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Common and distinct roles of frontal midline theta and occipital alpha oscillations in coding temporal intervals and spatial distances Mingli Liang^a, Jingyi Zheng^b, Eve Isham^a, Arne Ekstrom^a

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Abstract

12 Judging how far something is and how long it takes to get there are critical to memory and 13 navigation. Yet, the neural codes for spatial and temporal information remain unclear, 14 particularly the involvement of neural oscillations in maintaining such codes. To address these 15 issues, we designed an immersive virtual reality environment containing teleporters that displace participants to a different location after entry. Upon exiting the teleporters, participants 16 17 made judgements from two given options regarding either the distance they had travelled (spatial distance condition) or the duration they had spent inside the teleporters (temporal 18 19 duration condition). We wirelessly recorded scalp EEG while participants navigated in the virtual 20 environment by physically walking on an omnidirectional treadmill and traveling through 21 teleporters. An exploratory analysis revealed significantly higher alpha and beta power for short 22 distance versus long distance traversals, while the contrast also revealed significantly higher 23 frontal midline delta-theta-alpha power, and global beta power increases for short versus long 24 temporal duration teleportation. Analyses of occipital alpha instantaneous frequencies revealed 25 their sensitivity for both spatial distances and temporal durations, suggesting a novel and 26 common mechanism for both spatial and temporal coding. We further examined the resolution of distance and temporal coding by classifying discretized distance bins and 250ms time bins 27 28 based on multivariate patterns of 2-30 Hz power spectra, finding evidence that oscillations code 29 fine-scale time and distance information. Together, these findings support partially independent coding schemes for spatial and temporal information, suggesting that low-frequency oscillations 30 play important roles in coding both space and time. 31

32 *Keywords:* spatial cognition, temporal cognition, mobile EEG, virtual reality, theta 33 oscillations, alpha frequency

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34	Common and distinct roles of frontal midline theta and occipital alpha oscillations in coding
35	temporal intervals and spatial distances

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Introduction

38 Background

Tracking where we are in space and time is important for both navigation and episodic 39 memory (Eichenbaum & Cohen, 2014; Ekstrom & Isham, 2017; Robin & Moscovitch, 2014; 40 41 Tulving, 2002). However, it is not clear what neural mechanisms are recruited for spatial and temporal coding in humans and whether they share similar coding principles (Ekstrom & Isham, 42 2017; Frassinetti et al., 2009; Walsh, 2003). Movement, either physical or imagined, is a core 43 part of both our sense of space and time, and induces robust hippocampal low-frequency 44 45 oscillations (3-12Hz) in both rats (Vanderwolf, 1969) and humans (Bohbot et al., 2017; Ekstrom et al., 2005; Goyal et al., 2020; Jacobs, 2013; Watrous et al., 2011). Because movement 46 47 typically involves changes in both space and time, one possibility is that low-frequency 48 oscillations play a role in coding both variables.

49 Past investigations have established an important role for hippocampal theta oscillations 50 in coding spatial distance in humans but evidence is lacking for the role of neocortical theta 51 oscillations in distance coding. For example, hippocampal theta power increases linearly with 52 the amount of distance travelled in virtual reality (Bush et al., 2017; Vass et al., 2016), cross-53 regional theta connectivity plays a critical role in judgments of relative spatial distance (Kim et 54 al., 2018), and theta network connectivity differentiates distance from temporal contextual 55 retrieval (Watrous et al., 2013). However, it is not clear whether neocortical theta oscillations 56 can code spatial distance in a similar fashion, and if scalp EEG can reveal such a cortical theta 57 distance code.

58 In addition, while past studies have established a role for low-frequency oscillations in 59 spatial distance coding, their role in representing temporal durations remains less clear. The 60 medial temporal lobes (MTL) of rodents are capable of internally generating representations that 61 track time passage (Itskov et al., 2011; MacDonald et al., 2011; Pastalkova et al., 2008; Wang 62 et al., 2015). Given the strong presence of delta and theta oscillations in MTL, it is possible that 63 low-frequency oscillations contribute to temporal duration coding and that such a time code can 64 manifest in neocortical low-frequency oscillations as well. Past studies have also revealed a role 65 for cortical beta oscillations in supporting duration reproduction in humans, such as the finding that increased alpha-beta coupling strengths yield better time reproduction precision (Grabot et 66

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al., 2019), and higher beta power recorded with scalp EEG predicts longer reproduced 67 durations (Kononowicz & van Rijn, 2015). Therefore, both delta-theta and beta band 68 69 oscillations are strong potential candidates specifically dedicated to temporal duration coding, or both spatiotemporal coding, an issue we seek to resolve here. Beside low-frequency power 70 71 changes, another possible oscillatory timing mechanism is alpha frequency modulation. Alpha frequency variations manifest independently of changes in alpha power (Samuel et al., 2018), 72 73 and alpha frequency modulation has been implicated in the temporal resolution of visual 74 perception (Cecere et al., 2015; Samaha & Postle, 2015). Nonetheless, how alpha frequency 75 fluctuations relate to duration timing remains unclear and unresolved.

76 **Objectives**

77 In this current study, we aim at experimentally dissociating the spatial distance and 78 temporal duration information available to participants. Then, we examine whether and how 79 low-frequency oscillations support spatial distance and temporal duration coding, and whether 80 such spatiotemporal processing shares similar coding schemes. To address these research 81 questions, we developed a teleportation task in an immersive and ecologically enriched virtual 82 environment (Figure 1), largely similar to the experimental design in Vass et al. (2016) and 83 capable of disentangling spatial and temporal information. In this task, participants entered a 84 virtual teleporter, were presented with a black screen for a couple of seconds, and then exited at a different location in the virtual environment. After exiting, participants were prompted to make 85 86 a binary-choice judgment regarding the distance they were transported inside the teleporter (the 87 spatial distance task) or how long the duration was they spent inside the teleporter. By manipulating the distance and duration information independently, we disentangled participants' 88 memory for spatial distance from that of temporal duration. This in turn allowed us to examine 89 90 their neural correlates separately. In addition, participants navigated around the virtual reality by 91 physically walking on an omnidirectional treadmill while wearing a head mounted display. 92 allowing us to study the relationship between cortical oscillations and spatiotemporal processing 93 under more ecologically enriched conditions.

94 Hypotheses

We tested two primary hypotheses. First, for the **within-task difference hypothesis**, we tested whether cortical oscillatory power (2-30Hz) and occipital alpha frequencies responded differently within tasks, i.e., judging short vs. long spatial distance, or short vs. long temporal durations. Second, for the **between-task difference hypothesis**, we tested whether such oscillatory codes differed between tasks, i.e. for spatial distance vs. temporal duration

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- judgments, which might further support the ideas of independent codes (Watrous et al., 2013)
- 101 vs. a common magnitude estimation mechanism (Walsh, 2003) for spatiotemporal coding.
- 102 Together, these analyses allowed us to address to what extent the coding for spatial distance
- and temporal durations involves common vs. distinct neural mechanisms.
- 104

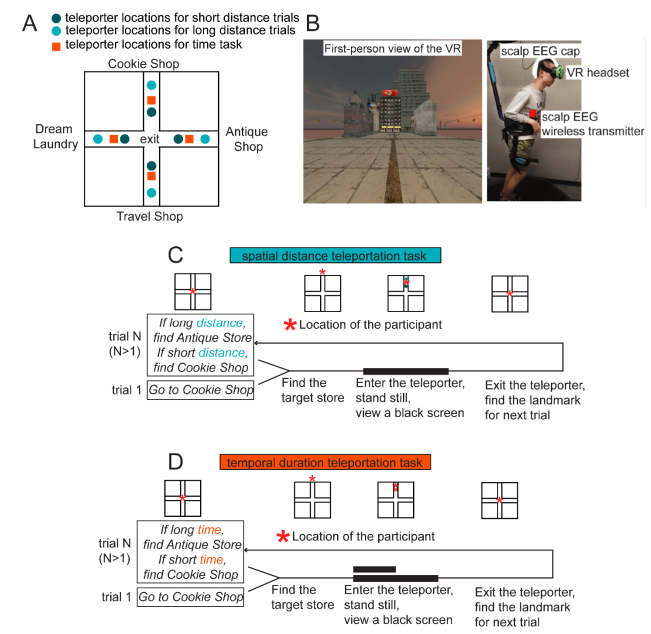




Figure 1. Spatial and temporal teleportation tasks, and virtual reality (VR) setup. (A) Layout of
 the VR and the possible entry locations of teleporters. (B) A view of the virtual environment, and
 the VR-scalp EEG setup. (C) Task flow in the spatial task. Participants were either teleported a

short or long distance inside teleporters. **(D)** Task flow in the temporal task. Participants either

- 110 experienced a short (4 seconds) or long (8 seconds) duration inside teleporters while standing
- . 111 still.
- 112
- 113

Materials and Methods

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115 This study was approved by the Institutional Review Board at the University of Arizona,

and all participants provided informed consent. The data analyzed in this study are available at

117 https://osf.io/3vxkn/.

118 **Participants**

119 We tested 19 adults (7 females, 12 males) from the Tucson community. Because this is the first investigation of its type (scalp-recorded oscillatory correlates of spatiotemporal 120 121 processing), it is difficult to estimate exact effects sizes needed to determine the sample size. 122 Therefore, we based our sample size on a previous study in which we observed movement-123 related changes in low-frequency oscillations during navigation (Liang et al., 2018). Participants 124 received monetary (\$20/hour) and/or class credit for compensation. Prior to testing, participants received a virtual reality training session, which involved 30 minutes of walking on the 125 126 omnidirectional treadmill with a head mounted display on. We implemented the training to 127 screen out participants with potential susceptibility to cybersickness. 128

129 Stimuli, Apparatus and Virtual Reality

130 The virtual environment was constructed with the Unity Engine and rendered with an HTC

131 Vive headset. Immersive walking experiences were simulated with an omnidirectional treadmill

132 (KAT VR Gaming Pro, KAT VR, Hangzhou China). Physical walking motions on the

133 omnidirectional treadmill were translated into movements in the virtual reality.

The size of the virtual environment was 560 x 560 virtual square meters. The layout of the virtual environment was a plus (+) sign (Figure 1A), with four arms extending from the center. Four target stores were placed at the end of each arm (Cookie Shop, Dream Laundry, Antique Store, and Travel Shop). Identical filler buildings were placed along each arm.

The entry point to the teleporters was rendered as a purple circle. When participants "collided" with teleporters in the virtual reality, they initiated a teleportation event. During teleportation, they stood still for a few seconds while viewing a black screen on the headmounted display, and eventually exited at the center of the plus maze.

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143 Behavioral Tasks

144 Participants completed two tasks: a spatial distance task and a temporal duration task.

145 In the spatial task, the teleporters displaced the participants with one of the two possible

spatial distances while the teleportation duration was kept constant. In the temporal task, the

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teleportation process could last a short (4 seconds) or long (8 seconds) duration, while the
teleporters transported the participants a fixed distance. Each task involved 48 trials. **Navigation phase.** At the beginning of a trial, participants started at the center of the plus
maze and navigated to a target store. The target store was either specified for the first trial, or it
needed to be determined for the following trials. When arriving at the target store, participants

entered a dummy teleporter in front of the target store. This involved showing a black screen for
4 seconds and rotating participant's camera angle by 180 degrees. This dummy teleporter was
set up to timestamp participants' arrival times on the EEG and was not used in any subsequent
analyses. If participants arrived at the wrong store, the dummy teleporters sent participants
back to the center of the plus maze and they searched for the store again. During the navigation
phase, no teleporters were visible except for four dummy teleporters in front of four target stores

to detect arrivals at the correct store.

159 **Teleportation phase.** After navigating to the target store, participants then walked up to 160 and entered a new teleporter spawned in front of the target store. In the spatial distance task, for long distance trials, the teleporters spawned 200 virtual meters away from the center of the 161 162 plus maze, and for short distance trials, the teleporters spawned 100 virtual meters away from the center. In the temporal duration task, the teleporters spawned 144 meters away from the 163 164 center. Upon entering the teleporter, participants stood still, with the camera fading to a 165 completely black screen in 200 milliseconds. They viewed the black screen for a specific 166 duration (spatial task: 5.656 seconds, and temporal task: 4 or 8 seconds). Then participants 167 reemerged at the center of the plus maze, with their camera fading from pure black to the view standing at the center of plus maze, in 200 milliseconds. 168

169 **Judgment phase.** After exiting the teleporter, written instructions were provided to the 170 participants by showing a billboard message overlaid on top of the virtual reality view. The 171 instructions were used to decide which target store to visit for the current trial. For the spatial task, instructions were: "If far distance, go find store A. If short distance, go find store B." For the 172 173 temporal task, instructions were: "If long time, go find store A. If short time, go find store B." 174 The instructions in the virtual reality disappeared when participants walked further than 55 175 meters away from the center of the plus maze. By asking participants to judge spatial distance and temporal durations, we ensured that they maintained these two task-relevant variables. 176

Parameters for the behavioral tasks. For the spatial task, the duration of viewing the
black screen was 5.656 seconds for both long distance and short distance trials. Short
distance was defined as teleporting 100 meters and long distance was defined as teleporting
200 meters and (Figure 1C). For the temporal task, the distance teleported was kept constant,

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at 141.4 meters. For short duration trials, participants viewed 4 seconds of a black screen 181 during teleportation, while for long duration trials, they viewed 8 seconds of a black screen 182 183 (Figure 1D). We selected these parameters for our spatial and temporal tasks to ensure the average teleportation speeds were the same between spatial and temporal tasks: the average 184 teleportation speed for the spatial task was $\frac{1}{2}$ × (200 meters/5.656s + 100 meters /5.656s) ≈ 185 26.52 m/s while the average speed for the temporal task was $\frac{1}{2} \times (141.4 \text{ meters}/8s + 141.4)$ 186 meters/4s) \approx 26.51 m/s). This is because movement speed has been shown to affect low-187 188 frequency oscillations (Caplan et al., 2003) and thus we attempted to control for movement 189 speed during teleportation.

190 The order of short/long trials were pseudorandomized across the 48 trials. Short and 191 long teleportation each had 24 trials, with each target store visited 12 times. Two sets of short/long orders were generated so that spatial and temporal tasks did not use the same set 192 193 of long/short sequences. The order of task types, and the short/long sequence sets, were 194 counterbalanced across participants. Before starting the main experiment, participants were 195 shown three examples each: long distance teleportation, short distance teleportation, short 196 temporal duration teleportation, and long temporal duration teleportation. Some participants 197 repeated this practice procedure until they reported understanding the differences between 198 short/long trials.

After each block of 12 trials, participants had the option to take a short break of 3 minutes. When participants took a break, we first asked participants to stand still and relax for 90 seconds on the omnidirectional treadmill while wearing the head-mounted display and viewing a black screen. Then we recorded the 90 second EEG data as the baseline. Pooling across the spatial and temporal tasks, we recorded, on average, 364.74 seconds (SD: 183.64 seconds) of EEG baseline data.

205

206 EEG Acquisition and Preprocessing

The continuous EEG was recorded with a 64-channel BrainVision ActiCAP system, which included a wireless transmission MOVE module, and two BrainAmp amplifiers (BrainVision LLC, Morrisville, NC). We recorded from 64 active electrodes, placed on the scalp according to the international 10-20 system. The reference electrode was located at FCz, and no online filter was applied to the recordings. Before the experimenter proceeded to start the recordings, impedances of all 64 electrodes were confirmed below 5k Ω .

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Preprocessing and analyses were performed with EEGLAB (Makeig et al., 2004), and 213 customized codes in MATLAB. No offline re-referencing or interpolation of electrodes was 214 performed on the continuous data. A 1650th order Hamming windowed sinc finite impulse 215 216 response (FIR) filter was performed for 1-50Hz bandpass filtering on the continuous data using 217 the EEGLAB pop_newfilt() function, with a transition bandwidth of 1 Hz, the passband edges of 218 1 and 50 Hz, and cut-off frequencies (-6dB) of 0.5 and 50.5 Hz. Artifact subspace reconstruction 219 (ASR) was then applied to the filtered continuous data, with the EEGLAB clean asr() function, 220 to repair large amplitude spikes that were 5 standard deviations away from the clean segments 221 of the continuous data.

222

223 EEG Epoching and Segmentation

224 The continuous EEG data were segmented using a time window aligned with the start 225 and ends of teleportation (not including the fade-to-black or fade-to-clear 200ms windows). 226 This segmentation procedure yielded 48 epochs with a length of 5.656 seconds for the spatial 227 task, and 48 epochs with a length of either 4 or 8 seconds for the temporal task. No baseline correction was applied. To keep the number of trials constant across participants, we did not 228 229 reject trials based on incorrect behavioral responses. We did not reject trials based a voltage 230 threshold because we mainly used independent component analysis to correct artifacts, as 231 described below.

232

233 Independent Component Analysis

234 Independent component analysis (ICA) with the infomax algorithm was performed in 235 EEGLAB to correct artifacts. Note that we ran ICA on the artificial "continuous data structure" by concatenating all the data in the distance task, time task, and the resting baseline task. 236 237 Our motivation was data in those three tasks should receive identical ICA correction procedure. 238 We used an automatic component selection procedure, ICLabel (Pion-Tonachini et al., 2019) 239 to avoid experimenter bias in identifying noisy components. Components were rejected automatically if they had labels of "Muscle", "Eye", "Heart", "Line Noise", or "Channel Noise" if 240 their probability was higher than 90% for being one of those labels. On average, 8.84 (13.81% 241 of all components, SD: 3.91) components were rejected. 242

243

244 Time Frequency Analysis

Power measures for delta, theta, alpha, and beta bands. We estimated the

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instantaneous power during the teleportation windows with 6-cycle Morlet Wavelets using code 246 247 from Hughes, Whitten, Caplan, and Dickson (2012). We sampled frequencies from 2 to 30 Hz in 20 logarithmic frequency steps, i.e., 2 Hz, 2.31 Hz, 2.66 Hz, 3.07 Hz, and 3.54 Hz for delta band, 248 249 4.08 Hz, 4.70 Hz, 5.42 Hz, 6.25 Hz and 7.21 Hz for theta band, 8.32 Hz, 9.59 Hz, and 11.06 Hz 250 for alpha band, 12.76 Hz, 14.71 Hz, 16.96 Hz, 19.56 Hz, 22.56 Hz, 26.01 Hz, and 30 Hz for 251 beta band. Zero paddings were added to both ends of the signal to alleviate edge artifacts. No 252 baseline correction was applied to the power estimates. Logarithmic transform with a base of 10 was applied to the obtain power values before averaging. Mean power for each band was 253 254 measured as log power averaged across timepoints within the teleportation window, across 255 frequencies within a band, and across trials of interest.

Cluster-based permutation tests for multiple comparison correction. Cluster-based
 permutation tests (Maris & Oostenveld, 2007) were used to determine the statistical significance
 between the mean power values for short vs. long trials. Correction for multiple comparisons
 was implemented in Fieldtrip. First, to identify uncorrected significant samples,

260 64(electrodes)*4(frequency bands) = 256 Wilcoxon signed rank two-tailed tests were performed

for the power contrasts, alpha = 0.05. Clusters were found by connecting significant sample

262 pairs (electrode x frequency bands) with spatiospectral adjacency (minimum neighbor of

channels was set to 0), and cluster-level statistics were computed using a weighted-sum

264 (Hayasaka & Nichols, 2004) of all the z values returned by Wilcoxon signed rank tests within a

cluster. Second, a surrogate distribution of cluster-level statistics was generated by randomly

shuffling condition labels 1000 times on the subject level and retrieving the maximum cluster-

level test statistic for each permutation. Third, p values of the observed cluster statistics were

obtained by benchmarking to the surrogate distribution. Empirical clusters with a p value smaller
than 0.025 (either left tail or right tail) were be reported.

We chose the nonparametric Wilcoxon signed rank tests over the parametric paired t tests because the normality assumption for t tests was violated. For all the power spectra contrast we conducted, all the power spectra differences showed a distribution different from normal distributions (one sample Kolmogorov-Smirnov test, alpha = 0.05, all p's < 0.01). In the results reported in which we employed the Wilcoxon signed rank tests, medians instead of means were

275 reported.

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Effect sizes calculation. Cohen's d was used as an estimate for effect sizes. For a
 within-participant paired comparisons between condition 1 and condition 2, we estimated the
 effect sizes using the following formula:

$$d = \frac{mean(condition1) - mean(condition2)}{std(condition1 - condition2)}$$

279

Frequency measures for alpha (8-12 Hz) band. To estimate alpha frequency, we used 280 a frequency sliding technique (Cohen, 2014) to estimate the alpha frequency fluctuations. We 281 first used a 125th order finite impulse response (FIR) 8-12 Hz bandpass filter (using MATLAB 282 283 firls() function) on the segmented EEG data, with a transition bandwidth of 1.2 and 1.8 Hz, the 284 passband edges of 8 and 12 Hz, and cut-off frequencies (-6dB) of 7.12 and 12.98 Hz. We then 285 employed the Hilbert Transform on the filtered segmented EEG data to obtain the instantaneous 286 phase estimates of alpha oscillations during teleportation windows. Instantaneous frequencies at timepoint *t* were estimated as 287

288 289 $f_t = \frac{\varphi_t - \varphi_{t-1}}{2\pi} \times s \tag{1}$

where *f* is the estimated instantaneous alpha frequency, φ is the estimated phase, and *s* is the EEG sampling rate. Here, we defined and estimated the instantaneous frequencies based on how many cycles the phase of alpha oscillations could go through in 1 second. Then, to smooth the frequency estimates, we applied a 10th order median filter. We dropped the frequency estimates for the first 100ms and last 100ms for every trial because of potential inaccurate estimates of frequencies at the edges of signal.

We selected the following occipital electrodes to analyze their alpha frequency based on two criteria: visible alpha prevalence in the raw traces and an identical cluster of occipital electrodes to what we used in our past study (Liang et al, 2018). These 18 electrodes corresponded to: Pz, P3, P7, O1, Oz, O2, P4, P8, P1, P5, PO7, PO3, POz, PO4, PO8, P6, P2, and Iz.

Alpha frequency for each behavioral task is measured as alpha frequency estimates averaged across timepoints during the windows of interest, averaged across electrodes of interest, and averaged across trials of interest. To compare the alpha frequency variations between two conditions, we submitted the averaged alpha frequencies of 19 participants to twotailed Wilcoxon signed rank tests (alpha = 0.05.) Six Wilcoxon signed rank tests were conducted,

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and the p values reported in the results section were FDR corrected (Benjamini & Yekutieli,

2001; Groppe, 2021), with the false discovery rate set to 0.05.

308

309 Classification Analyses

310 Binary classification of the duration/distance types. To further confirm the role of frontal midline delta-theta oscillations in spatial and temporal judgments, a binary support vector 311 312 machine (SVM) classifier was used to decode the types of teleportation using power of delta, theta, alpha and beta band, averaged at specific electrodes. For delta power, theta power, and 313 alpha power, 4 electrodes around frontal midline region were selected (Fz, FC1, Cz, FC2.) For 314 beta power, all available electrodes (64 electrodes) were chosen. Binary SVM classifiers were 315 316 implemented in MATLAB, with the function *fitcsvm()*, with the kernel function set up as linear. Three decoding tasks on a within-subject level were implemented: 1) decoding whether the trial 317 was from the teleportation trials that travelled short or long distance, 2) decoding whether the 318 319 trial was from short duration trials, or the 4-8 second portions of long durations trials in the time 320 task, and 3) decoding whether the trial was from short duration trials or the 0-4 second portions 321 of long durations trials. The ratio of train-test split for each iteration was 67%-33%. The trainingtesting sampling procedure was reiterated 1000 times for each participant, and for each 322 323 decoding task. An accuracy percentage score was calculated using the predicted and actual 324 labels of the testing data. The final decoding accuracy scores for 19 participants were 325 submitted to two-tailed Wilcoxon signed rank tests, against the null hypothesis that the decoding 326 accuracy was 50%. In total, 12 tests were conducted in the binary classification analysis, and 327 the p values were FDR corrected (Benjamini & Yekutieli, 2001; Groppe, 2021), with the false 328 discovery rate set to 0.05.

329 Additionally, we implemented a between-task classifier (space vs. time tasks) on an 330 intersubject level. We combined trials from the space task and the time task across 19 331 participants, resulting in a dataset of 912 trials. Then, we tested whether we could successfully 332 decode the task labels using the 912-trial dataset. By performing the classification on an intersubject level (with the task orders were counterbalanced), we avoided the possible 333 334 confound of systematic drift over the course of experiment, which could have affected our 335 decoding accuracy due to the blocked nature of the spatial vs. temporal judgments in our design 336 (Benwell et al., 2019). For features used for training classifiers, we employed the 2-30 Hz power spectra from 64 electrodes averaged within each trial, resulting in 20*64 = 1280 features. The 337 338 ratio of train-test split for each iteration was 67%-33%. The train-test split was repeated 100

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times. To determine the statistical significance of decoding accuracy, we submitted the
accuracies from 100 iterations to a two-tailed Wilcoxon signed-rank test against the null
hypothesis of 50%.

Fine-scale time decoding analyses. To examine whether continuous time codes were present in the scalp EEG signal, SVM classifier was trained to decode times beginning at the onset of teleportation using the 2-30 Hz power spectra from 64 electrodes. The SVM algorithm was implemented in MATLAB using the *fitcecoc(*) function, with coding style as '*onevsall*', and other parameters as default.

250-millisecond timebins were extracted by discretizing 2-30 Hz power estimates. The
size of timebins was chosen as the same one used by Bright et al. (Bright et al., 2020).
Therefore, short/long distance teleportation trials (5.656 seconds) yielded 22 bins (22 * 250ms =

5.5 seconds, the last 156ms of data were dropped), short temporal duration trials (4 seconds)
yielded 16 bins, and long temporal duration trials (8 seconds) yielded 32 bins. For the resting
baseline data (90 seconds long for each resting session), we broke 90 seconds into continuous
segments of 4 seconds, and from there, each 4s of baseline data were segmented into 16 bins.

354 Power estimates within each time bin were averaged over time, and the resulting power 355 spectra within each bin were used to trained classifiers. The number of features were 20 356 frequencies x 64 electrodes = 1280 features. For each classification iteration, train-test split ratio was 75%-25%. To increase the independence between training sets and testing sets, a 357 358 consecutive block of trials was reserved as the testing data, and the rest of data was used for training. Given our way of splitting the data, we were able to reiterate the classification 359 360 procedure limited number of times: for the distance task, the procedure was repeated 37 times: 361 for the short interval and long interval trials, 19 times; and for the baseline task, 16 times.

362 We calculated the accuracy score by summing how many correct predictions were made 363 in 100 iterations for each timebin label. The accuracy scores were then averaged across all iterations yielding a final accuracy score for each participant. Given that number of timebins 364 were different across the distance task, time task and baseline task, comparisons between 365 them would be difficult. We standardized the accuracy scores as the accuracy ratios by dividing 366 them against the chance level performance (ratios = $\frac{classification \ accuracy}{chance \ level}$). For the distance task 367 368 time decoder, the chance level was 1/22 = -4.55%; for decoding time in short temporal duration 369 trials, the chancel level was 1/16 = 6.25%; for decoding time in long duration trials, the chance 370 level was 1/32 = 3.125%, and for decoding time in the baseline data, the chance level was 1/16371 = 6.25%.

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To test whether we successfully decoded fine-scaled temporal information above chance, 372 373 we submitted the standardized accuracy ratios for 19 participants to a two-tailed Wilcoxon 374 signed rank tests against the null hypothesis that the accuracy ratios were different from 1. Ten signed rank tests were performed for this hypothesis, and the p values were FDR corrected 375 376 (Benjamini & Yekutieli, 2001; Groppe, 2021), with the false discovery rate set to 0.05. 377 To visualize the time decoder performance and the posterior probability distribution, we 378 calculated a N x N (N = the number of time bins) matrix to summarize the time decoder 379 prediction outputs. For element (i,j) in the matrix, the value represented the probability of a time 380 bin #i was predicted as time bin #j.

381 Calculation of Absolute Decoding Errors in the Fine-scale Time Analysis. After 382 retrieving the posterior probability distribution of decoding responses (the NxN matrix, where N 383 is the number of bins), we calculated the absolute decoding errors for each time bin, using the 384 following equation: $errors = \sum_{i=1}^{n} p_i * |i - j| * binSize$, where n is the number of bins, i are the possible decoder responses, p_i is the posterior probability for response i, the ground truth bin 385 index is j, and *binSize* is the size of time bin. After obtaining the decoding error curve (as a 386 function of the ground truth bin labels), we fitted the error curve with linear regression. The p 387 388 values of the slope were reported in the results section.

Fine-scale distance decoding analyses. To examine whether continuous distance codes were also present in the scalp EEG power, we discretized data from spatial distance teleportation trials into multiple small "distance" bins and trained SVM classifiers with 2-30Hz power spectra averaged within each distance bin.

393 To avoid the confounded decoding of fine-scale distance and time, we selected data with only maximal overlap in conceptual distance updating but with zero overlap in the temporal 394 395 dimension. We selected the 0-2.828s portions of short distance trials, and 2.828s-4.242s 396 portions of long distance trials. While they did not overlap in time ranges, they conceptually covered the same range of spatial distance (see Figure 6A). After the data selection, the 2-397 398 30Hz power series of both short and long distance trials were discretized into 11 distance bins, 399 with each distance bin covering 4.42m of distance. For short distance trials, each distance bin 400 occupied 248ms (with a sample rate of 500Hz, 248ms = 124 sampling points), and for long distance trials, each distance bin occupied 248/2 = 124ms (124ms = 62 sampling points). 401 Power estimates within each time bin were averaged. We trained multiclass SVM classifiers 402 403 with 1280 power spectra features (64 electrodes x 20 frequency). For each classification iteration, 75% of the trials were selected as the training data and 25% of the trials were 404

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405	reserved as the testing data. To increase the independence between training sets and testing
406	sets, a consecutive block of trials was reserved as the testing data, and the rest of data was
407	used for training. We were able to reiterate the classification procedure 37 times. The resulting
408	classification accuracy ratios were averaged across the 37 iterations for each participant, and
409	the 19 participant scores were submitted two-tailed Wilcoxon signed rank tests testing whether
410	they were significantly different than 1.
411	
412 413	Results
414	Participants correctly judged spatial and temporal teleportation durations with high
415	accuracy
416	Participants performed well above chance in both the spatial and temporal teleportation
417	tasks. For the spatial task, out of 48 trials, participants on average made 0.68 errors (SD = 0.89)
418	in judging how far the distance they travelled at the first attempt. For the temporal task, out of
419	48 trials, participants on average made 1.79 errors (SD = 2.51) in judging how long they spent
420	inside teleporters. On average, participants finished the spatial task within 53.46 (SD = 12.73)
421	minutes and the temporal teleportation task within 52.35 (SD = 9.24) minutes.
422	
423	Within-task comparisons: Longer distances traveled associated with decreases in alpha
424	and beta power compared to shorter distance traversals
425	We first tested the within-task difference hypothesis in the spatial distance task. We
426	compared delta, theta, alpha and beta power between short distance and long distance
427	teleportation trials, and used a cluster-based permutation test for multiple comparison correction.
428	When comparing short distance vs. long distance trials, the permutation test returned a cluster
429	with a p value of 0.015. For short distance trials, we found higher alpha power at central
430	electrodes (Figure 2A, Pz, CP2, Cz, CPz, Cohen's d: 0.55, averaged log10 alpha power for
431	short distance: [median \pm SD] = 4.99 \pm 0.34, long distance: 4.91 \pm 0.32) and higher beta power over
432	central-posterior electrodes (Cohen's d: 0.91, averaged beta power for short distance:
433	[median \pm SD] = 4.51 \pm 0.26, long distance: 4.50 \pm 0.26.) These findings support a possible role for
434	alpha and beta power changes in spatial distance coding.
435	

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436 Within-task comparisons: Longer temporal durations were associated with frontal delta-

437 theta-alpha power and global beta power decreases compared to shorter temporal

438 durations

439 We then tested the within-task difference hypothesis for temporal duration teleportation by 440 comparing the power spectra between short duration and long duration trials (Figure 2B). The 441 cluster-based permutation test returned a positive cluster (p < 0.001). This effect was most pronounced over frontal midline electrodes for delta power (Cohen's d: 1.03, short duration: 442 443 $[median\pm SD] = 4.47\pm 0.22$, long duration: 4.42 ± 0.23), over frontal electrodes for theta power 444 (Cohen's d: 0.97, short duration: [median \pm SD] = 4.86 \pm 0.19, long duration: 4.83 \pm 0.20), and over frontal electrodes for alpha power (Cohen's d: 0.98, short duration: [median \pm SD] = 4.35 ± 0.24 , 445 long duration: 4.32±0.25). We also found global beta power changes (Cohen's d: 1.63, short 446 duration: $[median\pm SD] = 4.59\pm 0.25$, long duration: 4.55 ± 0.26 .) 447 448 To further confirm the role of frontal midline theta oscillations in duration timing, we trained 449 a binary classifier to decode types of temporal durations in the teleporter (Figure 3). We 450 successfully decoded whether a trial was a short duration trial or the 4-8 s portion of a long 451 duration trial (Figure 3A, classifiers trained with frontal-midline delta power: [median±SD] = 452 64.40±9.89%, frontal-midline theta: 65.42±11.68%, frontal-midline alpha: 69.34±10.89%, global beta: 88.76±7.51%, all p_{corrected} = 0.002). However, we could not decode the distance travelled in 453 454 the teleporter significantly above chance (Figure 3B, classifiers trained with frontal-midline delta 455 power: $52.93 \pm 5.09\%$, p_{corrected} = 0.06, theta: $52.26 \pm 6.77\%$, p_{corrected} = 1, alpha: $50.88 \pm 4.47\%$, 456 $p_{corrected} = 1$, beta: 52.94±5.95%, $p_{corrected} = 1$), suggesting frontal midline delta-theta-alpha power, 457 and global beta power alone contained sufficient information regarding the temporal duration 458 being coded but not the distance traveled.

459 As an additional control analysis, we trained the same classifier with frontal-midline delta-460 theta-alpha power and global beta power to discriminate the 0-4s portion of the long duration trials from the short duration trials. This served as a control because participants could not have 461 462 known what types of durations they experienced until they crossed the 4s threshold within the teleporter. Indeed, the classifier was not able to decode whether the trials were short duration 463 trials (4s) or the 0-4s portion of long duration trials (Figure 3C, delta: 50.33±5.88%, theta: 464 465 49.16±5.33%, alpha: 50.34±7.25%, beta: 48.04±5.94%, all p_{corrected} > 0.05). Together, these 466 findings support a general role for global beta power changes in spatiotemporal processing, and 467 a unique role of frontal midline delta-theta-alpha oscillations, in coding temporal durations.

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469 Between task comparisons: Spatial and temporal teleportation did not induce focal

470 differences in delta, theta, alpha, or beta power

To test our between-task hypothesis regarding differences in oscillatory codes between spatial and temporal tasks, we compared the power spectra among spatial, temporal and

473 baseline tasks (Figure 2C, 2D.)

474 For both contrasts (*distance task* > *baseline; time task* > *baseline*), the cluster-based permutation tests returned a significant positive cluster with p values < 0.001. The effect was 475 476 most pronounced over frontal midline electrodes for delta power (Cohen's d for distance vs 477 baseline: 0.60, distance-baseline: [median \pm SD] = 0.12 \pm 0.27, Cohen's d for time vs baseline: 0.77, time-baseline: [median \pm SD] = 0.12 \pm 0.15), over frontal electrodes for theta power (Cohen's 478 d for distance vs baseline: 1.04, distance-baseline: [median±SD] = 0.07±0.07, Cohen's d for 479 480 time vs baseline: 1.01, time-baseline: [median±SD] = 0.05±0.08), and over frontal and occipital electrodes for alpha power (Cohen's d for distance vs baseline: 0.82, distance-baseline: 481 482 $[median\pm SD] = 0.20\pm 0.18$, Cohen's d for time vs baseline: 0.76, time-baseline: $[median\pm SD] =$ 0.10±0.21). We also found widespread increases in beta power (Cohen's d for distance vs 483 baseline: 1.81, distance-baseline: [median \pm SD] = 0.16 \pm 0.08, Cohen's d for time vs baseline: 484 485 1.80, time-baseline: [median \pm SD] = 0.14 \pm 0.07). The findings suggest that compared to a passive baseline, participants showed distinct oscillatory profiles while maintaining 486 spatiotemporal information during the teleportation tasks, which was consistent with their high 487 performance in the behavioral tasks. 488 489 Next, we asked whether the power spectra profiles differed between the spatial distance

and temporal duration task (Figure 2E). The cluster-based permutation test did not reveal any
 clusters with a p-value lower than threshold. This suggests that the spatial and temporal
 teleportation tasks did not differ in overall power when compared within each of the canonical
 frequency bands (delta, theta, alpha and beta bands).

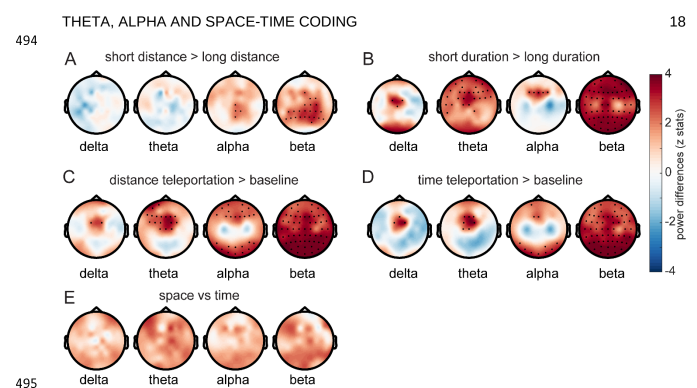


Figure 2. Oscillatory fluctuations present during spatial distance and temporal duration 496 teleportation. (A) Short distance teleportation trials resulted in increased alpha and beta power 497 498 compared to long distance trials. (B) Short duration teleportation trials resulted in increased frontal midline delta-theta-alpha power increases, and global beta power increases compared to 499 long duration trials. (C,D) Spatiotemporal coding was associated with frontal delta-theta, frontal 500 501 and posterior alpha, and global beta power increases compared to resting baseline. (E) No 502 power differences were observed within the canonical frequency bands between the distance task and the time task. **Notes:** Black dots are electrodes considered significant after multiple 503 504 comparison correction. Colors represent the Wilcoxon signed rank tests z statistics.

505

506 Between task comparison: Successful decoding of spatial and temporal trials based on

507 single-trial multivariate patterns of power

508 It could be possible that spatial and temporal coding did not differ in terms of power

509 changes in focal frequency bands; instead, spatiotemporal coding might differ in the multivariate

510 patterns across electrodes and frequencies in a manner that generalizes across participants.

511 To test this possibility, we used multivariate power features to classify whether trials were from

the spatial or temporal task. The classifier revealed above chance classification of task labels

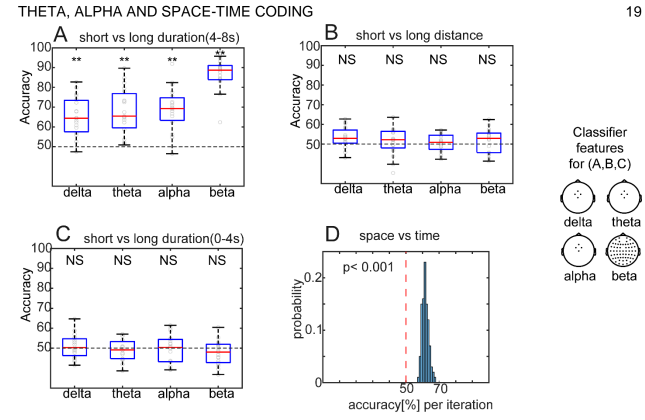
513 (Figure 3D, median = 61.46%, SD over 100 iterations = 1.97%, Wilcoxon signed-rank test, z =

514 8.68, p < 0.001.) These findings suggest the single-trial multivariate patterns significantly

515 differed between spatial and temporal tasks in a manner that generalized across participants.

516 The findings together support a notion of a partially independent space-time code.

517



518

519 Figure 3. Successful within-task (A-C) and between-task (D) decoding using power as features. 520 (A) Different durations (short vs. long) could be decoded from frontal delta, theta, alpha and global beta power separately. (B) Different distances (short vs. long) could not be decoded from 521 frontal midline delta-theta, alpha or global beta power. (C) As a control analysis, decoders were 522 not able to differentiate whether a trial was from short duration trials, or from the 0-4s segments 523 524 of long duration trials. (D) When aggregating trials across participants, we were able to decode whether a trial was in the space or time condition based on the single-trial multivariate patterns 525 of power. The histogram of classification accuracies based on 100 iterations is shown. Notes: **, 526 all $p_{FDR} = 0.002$. Each circle represents a participant in A-C. 527 528

529 Alpha frequency modulation: A common mechanism for spatial and temporal judgments

530 We hypothesized that occipital alpha frequency modulation could be an additional form of

distance and duration coding in our teleportation task, as suggested by (Cao & Händel, 2019;

532 Samaha & Postle, 2015). To test this idea, we first assayed whether there were differences in

occipital alpha frequencies during the teleportation tasks compared to the task-irrelevant resting

baseline. Both spatial and temporal teleportation tasks showed faster occipital alpha

frequencies than the baseline (Figure 4A, spatial task: $[median\pm SD] = 10.23\pm 0.30$ Hz, temporal

task: 10.13±0.25Hz, baseline: 10.00±0.26Hz; spatial task vs. baseline: Wilcoxon signed rank

test, z = 3.74, $p_{corrected} = 0.001$; temporal task vs. baseline: z = 3.78, $p_{corrected} = 0.001$.) These

538 findings suggest that occipital alpha frequencies were significantly altered during spatiotemporal

539 coding compared to a resting baseline.

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540 Second, we asked whether occipital alpha frequency differed between the spatial and 541 temporal tasks. Comparing across all participants, the spatial distance task showed significantly 542 faster occipital alpha compared to the temporal teleportation task (Figure 4A, z = 2.62, p_{corrected} = 0.026) The findings of differences in alpha frequencies between spatial and temporal 543 teleportation tasks might reflect another distinction in oscillatory codes for spatiotemporal 544 545 information. 546 Therefore, we asked whether the observed occipital alpha frequencies were sensitive to 547 distance and duration information. We first compared the averaged alpha frequency at occipital 548 electrode sites for short vs. long distance trials. When comparing across participants, results 549 revealed that occipital alpha oscillations were of higher frequency for short distance trials 550 compared to long distance trials (Figure 4B, short distance: [median±SD] = 10.26±0.29Hz, long 551 distance: 10.20 ± 0.30 Hz, z = 3.38, p_{corrected} = 0.003). Occipital alpha frequency also varied 552 between short and long temporal duration trials. Occipital alpha frequency was faster for short 553 duration trials than the 4-8s portion of long duration trials (Figure 4C, short temporal duration: 554 $[median\pm SD] = 10.28\pm 0.24Hz$, long temporal duration(4-8s): $10.00\pm 0.31Hz$, z = 3.58, $p_{corrected} =$ 555 0.002.) 556 As a control analysis, we tested whether there were differences in occipital alpha 557 frequencies for short duration trials vs. the 0-4s portion of long duration trials. The alpha

frequencies for short duration trials vs. the 0-4s portion of long duration trials. The alpha frequencies did not differ (Figure 4D, short temporal duration: [median \pm SD] = 10.28 \pm 0.24Hz, long temporal duration (0-4s): 10.24 \pm 0.23Hz, z = 0.76, p_{corrected} = 1.) Together, these findings support alpha frequency modulation as a shared mechanism for coding spatial distance and temporal durations.

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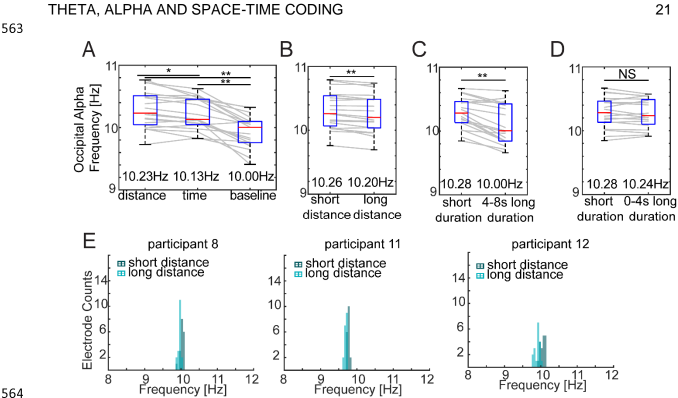


Figure 4. Occipital alpha frequency modulation as a shared mechanism for both spatial and 565 temporal coding. Medians across participants are shown under the box plots. (A) The spatial 566 and temporal tasks showed faster alpha frequency than baseline, and the distance task showed 567 568 faster alpha frequency than the time task. (B) In the distance task, traveling a short distance resulted in faster alpha than traveling a long distance. (C) In the time task, short duration trials 569 resulted in faster alpha than long duration trials. (D) No differences were found between short 570 571 duration trials and the 0-4s portion of long duration trials. (E) Histograms of alpha frequencies at 572 18 occipital electrodes during the distance task. Data from three example participants were shown. **Notes**: **: $p_{FDR} < 0.01$. *: $p_{FDR} < 0.05$. NS: not significant. 573

574 575

576 **Fine-scale temporal information was decoded from multivariate patterns of 2-30Hz power**

577 spectra

578 We next tested whether temporal duration codes might be present in the EEG data at a

579 finer scale, inspired by Bright et al. (2020), for example, at the level of 250 milliseconds.

580 Therefore, we trained classifiers on 2-30Hz power to decode times since onset of teleportation.

- 581 We were able to decode fine-scale temporal information from distance teleportation trials
- significantly above chance (Figure 5A, accuracy: [median±SD] = 10.34±1.32%, accuracy ratios:
- [median \pm SD] = 2.27 \pm 0.29%, Wilcoxon signed rank test, z = 3.82, p_{corrected} < 0.001), from short
- duration trials (Figure 5B, accuracy: [median±SD] = 13.87±1.67%, accuracy ratios: [median±SD]
- $= 2.22 \pm 0.27$, z = 3.82, $p_{corrected} < 0.001$), and from the long duration trials as well (Figure 5C,
- accuracy: $[median\pm SD] = 6.99\pm 0.95\%$, accuracy ratios: $[median\pm SD] = 2.24\pm 0.30$, z = 3.82,
- $p_{corrected} < 0.001$). As a control analysis, we applied the fine-scale time decoder for data
- 588 obtained in the baseline task. The decoder was able to decode time from the baseline data

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marginally better than chance after multiple comparison correction (accuracy: [median±SD] = 589 590 $7.50 \pm 1.87\%$, accuracy ratios: [median \pm SD] = 1.20 ± 0.30 , z = 2.37, p_{corrected} = 0.052). However, time decoding performance for the baseline task was significantly worse than those in the 591 592 temporal and distance tasks (baseline < distance task, baseline < short duration trials, baseline 593 < long duration trials; all z = -3.82, p < 0.001.) These findings suggest the intriguing possibility 594 that fine-scaled temporal codes are embedded in low-frequency oscillations. 595 We note that following entry into the teleporter, participants exhibited a P300-like ERP 596 response (Polich, 2007) at Cz electrode. Therefore, we repeated the fine-scaled time 597 classification analyses, with the grand averaged EEG traces subtracted from every trial. After 598 removing the grand ERP responses, we were still able to successfully decode continuous-like temporal information from the distance teleportation trials (accuracy: $[median\pm SD] =$ 599 600 11.08 \pm 1.29%, accuracy ratios: [median \pm SD] = 2.44 \pm 0.28, Wilcoxon signed rank test, z = 3.82, $p_{corrected} < 0.001$), from the short duration trials (accuracy: [median±SD] = 15.52±1.93%, 601 602 accuracy ratios: [median \pm SD] = 2.48 \pm 0.31, z = 3.82, p_{corrected} < 0.001), and from the long duration trials (accuracy: $[median\pm SD] = 8.22\pm 1.08\%$, accuracy ratios: $[median\pm SD] =$ 603 2.63 ± 0.35 , z = 3.82, p_{corrected} < 0.001.) 604 Further, to exclude the possible contribution of movement-related in early onsets of a trial, 605 606 we removed the first second of teleportation epochs and repeated the fine-scale time decoding 607 analyses. We were again able to successfully decode fine-scale time information from distance 608 teleportation trials above chance (accuracy: $[median\pm SD] = 8.80\pm 1.05\%$, accuracy ratios: $[median\pm SD] = 1.58\pm 0.19$, z = 3.82, $p_{corrected} < 0.001$), from short duration trials (accuracy: 609 $[median\pm SD] = 12.65\pm 1.97\%$, accuracy ratios: $[median\pm SD] = 1.52\pm 0.24$, z = 3.82, $p_{corrected} < 1.52\pm 0.24$, $p_{corrected} < 1.52\pm 0.24$, z = 3.82, $p_{corrected} < 1.52\pm 0.24$, $z = 3.82\pm 0.24\pm 0.24$, 610 611 0.001), and from long duration trials above chance as well (accuracy: [median±SD] = 612 $5.67 \pm 0.89\%$, accuracy ratios: [median \pm SD] = 1.59 ± 0.25 , z = 3.82, p_{corrected} < 0.001.) 613

614 Decoding errors linearly increased as time progressed forward

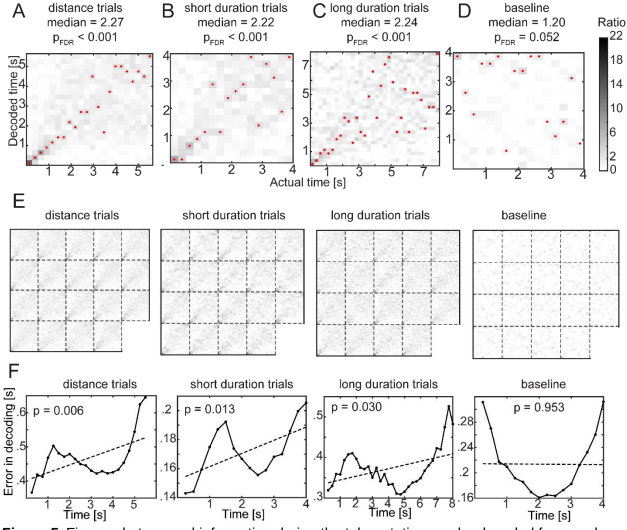
615 We noticed a qualitative pattern that the decoding responses were less precise as time 616 progressed forward in the posterior probability distribution of time decoding responses. To 617 guantitatively test this, we calculated the absolute decoding errors for each timebin and fitted 618 the error curves with a linear regression model (Figure 5F). Results of the linear regression 619 fitting indicated that the decoding errors were significantly larger for later time bins; this effect 620 was found in the distance trials, short duration trials, long duration trials, but not in the baseline 621 task (for distance trials: slope [estimate, standard error (SE)] = [0.02, 0.01], t = 3.03, p = 0.007; 622 for short duration trials: slope [estimate, SE] = [0.01, 0.003], t = 2.70, p = 0.017; for long

23

THETA, ALPHA AND SPACE-TIME CODING duration trials: slope [estimate, SE] = [0.01, 0.003], t = 2.56, p = 0.016; for the baseline task:

slope [estimate, SE] = [-0.0004, 0.01], t = -0.04, p = 0.97.) The results suggest that the fine-

- scale temporal information revealed by the decoders are aligned with the human behavioral
- 626 findings of increased variability for longer reproduced durations (lvry & Hazeltine, 1995; Rakitin
- 627 et al., 1998). We discuss the implications in discussion.
- 628



630 Figure 5. Fine-scale temporal information during the teleportation can be decoded from scalp 631 EEG 2-30 Hz power spectra. Heat maps visualize the posterior probability distribution of the decoder responses. High classification accuracy is indicated by dark colors on the diagonal. 632 (A,B,C,D) Fine-scale timing information can be decoded from 2-30 Hz power in the distance 633 task and time task, with accuracies significantly higher than chance level and higher than the 634 635 baseline task. Medians of accuracy ratios across 19 participants were reported. (E) Decoder response probability distributions from 19 participants. Each sub square displays the time 636 decoding heatmap from one participant. (F) Decoding errors linearly increased as time 637 progressed in the spatial and temporal tasks, but not in the baseline task. Dashed lines indicate 638 639 the linear regression fitting models of the decoding errors. Notes: Units of the colorbar are accuracy ratios. Red dots mark the highest posterior probability in decoder responses. 640

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Fine-scale distance information was also present in multivariate patterns of 2-30Hz power

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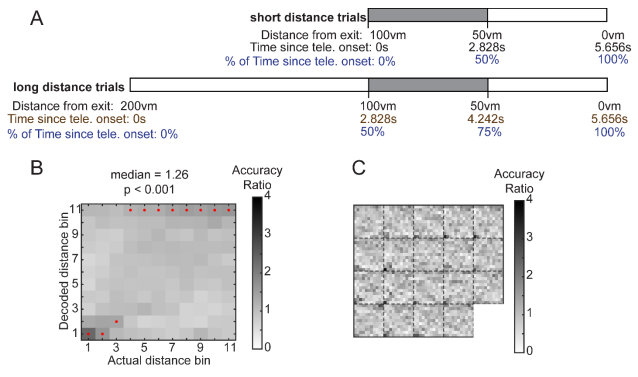
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643 Given our findings with fine-scale temporal information, we also tested whether fine-

scale distances could be decoded using the same approach. Indeed, we found that the

classifiers were able to decode fine-scale distance information from the spatial task (Figure 6A,

- accuracy: [median±SD] = 11.45±1.60%, accuracy ratios: [median±SD] = 1.26±0.18, Wilcoxon
- signed rank test, z = 3.70, p < 0.001). The findings of the fine-scale distance code support the
- 648 possibility that participants linearly updated their spatial position inside teleporters. The
- 649 demonstrations of both fine-scale distance and temporal codes in the multivariate power spectra
- 650 patterns reveal another common aspect that exists in spatiotemporal coding.



651 652

Figure 6. Fine-scale distance information during teleportation can be decoded from multivariate 653 654 power patterns. Heat maps visualize the posterior probability distribution of the decoder 655 responses. (A) Decoding fine-scale distance information while taking care of the temporal 656 confound. To minimize the dependence between temporal and distance information, we selected data (the shaded portions) from both short distance trials and long distance trials that 657 658 had zero overlaps in the temporal dimension. (B) Fine-scale distance information could be decoded in the distance task. (C) Posterior probability distributions plotted for each participant. 659 Each sub square displays the distance decoding heatmap from a participant. Notes: Red dots 660 661 mark the highest posterior probability in decoder responses. 662

- 663
- 664

Discussion

- In the current study, we tested whether neural oscillations recorded at the scalp supported
- 666 maintenance of spatial distance and temporal duration information. Decades of research
- support a role for low-frequency oscillations, both in cortex and hippocampus, in coding spatial
- 668 information during navigation (McFarland et al., 1975; Vanderwolf, 1969; Kropff et al., 2021; for
- reviews, see Jacobs, 2013; Watrous et al., 2011). To attempt to disentangle space and time,

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whose changes are strongly intertwined in movement speed, participants experienced 670 teleportation of different spatial distance and temporal durations in the absence of any optic flow 671 672 or other sensory input to provide cues about speed, similar to the design in Vass et al. (2016). Results from power spectra analyses suggested the sensitivity of central-posterior alpha power 673 674 and global beta power for spatial distances, and a role of frontal theta and global beta power changes for temporal duration. Furthermore, the analysis of instantaneous alpha frequencies 675 676 revealed a robust association between alpha frequency and magnitudes of distances and 677 durations, suggesting alpha frequency modulation as a potential common mechanism for spatial 678 and temporal coding. Classifiers trained on power spectra further support the hypothesis that 679 both distance and temporal information could be decoded from scalp EEG signals at a fine-680 scale resolution.

Given that hippocampal delta-theta power display a distance code (Bush et al., 2017; 681 Vass et al., 2016), and a connectivity between rodent's prefrontal and hippocampal theta during 682 683 mobility (Siapas et al., 2005; Young & McNaughton, 2009), we are surprised to find that the cortical delta-theta power did not exhibit significant differences between short distance and long 684 685 distance trials. This null finding cannot be explained by the failure of task design, or the absence of spatial coding during the teleportation period. This is because participants demonstrated high 686 687 accuracy in identifying distances travelled upon exiting the teleporters, and power spectra 688 analyses revealed significantly different oscillatory profiles for the distance task compared to 689 baseline (Figure 2C). What could lead to such a disconnect? Here, we offer three speculations on the null findings linking cortical theta and spatial distance coding. One possibility is that 690 prefrontal theta oscillations are phase locked but not amplitude locked to hippocampal theta 691 (Young & McNaughton, 2009), and therefore phase information in frontal theta but not power 692 693 changes code spatial distance duration (see Watrous et al., 2013, for an example of this). This 694 is an issue we cannot address in the current study because scalp EEG does not give reliable 695 access to hippocampal signals. A second possibility is that frontal midline theta may be locked 696 to the temporal-processing or memory-related components, but not the movement-related 697 components, of hippocampal (HPC) theta oscillations (Goyal et al., 2020; Watrous et al., 2013). 698 A third possibility is that hippocampal movement-related theta oscillations manifest in the cortex within the traditional alpha band (8-12Hz) consistent with the alpha frequency modulation we 699 700 observed for both spatial and temporal judgments. The third interpretation is consistent with 701 recent reports (Aghajan et al., 2017; Bohbot et al., 2017; Goyal et al., 2020) that hippocampal 702 movement-related theta oscillations, particularly during real-world movements, manifest most

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prominently above 8Hz, which would align with the frequency range of traditional alpha band (8-

12 Hz) rather than theta band (4-8 Hz).

705 Our results supporting a role for frontal delta-theta power but not distance coding have 706 important implications. In the power spectra analysis, we found frontal midline delta-theta and 707 frontal alpha power sensitive to the temporal durations, while central-posterior alpha power was 708 sensitive to the distance information. The results provide further evidence for partially 709 independent codes for space and time in the human brain. Our findings demonstrating cortical 710 beta oscillations sensitive to temporal duration align with previous reports of timing-related beta 711 power in time production domain (Grabot et al., 2019; Kononowicz & van Rijn, 2015), and 712 movement-related frontal midline delta-theta increases (Liang et al., 2018). On the other hand, 713 our findings regarding central-occipital alpha oscillations related to distance are consistent with 714 notions that human navigation is enriched with regarding to visual input (Ekstrom, 2015), with 715 occipital alpha oscillations particularly sensitive to visual-related changes (such as optic flow, 716 Cao & Händel, 2019). As proposed by Goyal et al. (2020), a theoretical link might therefore exist 717 between HPC movement-related theta and occipital alpha oscillations. For example, eye closure 718 induces alpha power increases both at occipital sites and in hippocampus (Geller et al., 2014). 719 Our current results would suggest differing roles in navigation for frontal midline theta (4-8 Hz) 720 and occipital alpha (8-12 Hz), which were both found relevant to movement (Liang et al., 2018), 721 and frontal midline theta and occipital alpha oscillations could possibly cooperate to support 722 task-dependent spatial or temporal processing. Therefore, a helpful next step would be to 723 determine how these signals coordinate between hippocampus and cortex in our task using 724 ECoG.

725 We note that when we compared the power spectra of the spatial and temporal 726 teleportation task, we did not find significant differences. Yet, we were able to classify whether a 727 trial was from the spatial or temporal task with an accuracy better than chance in a manner that 728 was generalizable across participants. This suggests the classifiers captured higher-order 729 differences (perhaps the underlying connectivity patterns) between the oscillatory coding of 730 space and time, other than the mean of power fluctuations. One future direction is to examine 731 the affinity of connectivity patterns for spatial coding and temporal coding, using a similar 732 behavioral task used in this study. We predict that the networks for spatiotemporal coding 733 should diverge, both measured using scalp EEG data, and using intracranial EEG data (as 734 proven by Watrous et al., 2013).

In addition to our findings that spatial distance and temporal duration involve differences in
 oscillatory codes, both for short vs. long teleportation durations and in their multivariate patterns,

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we also found a common role for alpha frequency modulation in supporting spatiotemporal 737 coding. Specifically, we found faster occipital alpha for smaller magnitudes of 738 739 durations/distances. What roles could endogenous alpha frequency modulation possibly play 740 here? One explanation is the processing-speed theory, whereby occipital alpha frequency 741 indexes the processing speed of incoming sensory information (Klimesch et al., 1996). We 742 speculate that the sensory processing speed differed between short and long duration trials 743 because of their different cognitive demands. To complete the temporal task, participants only 744 needed to track time passage in the teleporter up to 4s, and not beyond 4s, and therefore the 745 cognitive demands differed between 0-4s and 4-8s portions of the temporal task.

746 In contrast to the processing-speed account, another possibility however, relates to a 747 perceptual resolution account. For example, it could be that occipital alpha frequency is linked 748 to the perceptual resolution of duration timing. For example, individuals with 10 Hz resting 749 occipital alpha oscillations might discriminate two temporal durations with a minimum of 100ms 750 (1/10) differences, and those with 12 Hz resting alpha could discriminate two durations with 83.33ms minimal differences (1/12). This perceptual resolution account is also supported by 751 752 Samaha and Postle (2015) showing that occipital alpha frequency reflects the "refresh rate" of 753 visual perception and occipital alpha represents the perceptual unit of temporal processing 754 (Cecere et al., 2015). Future studies should investigate the potential causal links between 755 occipital alpha frequency and spatiotemporal processing, given recent findings that tACS-756 induced alpha frequency shifts led to shifts in subjective time experiences (Mioni et al., 2020) 757 and that clinical Alzheimer populations show irregularities in parietal alpha oscillations (Montez 758 et al., 2009).

759 Given that we found alpha frequency modulation and beta power fluctuations related to 760 both spatial and temporal judgments, our results also provide evidence for a common 761 mechanism for spatial and temporal coding involving magnitude estimation. Although distance-762 related beta power has rarely been studied in a scalp EEG setting, the timing-related beta 763 power we observed has been noted in predicting the accuracy and precision of time production 764 (Grabot et al., 2019; Kononowicz & van Rijn, 2015). Our findings suggest that beta oscillations 765 may reflect a common magnitude representation underlying both spatial and temporal processing, and that such distance and fine-scale temporal information could be widely 766 767 accessible in neocortical regions, including early sensory and motor cortices. Future studies can 768 bridge the gap of research between spatial and temporal processing, and further elaborate the 769 roles of beta oscillations in spatial coding vs. temporal coding, with a variety of tasks such as

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estimating and reproducing spatial distance with a path integration task (Harootonian et al.,

771 2020).

Another important finding from our study is the ability to decode fine-scale distance and 772 773 temporal information from cortical low-frequency power spectra. Interestingly, when attempting 774 to decode temporal information, we showed that the decoding error linearly increased as the 775 time bins progressed forward. These findings are closely aligned with the behavioral findings in 776 which humans show larger variability in time reproduction responses for longer intervals (Ivry & 777 Hazeltine, 1995; Rakitin et al., 1998). One intriguing possibility is that the cortical low-frequency 778 oscillations support a fine-scale representation of temporal intervals. Future studies can test this 779 possibility by linking the decodability of fine-scale time information and the accuracy/precision of 780 time reproduction in human participants.

781 Notably, our findings of decodable fine-scale temporal information are qualitatively similar 782 to the findings done with entorhinal temporal context cells (Bright et al., 2020). The tenet of a 783 unified math model of space and time (Howard et al., 2014) is that the neural representations are the Laplace transform of space and time, coded through the exponentially decayed firing 784 785 rates of neurons. However, the theory does not directly predict or rule out the involvement of 786 neural oscillations in coding space and time. Here we demonstrated that neural oscillations 787 could yield a similar time representation possibly with scale invariance, and we suggest that neural oscillations could be a synergistic component on top of single neuron firing rates for 788 789 spatiotemporal coding. Another question that should be clarified through future studies is 790 whether the neural representations of spatial distance also possess scale invariance like the 791 representations of time (i.e., reproducing longer distances are associated with greater variability 792 in responses.) Behavioral findings suggest path integration errors systematically scaled with 793 path lengths (Harootonian et al., 2020), which will predict linearly increases in decoding errors 794 as distances increase. Future studies should further test the links between oscillatory 795 representations of fine-scale space and time, and the behavioral phenomena of spatiotemporal 796 reproduction, using a reproduction paradigm, such as reproducing space and time in virtual 797 reality (E. M. Robinson & Wiener, 2020).

798

799 Limitations

800 It is worth considering some potential limitations with our paradigm which we 801 nonetheless believe do not undermine or challenge our findings. One concern could be that 802 because participants knew how far they would travel before entering the teleporter, distance 803 coding was therefore transient and completed *before* entering the teleporters, thus nullifying the

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804 existence of distance coding during the teleportation. We note, however, that maintenance of 805 distance information *during* the teleportation was still necessary for accurate performance in the 806 spatial teleportation task. When participants entered the teleporter, while they knew beforehand whether it was a short or long distance, they had to maintain this information during teleportation 807 808 to make the correct decision upon exiting the teleporter. Our interpretation of perceiving spatial 809 distance prior to decisions about movement is consistent with a rich literature in human spatial 810 navigation, suggesting that humans first estimate distance based on perceptual cues and then 811 attempt to maintain this in working memory as they actively navigate to different goals (Knapp & 812 Loomis, 2004; Philbeck et al., 1997; Philbeck & Loomis, 1997). Using a similar spatial distance 813 teleportation design, Vass et al. (2016) showed that the spatial distance teleportation task 814 resulted in different oscillatory profiles from those during the resting state (viewing a laptop 815 black screen outside the experimental context). We similarly found a clear difference between 816 teleportation and a resting baseline task. These findings suggest that the spatial teleportation 817 task triggered distance information processing absent in a resting state condition. 818 Another concern could be that movement-related noise from the navigation phase

permeated into the EEG data during the teleportation, thus confounding the findings we presented here. Note that the amount of noise, if any, should be identical between short and long trials, and between the spatial and temporal tasks, given that participants stood still after they entered the teleporter. Therefore, noise should not confound the findings regarding the contrasts of EEG responses between short and long trials, or between the spatial and temporal tasks.

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- 826 827

Conclusions

Our study addressed an important issue regarding whether spatial and temporal 828 829 processing share common or distinct mechanisms (Eichenbaum & Cohen, 2014; Ekstrom et al., 830 2011; Frassinetti et al., 2009; Gauthier et al., 2019, 2020; Watrous et al., 2013). Our findings 831 suggest that spatial and temporal judgments during navigation differ as a function of power changes within specific frequency bands: while spatial judgments resulted in changes in cortical 832 833 alpha and beta power, while different temporal durations were linked to changes in frontal 834 midline delta-theta, frontal and posterior alpha, and global beta power. Consistent with the idea 835 of separable representations for space and time, spatial and temporal discounting are behaviorally distinctive from each other (E. Robinson et al., 2019), estimating spatial distance 836 837 are subject to large errors (Zhao, 2018) while estimating suprasecond durations can be

THETA, ALPHA AND SPACE-TIME CODING 30 performed with high accuracy (Grabot et al., 2019), and spatial and temporal estimation errors 838 839 distort in opposing manners (Brunec et al., 2017). Previous reports have also hinted at a dissociation between space and time at the neural level although using different paradigms in 840 841 which temporal information, in particular, involved order and not duration (Ekstrom et al., 2011; 842 Watrous et al., 2013). More generally, evidence exists for and against the notion that space and time processing are of the same nature, and we also found evidence for alpha frequency 843 844 modulation as a common mechanism for spatial and temporal coding. Thus, one implication of 845 our study is that there are both distinct and common mechanisms related to how we process 846 spatial distance and temporal durations. 847 **Acknowledgements** 848 849 This research was supported by National Science Foundation (NSF BCS-1630296, A. D. 850 851 E.). We thank Stephanie Doner for the assistance in scalp EEG data collection, Eva Robinson

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