $TTC_{0.86}$ pmEC spatial stability predicts corresponding grid-like signal across participants. Each dot represents a single participant. Regression line (black) and standard error (gray shade). D) Within-participant V1 grid-like signal does not predict TTC estimation RMSE. Separate regression lines are plotted for each participant. AB) Depicted are the mean and SEM across participants (black dot and line) overlaid on single participant data (colored dots). Target TTCs are color coded. CD) Participants are color-coded.

Figure S4: Bayesian observer model parameters and simulations. A) Model parameter differences across data partitions. Distribution of differences in parameter values across data partitions centered around zero across participants. The magnitude of these differences were relatively small. Model parameters: w_m (measurement noise parameter, green), w_p (TTC production noise, blue), o_b (response bias, red). B) Median parameter value across data partitions for each participant. Parameter values showed differences across participants. Parameters were color coded. C) Mean Absolute Error (MAE) across Bayesian model simulations with different prior mean values. Group average model performance was best for simulations that set the prior mean value at 0.82 s, the average across sampled durations. Model simulations with different prior mean values were color coded. Statistics reflect p<0.05 at FDR-corrected levels (*). D) Bayesian model performance using different prior mean values. For each model simulation, we show the simulated vs observed RMSE for each participant (Left) and the pattern of RMSE across t_{TTC} . The model simulation with a prior mean value equal to the average across sampled durations (0.82 s) displayed a quadratic RMSE pattern across t_{TTC} that most resembled participants' observed behavior. Separate regression lines are plotted for each participant. t_{TTC} are color coded. Grey diagonal dashed line represents the identity line. Better model performance can also be observed as greater overlap between the individual regression lines and the identity line. Magenta vertical dashed line represents the scaled prior mean value used in the model simulation. BCD) Single-participant data plotted as dots.

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| One-tailed one-sample Wilcoxon signed-rank tests | | | | | | | | |
|--|--------------------------|----------------|----------|-------------------------|-----------------|--------------|--|--|
| Target TTC | n | V statistic | р | <i>p</i> _{FDR} | Effect size (r) | CI | | |
| 0.55 | 34 | 286 | 0.580 | 0.908 | -0.04 | [-0.35, 1] | | |
| 0.67 | 34 | 220 | 0.908 | 0.908 | -0.26 | [-0.53, 1] | | |
| 0.86 | 34 | 448 | 0.004 | 0.018 | 0.51 | [0.23, 1] | | |
| 1.2 | 34 | 224 | 0.896 | 0.908 | -0.25 | [-0.52, 1] | | |
| | Τv | vo-tailed pair | ed Wilco | kon signe | d-rank tests | | | |
| Target TTC | Target TTC n V statistic | | | <i>p</i> _{FDR} | Effect size (r) | CI | | |
| 0.86 vs. 0.55 | 34 | 425 | 0.029 | 0.023 | 0.43 | [0.07, 0.69] | | |
| 0.86 vs. 0.67 | 34 | 441 | 0.013 | 0.022 | 0.48 | [0.14, 0.72] | | |
| 0.86 vs. 1.2 | 34 | 439 | 0.014 | 0.022 | 0.48 | [0.13, 0.72] | | |

pmEC hexadirectional modulation aligned to pmEC grid orientation

Table S1: Independent ROI analysis for 6-fold symmetry in pmEC

preSMA n-fold modulation aligned to pmEC grid orientation

| One-tailed one-sample Wilcoxon signed-rank tests | | | | | | | |
|--|---|----|-----|-------|-------|------------|--|
| Target TTC | arget TTC Symmetry n V statistic p Effect size (r) Cl | | | | | | |
| 0.86 | 4 | 34 | 363 | 0.135 | 0.22 | [-0.10, 1] | |
| 0.86 | 6 | 34 | 422 | 0.016 | 0.42 | [0.12, 1] | |
| 0.86 | 8 | 34 | 165 | 0.989 | -0.45 | [-0.67, 1] | |

Table S2: Independent ROI confirmatory analyses for 6-fold symmetry in preSMA.

A) pmEC 4-fold symmetry control

| One-tailed one-sample Wilcoxon signed-rank tests | | | | | | | |
|--|--|-----|-------|-------|----------------|------------|--|
| Target TTC | n V statistic p p_{FDR} Effect size (r) CI | | | | | | |
| 0.55 | 34 | 305 | 0.453 | 0.868 | 0.03 | [-0.29, 1] | |
| 0.67 | 34 | 295 | 0.520 | 0.868 | $-8.4x10^{-3}$ | [-0.32, 1] | |
| 0.86 | 34 | 259 | 0.745 | 0.868 | -0.13 | [-0.42, 1] | |
| 1.2 | 34 | 232 | 0.869 | 0.868 | -0.22 | [-0.50, 1] | |

B) pmEC 8-fold symmetry control

| One-tailed one-sample Wilcoxon signed-rank tests | | | | | | | | |
|--|----|-------------|-------|-----------|-----------------|------------|--|--|
| Target TTC | n | V statistic | р | p_{FDR} | Effect size (r) | CI | | |
| 0.55 | 34 | 269 | 0.688 | 0.694 | -0.10 | [-0.40, 1] | | |
| 0.67 | 34 | 312 | 0.407 | 0.694 | 0.05 | [-0.27, 1] | | |
| 0.86 | 34 | 297 | 0.507 | 0.694 | $-1.7x10^{-3}$ | [-0.31, 1] | | |
| 1.2 | 34 | 268 | 0.694 | 0.694 | -0.10 | [-0.40, 1] | | |

C) V1 6-fold symmetry control

| One-tailed one-sample Wilcoxon signed-rank tests | | | | | | | | |
|--|----|---|-------|-------|----------------|------------|--|--|
| Target TTC | n | n V statistic p p_{FDR} Effect size (r) | | | | | | |
| 0.55 | 34 | 295 | 0.520 | 0.554 | $-8.4x10^{-3}$ | [-0.32, 1] | | |
| 0.67 | 34 | 306 | 0.446 | 0.554 | 0.03 | [-0.29, 1] | | |
| 0.86 | 34 | 343 | 0.223 | 0.554 | 0.15 | [-0.17, 1] | | |
| 1.2 | 34 | 290 | 0.554 | 0.554 | -0.03 | [-0.34, 1] | | |

Table S3: A) Independent ROI control analysis for 4-fold symmetry in pmEC. B) Independent ROI control analysis for 8-fold symmetry in pmEC. C) Independent ROI control analysis for 6-fold symmetry in V1.

A) pmEC visual grid-like modulation spatial stability

| One-tailed one-sample Wilcoxon signed-rank tests | | | | | | | | |
|--|----|--|----------------|----------------|---|--------|--|--|
| Target TTC | n | V statisticp p_{FDR} Effect size (r)CI | | | | | | |
| 0.55 | 34 | 595 | $5.8x10^{-11}$ | $5.8x10^{-11}$ | 1 | [1, 1] | | |
| 0.67 | 34 | 595 | $5.8x10^{-11}$ | $5.8x10^{-11}$ | 1 | [1, 1] | | |
| 0.86 | 34 | 595 | $5.8x10^{-11}$ | $5.8x10^{-11}$ | 1 | [1, 1] | | |
| 1.2 | 34 | 595 | $5.8x10^{-11}$ | $5.8x10^{-11}$ | 1 | [1, 1] | | |

B) pmEC visual grid-like modulation temporal stability

| One-tailed paired Wilcoxon signed-rank tests | | | | | | | | |
|--|----|--|-------|-------|------|-----------|--|--|
| Target TTC | n | V statisticp p_{FDR} Effect size (r)CI | | | | | | |
| 0.86 vs. 0.55 | 34 | 429 | 0.004 | 0.012 | 0.53 | [0.25, 1] | | |
| 0.86 vs. 0.67 | 34 | 337 | 0.042 | 0.042 | 0.36 | [0.05, 1] | | |
| 0.86 vs. 1.2 | 34 | 392 | 0.009 | 0.013 | 0.48 | [0.20, 1] | | |

Table S4: A) Clustering of grid orientations across pmEC voxels for each TTC_r. B) Percent of pmEC temporally stable voxels for each TTC_r.

| One-tailed paired Wilcoxon signed-rank tests | | | | | | | | |
|--|----|---|----------------|---------------|------|-----------|--|--|
| Prior mean (s) | n | n V statistic p p_{FDR} Effect size (r) C | | | | | | |
| 0.82 vs. 0.52 | 34 | 589 | $8.1x10^{-10}$ | $2.4x10^{-9}$ | 0.98 | [0.96, 1] | | |
| 0.82 vs. 0.64 | 34 | 582 | $5.1x10^{-9}$ | $6.4x10^{-9}$ | 0.96 | [0.92, 1] | | |
| 0.82 vs. 1.14 | 34 | 581 | $6.4x10^{-9}$ | $6.4x10^{-9}$ | 0.95 | [0.91, 1] | | |

Table S5: Comparison of group average Mean Absolute Error (MAE) across model simulations with different prior mean values

Fixation error across directions per target TTC

| Mixed-Effect Model results | | | | | | | |
|----------------------------|----|------|---------|-------------------------------|--------------|--|--|
| Target TTC (s) | DF | F | p-value | Effect size (ϵ^2) | CI | | |
| 0.55 | 23 | 0.75 | 0.79 | $-7.5x10^{-3}$ | [0.00, 0.00] | | |
| 0.67 | 23 | 1.33 | 0.14 | 9.6 <i>x</i> 10 ⁻³ | [0.00, 0.00] | | |
| 0.86 | 23 | 1.31 | 0.15 | 9 <i>x</i> 10 ⁻³ | [0.00, 0.00] | | |
| 1.2 | 23 | 0.98 | 0.48 | $-4.5x10^{-4}$ | [0.00, 0.00] | | |

Table S6: Fixation error across directions per target TTC.