- 1 Title: Imaging decision-related neural cascades in the human brain
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#### 20 Abstract

21 Perceptual decisions depend on coordinated patterns of neural activity cascading across 22 the brain, running in time from stimulus to response and in space from primary sensory 23 regions to the frontal lobe. Measuring this cascade and how it flows through the brain is 24 key to developing an understanding of how our brains function. However observing, let 25 alone understanding, this cascade, particularly in humans, is challenging. Here, we report 26 a significant methodological advance allowing this observation in humans at 27 unprecedented spatiotemporal resolution. We use a novel encoding model to link 28 simultaneously measured electroencephalography (EEG) and functional magnetic 29 resonance imaging (fMRI) signals to infer the high-resolution spatiotemporal brain 30 dynamics taking place during rapid visual perceptual decision-making. After 31 demonstrating the methodology replicates past results, we show that it uncovers a 32 previously unobserved sequential reactivation of a substantial fraction of the pre-response 33 network whose magnitude correlates with decision confidence. Our results illustrate that 34 a temporally coordinated and spatially distributed neural cascade underlies perceptual 35 decision-making, with our methodology illuminating complex brain dynamics that would 36 otherwise be unobservable using conventional fMRI or EEG separately. We expect this 37 methodology to be useful in observing brain dynamics in a wide range of other mental 38 processes.

39

# 41 Introduction

42	The detailed spatiotemporal brain dynamics that underlie human decision-making are
43	difficult to measure. Invasive techniques with sufficient temporal or spatial resolution,
44	such as depth electrodes or cortical arrays used with epilepsy patients, are only feasible in
45	rare cases and, in addition, do not capture activity from the entire brain. In comparison,
46	non-invasive measures such as electroencephalography (EEG) and
47	magnetoencephalography (MEG) suffer from poor spatial resolution, and blood oxygen
48	level dependent functional MRI (BOLD fMRI) from poor temporal resolution and
49	indirect coupling to neural activity (e.g. fMRI) <sup>1</sup> . In spite of this, EEG, MEG, and fMRI
50	have been used individually to study perceptual decision-making in the human brain,
51	although, by themselves they provide a limited view of the underlying brain dynamics <sup>2</sup> .
52	Recently, methods enabling simultaneous acquisition of EEG and fMRI
53	(EEG/fMRI) have led to varied analytic approaches aimed at integrating the
54	electrophysiological and hemodynamic information contained in the joint measurements.
55	Such approaches offer the potential to provide a comprehensive picture of global brain
56	dynamics, and will likely offer new insights into how the brain makes rapid decisions <sup>3,4</sup> .
57	Some of the techniques that have been proposed for combining multi-modal brain signals
58	have separately analyzed the EEG and fMRI data and subsequently juxtaposed the
59	results <sup>5,6</sup> , while others attempt for a truly integrated approach in order to fully exploit the
60	joint information contained in the data sets <sup>7</sup> . In general, simultaneous EEG/fMRI and the
61	associated analysis techniques have been used to identify neuronal sources of EEG trial-
62	to-trial variability, linking them to cognitive processes such as attention $^{8}$ and inhibition $^{9}$ .

63	Many previous studies have used known EEG markers (P1, N2, N170, P300, $\alpha$ -
64	rhythm) or data driven approaches such as Independent Component Analysis (ICA) to
65	combine EEG with fMRI data <sup>4,8-16</sup> . One promising approach has been to use supervised
66	machine-learning techniques (e.g. classifiers) to find relevant projections of the EEG
67	data, where single-trial variability of the electrophysiological response along these
68	projections can be correlated in the fMRI space. Goldman, et al. <sup>17</sup> , Walz, et al. <sup>18</sup> and
69	Fouragnan, et al. <sup>19</sup> have demonstrated this technique on visual and auditory paradigms.
70	This methodology has been shown to localize cortical regions that modulate with the task
71	while preserving the temporal progression of task-relevant neural activity.
72	Here we combine a classification methodology with an encoding model that
73	relates the trial-to-trial variability in the EEG to what is observed in the simultaneously
74	acquired fMRI. Encoding models have become an important machine learning tool for
75	analysis of neuroimaging data, specifically fMRI <sup>20</sup> . In most cases encoding models have
76	been used to learn brain activity that encodes or represents features of a stimulus, such as
77	visual orientation energy in an image/video <sup>21-23</sup> , acoustic spectral power in sound/speech
78	<sup>24</sup> , or visual imagery during sleep <sup>25</sup> . In the method presented here, we employ an
79	encoding model to directly relate the simultaneously collected data from the two
80	neuroimaging modalities-instead of features derived from the stimulus, they are derived
81	from EEG component trial-to-trial variability. Specifically, we learn an encoding in the
82	spatially precise fMRI data from the temporally precise trial-to-trial variability of EEG
83	activity predictive of the level of stimulus evidence. This approach leverages the fact that
84	the level of stimulus evidence, as measured via EEG, persists across the trial <sup>26,27</sup> , and

that by discriminating this information in a time-localized way, one can temporally "tag"
specific cortical areas by their trial-to-trial variability.

87	Using our framework for learning the BOLD signal encoding of task-relevant and
88	temporally precise EEG component variability, we unravel the cascade of activity from
89	the representation of sensory input to decision formation, decision action, and decision
90	monitoring. A particularly novel finding is that after the activation of decision
91	monitoring regions (i.e. ACC), we see a reactivation of pre-response networks, where the
92	strength of this reactivation correlates with measures of decision confidence. This
93	specific reactivation, as well as the entire spatio-temporal cascade, is completely
94	unobservable using conventional fMRI-only or EEG-only methodologies.

95

#### 96 *Results*

97 In this study, we used a visual alternative forced choice (AFC) task where 98 subjects were shown brief presentations of pictures corrupted by noise and instructed to 99 rapidly discriminate between object categories. On any given trial, the level of noise, or 100 stimulus evidence, was varied randomly. The task itself, as well as similar visual decision-making tasks<sup>28</sup>, is believed to engage an extensive set of cortical areas in a 101 102 coordinated fashion, including regions that are responsible for sensory encoding, 103 evidence accumulation, decision formation, and response and decision monitoring. 104 However, the dynamic interplay of these regions has never been observed in humans. 105 Here we exploit previously reported findings regarding the sensitivity of the EEG and 106 fMRI signals to the level of stimulus evidence during a perceptual decision-making task. 107 Specifically, previous work has shown differential neural responses to high vs. low

108	stimulus evidence in trial averaged EEG event-related potentials (ERPs), where this
109	difference persists across the trial <sup>26,27</sup> . Similarly, fMRI studies have shown that for
110	perceptual decision making tasks a number of spatially-distributed cortical areas
111	significantly correlate with the level of stimulus evidence <sup>29,30</sup> . We leverage the fact that
112	the level of stimulus evidence is expressed temporally in the EEG and spatially in the
113	fMRI to "tag" voxels with a time. Specifically, using a classification methodology (i.e.
114	discriminative components) we identify temporally precise expressions of the level of
115	stimulus evidence that then can be spatially localized through an encoding model of the
116	fMRI BOLD data.
117	We collected simultaneous EEG/fMRI data from 21 subjects as they performed a
118	3-AFC task discriminating between faces, cars, and houses (Fig. 1A). Subjects were
119	instructed to discriminate the object class after briefly viewing an image corrupted by
120	varying levels of noise (Fig. 1B) and respond by pressing one of three buttons. Overall,
121	subjects responded with accuracies of $94 \pm 5\%$ and $58 \pm 12\%$ and with response times of
122	$634 \pm 82$ ms and $770 \pm 99$ ms for high and low stimulus evidence trials, respectively (Fig.
123	1 C, D). Subject accuracies and response times across stimulus types (faces, cars, houses)
124	for low stimulus evidence trials were similar; however, for high stimulus-evidence trials
125	subject accuracies were higher and response times were shorter for faces than for cars or
126	houses (See Supplemental Information Fig. S1).
127	

128 GLM based analysis of BOLD fMRI shows superposition of cortical areas correlated
129 with stimulus evidence

130	A traditional general linear model (GLM) analysis of the fMRI (see Methods)
131	revealed differences in BOLD activation between the two stimulus evidence conditions
132	(Fig. 1F, SI Table 1). Brain regions showing greater BOLD activation to high vs. low
133	stimulus evidence trials included areas associated with early visual perception and the
134	default mode network <sup>26</sup> , such as fusiform gyrus, parahippocampal gyrus, lateral occipital
135	cortex, superior frontal gyrus, and posterior cingulate cortex. Regions with greater BOLD
136	activation to low vs. high stimulus evidence trials included areas in the executive control
137	and difficulty networks' such as dorsal lateral prefrontal cortex, anterior cingulate cortex,
138	intraparietal sulcus, and insula. Overall, these GLM results for the BOLD data
139	reproduced previous results in the literature where similar stimuli and paradigms were
140	used <sup>29</sup> (Fig. S2A).
141	
142	Extracting temporally localized EEG signatures of stimulus evidence variability
143	The traditional fMRI results showed multiple brain regions correlated with the
144	difficulty, or stimulus evidence, of the trial; however, this traditional approach does not

145 enable one to infer the relative timing of these fMRI activations. To infer timing at a scale of tens of milliseconds, we used linear classification<sup>31,32</sup> of the EEG to extract trial-146 147 to-trial variability related to stimulus evidence at specified post-stimulus time points. 148 The basic idea is illustrated in Figure 2, where hypothetical neural activity is 149 shown for two different regions that are constituents of the perceptual decision-making 150 network. Averaging over trials would clearly reveal a difference in the mean neural 151 activity between high and low stimulus evidence. However, the two regions contribute 152 differentially to the network, with one region encoding the stimulus evidence (Region 1)

153 and the other integrating it over time (Region 2); both are sensitive to the level of 154 stimulus evidence, though varyingly so at different times in the trial. By taking 155 advantage of this sensitivity to the stimulus evidence, we can learn EEG discriminant 156 components, i.e. spatial filters, that best classify trials at different time windows given the 157 neural data. We used the trial-to-trial variability along these component directions as 158 features to uniquely tag fMRI voxels with the specific time window of the component. 159 This tagging is done by building an encoding model of the features, given the BOLD 160 signal, details of which are described in the following section. 161 We constructed EEG components by learning linear classifiers at 25ms steps, 162 starting from stimulus onset to 50ms past the average low stimulus evidence response 163 time. We chose a time step of 25ms due to an empirical analysis showing a half width of 50ms in the temporal autocorrelation of the EEG data, though in principle this 164 165 methodology allows for temporal resolution up to the EEG sampling rate. Each classifier 166 was associated with a set of discriminant values, which can be represented as a vector  $y_{\tau}$ ; 167 each element of the vector is the distance of a given trial to the discrimination boundary 168 for the classifier at time step  $\tau$  (Fig. 2). This distance can be interpreted as a measure of the EEG classifier's estimate of the level of stimulus evidence for that trial<sup>17,18,31-34</sup>. 169 170 Results of the EEG analysis show discriminating information for stimulus 171 evidence spanning the trial (see Fig. 4A), beginning roughly 175ms post-stimulus to past 172 the average response times. A dip occurs around 300ms, indicating stimulus evidence is 173 less discriminative at this time and serves to demarcate early and late cognitive processes. 174 The early process corresponded to the time of the D220 ERP component, which has been 175 shown to modulate with the degree of task difficulty, whether via stimulus noise or task

176	demands <sup>35</sup> . The later and more prolonged component is likely related to more complex
177	cognitive and motor preparatory processes that differ between high and low stimulus
178	evidence trials. Importantly, although the early and late EEG components were both
179	discriminative, we found their trial-to-trial variability to be uncorrelated (Figs. 4B and
180	S3E), indicating that while the discriminating information (level of stimulus evidence)
181	persists across the trial, it couples differently to processes across time.
182	
183	An encoding model links fMRI activations with temporally distinct EEG trial-to-trial
184	variability
185	After extracting the trial-to-trial variability from the EEG discriminant
186	components, feature vectors $y_\tau$ are collected across time steps, $\tau,$ along with a response
187	time vector to construct a matrix $Y$ . This matrix is the temporally precise representation
188	of the trial-to-trial EEG variability that reflects high vs. low stimulus evidence. An
189	encoding model is then fit, namely a model in which weights are estimated for each time-
190	localized EEG window, to predict the trial-to-trial variability of the BOLD response for
191	each fMRI voxel. Figure 3 shows a schematic of the encoding model framework we used
192	and compares it to a traditional encoding model constructed by using features derived
193	directly from the stimulus. Rather than constructing a map that directly relates each voxel
194	to a type of stimulus feature, such as whether it encodes edges, motion or some semantic
195	concept such as "animal" <sup>21-23,36-38</sup> , our model is used to construct maps that label voxels
196	by the time window of the variability they encode $-i.e.$ it "tags" each voxel with a
197	"time", or set of times, when it encodes the variability in the given EEG discriminant
198	component(s).

199 It is important to note that this approach does not attempt to improve source 200 localization typically done for EEG/MEG studies. Our approach instead provides the 201 temporal resolution of EEG (ms) and the spatial resolution of fMRI (mm) without the 202 need to solve the ill-posed inverse solution and make the many associated assumptions 203 required for reliable source-localization results<sup>39</sup>.

204 An example of the quality of the encoding model is shown in Fig. 4C (see also 205 Fig. S2B) where significant voxels from the encoding model are shown in yellow. Fig. 206 4D shows the trial-to-trial variability of BOLD signal at a specific voxel, comparing it to 207 the variability predicted by the encoding model. Additional validity of the encoding 208 model and single subject results are presented in the Supplemental Information (Fig. 209 S4A/B). The encoding model was also evaluated as a decoding model (see Methods) with 210 the BOLD activity used to predict the trial-to-trial variability in the EEG for unseen 211 data-data on which the encoding model was not trained. Fig. 4E shows these results, 212 expressed as the correlation between the measured and predicted EEG trial-to-trial 213 variability across the 800ms epoch. The shape of the curve is highly consistent with that 214 observed for the EEG data itself (comparing Fig. 4A and Fig. 4E) (additional analysis of 215 the fidelity of the model is provided in the SI, Fig. S3). 216 Given the encoding model, we unwrap the BOLD activity across time by 217 identifying weights that are consistent across subjects in space and time (see Methods). 218 Fig. 5 shows these results for a group level analysis. We observe a progression of activity 219 (see Movie S1), at 25ms resolution, which proceeds simultaneously down the dorsal and

- ventral streams of visual processing for the first 250ms. After that the cascade becomes
- 221 more complex with activation in the IPS at 425ms and 750ms (see Fig. 6A), reactivation

222	of the SPL at 675ms and activation of ACC at 600ms along with other regions found in
223	the traditional fMRI results. (see Fig. S5, Tables S2 and an additional analysis using
224	dynamic causal modeling <sup>40</sup> ). The reactivation pattern is particularly significant since it
225	would not be observable via a traditional fMRI general linear model (GLM) analysis,
226	which integrates over time and thus superimposes these activities. For example, the
227	changing sign of the middle temporal gyrus (MT) encoding weights in Fig. 6A
228	manifested as no activity in the MT for the traditional fMRI GLM analysis-the change
229	in sign canceled the effective correlation in the GLM (see Fig. 1F and Fig. S1). The areas
230	of activation we find are consistent with previous reports in the literature for human
231	subjects <sup>29,30</sup> ; however, here we are able to link activations across time in a way that was
232	previously only possible with invasive techniques.

233

## 234 Cortical reactivation correlates with decision confidence

235 Further analysis of the spatiotemporal dynamics (see Fig. 6B), shows that the 236 reactivation pattern in the network occurs after decision-monitoring areas become 237 engaged (i.e. after ACC). Spontaneous reactivation, or "replay", of neural activity in the human brain has been observed and believed to be important for memory consolidation<sup>41</sup> 238 239 and more recently has been hypothesized to play a role in perceptual decision-making by enabling the formation of decision confidence<sup>42</sup>. To test the hypothesis that the 240 241 reactivation activity we see is in fact related to decision confidence, we used a hierarchical drift diffusion model (DDM)<sup>43,44</sup> to fit the behavioral data for high and low 242 243 stimulus evidence conditions (see Methods). Specifically, our model enables us to define a proxy for decision confidence based on the DDM fits to the behavior<sup>45,46</sup>. Correlating 244

245	the reactivation level to this confidence proxy shows a strong and significant monotonic
246	relationship between confidence and the level of reactivation (high stimulus evidence-
247	slope=0.037±0.008, t=4.657, p=3.2x10 <sup>-6</sup> ; low stimulus evidence-slope=0.062±0.008,
248	t=7.754, p= $8.88 \times 10^{-15}$ ), with low stimulus evidence trials reactivated more strongly than
249	high stimulus evidence trials (difference in slopes=-0.025±0.011, t=2.189, p=0.029)(see
250	Fig. 7 and Fig. S7). Additionally, reactivation amplitude correlates with behavioral
251	accuracy (Fig. S8) (high stimulus evidence, slope=0.0115±0.0047, t=2.41, p=0.016; low
252	stimulus evidence, slope=0.0104±0.0047, t=2.19, p=0.028). Recursive feature elimination
253	showed that the IPS/SPL and dorsal lateral prefrontal cortex (DLPFC) clusters
254	contributed the most to reactivation/confidence proxy correlation (Fig. 7C).
255	
256	
257	Discussion
258	We have shown that linking simultaneously acquired EEG and fMRI using a novel
250	anading model anghles imaging of high resolution anotistem paral dynamics that

encoding model enables imaging of high-resolution spatiotemporal dynamics that

260 underlie rapid perceptual decision-making — decisions made in less than a second. This

261 method, which resolves whole-brain activity with EEG-like temporal resolution, was

shown to uncover reactivation processes that would otherwise be masked by the temporal

averaging and slow dynamics of traditional fMRI. More broadly, our results

demonstrated a general non-invasive data-driven methodology for measuring high

spatiotemporal latent neural processes underlying human behavior.

266 This approach temporally "tags" the BOLD fMRI data by encoding the trial-to-267 trial variability of the temporally precise task relevant components in simultaneously 268 acquired EEG. In effect, the EEG discrimination indexes the activity of interest at high 269 temporal resolution, defining a feature space, and the trial-to-trial variability of these 270 discriminant components becomes the specific feature values used in the encoding model. 271 For the case presented here, this variability was used to tease apart the cascade of activity 272 modulated by stimulus evidence across the trial, and this allowed us to observe, as never 273 seen before, the spatiotemporal brain dynamics underlying a perceptual decision. 274 Previous studies have sought to generalize the timing diagram of a perceptual decision through multi-unit recordings in non-human primates<sup>47,48</sup> or more broadly in 275 humans<sup>29,30</sup> using fMRI. Our results confirmed the general temporal ordering of 276 277 activations found previously (early visual processing, decision formation, decision 278 monitoring). However, there was a possibility the temporal order we observed using our 279 technique was an artifact of our methodology. To assess this possibility, we performed 280 additional analyses using dynamic causal modeling (DCM) to further validate the 281 temporal activation sequence (see Fig. S6) and show, using a different set of assumptions 282 and method, that the temporal sequence we observe is highly likely under a set of 283 alternative sequences. We found that the most likely model is the one consistent with the 284 time course inferred from our encoding model. The DCM results provide additional 285 evidence that the temporal profile uncovered by the encoding model is a likely temporal 286 decomposition of the superimposed fMRI activations. 287 The approach we present requires that EEG and BOLD data be collected 288 simultaneously and not in separate sessions in order to exploit the correlations in trial-to-289 trial variability to "tag" the BOLD data. To show the importance of collecting the data

simultaneously, we ran a control analysis that randomly permuted the trials within their

291 stimulus evidence class, thus effectively simulating an EEG and BOLD dataset collected 292 separately. By destroying the link between the EEG and BOLD trials, the encoding 293 model failed to find any consistent activation (Fig. S11/12), indicating the necessity of 294 simultaneous acquisition. 295 Alternative techniques for fusing simultaneous EEG-fMRI typically do not 296 exploit EEG across the trial and instead only analyze specific ERP components or time windows of interest <sup>4,8,10,12-19,49,50</sup>. Results from these techniques identify regions that 297 298 modulate with the specific components, but yield limited information about the timing of 299 other task-relevant regions seen in traditional fMRI contrasts. The methodology developed here extends the work of Goldman, et al.<sup>17</sup> and Walz, et al.<sup>18</sup> by combining 300 301 their EEG data reduction techniques with techniques developed for encoding stimulus features onto BOLD data<sup>20-23,36,38</sup>, ultimately providing a framework for labeling voxels 302 303 in task-relevant fMRI contrasts with their timing information (Fig. S2C/E/F). 304 Clearly, other EEG components that are task-related can be isolated and could 305 potentially be used to "tag" BOLD data. The sliding window linear classification used 306 here acts to reduce the EEG data along a dimension that categorizes stimulus evidence; 307 however, this could be replaced by any other data reduction technique, such as 308 temporally windowed ICA or PCA. Variability along these component directions could 309 then be used in the encoding model to link with the simultaneously collected BOLD data. 310 The choice of data reduction technique (i.e. feature space) would be highly dependent on 311 the nature of the inferences. 312 Our methodology enabled us to observe reactivation of the pre-response network,

313 spatiotemporal dynamics that would be masked using traditional fMRI analysis.

314	Interestingly, the reactivation terminated in a network that included the MFG, SPL, and
315	IPS, similar areas previously reported to be reactivated in metacognitive judgments of
316	confidence in perceptual decisions <sup>42,51,52</sup> . In addition, these areas contributed the most to
317	the correlation to confidence proxy (Fig. 7C). Gherman and Philiastides <sup>53</sup> observed this
318	network using a multivariate single-trial EEG approach, coupled with a distributed source
319	reconstruction technique. Fleming, et al. <sup>42</sup> and Heereman, et al. <sup>54</sup> used BOLD fMRI to
320	show that areas in this network negatively correlate with subjective certainty ratings.
321	Unique to our findings, we saw this reactivation on a single-trial basis after engagement
322	of the ACC, which has been shown to be involved in decision monitoring <sup>53,55</sup> , and also
323	observed the dynamic sequence leading up to this network reactivation. Our results
324	showed that reactivation/replay occurred on a trial-to-trial basis after a decision, was
325	stronger for difficult decisions, and correlated with decision confidence.
326	A potential confound in our analysis is that the timing of the reactivation overlaps
327	with some of the response times. To check if the reactivation was pre or post response,
328	we implemented a response-locked encoding model analysis (Fig. S9). The response-
329	locked results showed significant activation pre-response that overlaps with the
330	reactivation network from the stimulus locked analysis. In addition, trial-to-trial
331	reactivation taken from pre-response clusters correlates with confidence proxy similarly
332	to the stimulus locked results (Fig. S10). This provides further evidence that the
333	reactivation is occurring pre-response.
334	The encoding model we developed was able to decompose traditional fMRI
335	activation maps into their temporal order with significant voxel overlap between the

336 encoding model results and traditional results. The encoding model was also able to show

337 regions that were activated at multiple time points throughout the decision, indicating 338 temporal dynamics that were hidden previously. The regions of activation we found are 339 consistent with earlier findings; however, the work here provided the precise temporal 340 decomposition of these previously reported, temporally superimposed regions of 341 activation. In general, we have shown that simultaneously acquired EEG/fMRI data 342 enables a novel non-invasive approach to visualize high resolution spatial and temporal 343 processing in the human brain with the potential for providing a more comprehensive 344 understanding of the neural basis of complex behaviors. 345

- 346 *Methods*
- 347 Subjects

348 21 subjects (12 male, 9 female; age range 20-35 years) participated in the study. The

349 Columbia University Institutional Review Board (IRB) approved all experiments and

- 350 informed consent was obtained before the start of each experiment. All subjects had
- 351 normal or corrected-to-normal vision.

352 Stimuli

We used a set of 30 face (from the Max Planck Institute face database), 30 car, and 30

house (obtained from the web) gray scale images (image size 512x512 pixels, 8

bits/pixel). They were all equated for spatial frequency, luminance, and contrast. The

356 stimulus evidence (high or low) of the task was modulated by systematically modifying

the salience of the image via randomization of image phase (35% (low) and 50% (high)

358 coherence)<sup>56</sup>.

359 Experimental task

360	The stimuli were used in an event-related three-alternative forced choice (3-AFC) visual
361	discrimination task. On each trial, an image either a face, car, or house was presented
362	and subjects were instructed to respond with the category of the image by pressing one of
363	three buttons on an MR compatible button controller. Stimuli were presented to subjects
364	using E-Prime software (Psychology Software Tools) and a VisuaStim Digital System
365	(Resonance Technology) with 600x800 goggle display. Over four runs, a total of 720
366	trials were acquired (240 of each category with 120 high coherence trials) with a random
367	inter-trial interval (ITI) sampled uniformly between 2-2.5s. Each run lasted for 560
368	seconds.
369	fMRI acquisition
370	Blood-oxygenation-level-dependent (BOLD) T2*-weighted functional images were
371	acquired on a 3T Philips Achieva scanner using a gradient-echo echo-planar imaging
372	(EPI) pulse sequence with the following parameters: Repetition time (TR) 2000ms, echo
373	time (TE) 25ms, flip angle 90°, slice thickness 3mm, interslice gap 1mm, in-plane
374	resolution 3x3mm, 27 slices per volume, 280 volumes. For all of the participants, we also
375	acquired a standard T1-weighted structural MRI scan (SPGR, resolution 1x1x1mm).
376	EEG acquisition
377	We simultaneously and continuously recorded EEG using a custom-built MR-compatible
378	EEG system <sup>57,58</sup> , with differential amplifiers and bipolar EEG montage. The caps were
379	configured with 36 Ag/AgCl electrodes including left and right mastoids, arranged as 43
380	bipolar pairs. Bipolar pair leads were twisted to minimize inductive pickup from the
381	magnetic gradient pulses and subject head motion in the main magnetic field. This
382	oversampling of electrodes ensured data from a complete set of electrodes even in

383 instances when discarding noisy channels was necessary. To enable removal of gradient

artifacts in our offline preprocessing, we synchronized the EEG with the scanner clock by

385 sending a transistor– transistor logic pulse at the start of each image volume. All

electrode impedances were kept below 20 k $\Omega$ , which included 10 k $\Omega$  resistors built into

- 387 each electrode for subject safety.
- 388 Functional image pre-processing.
- 389 Image preprocessing was performed with FSL (www.fmrib.ox.ac.uk/fsl/). Functional
- images were spatially realigned to the middle image in the times series (motion-

391 correction), corrected for slice time acquisition, spatially smoothed with a 6mm FWHM

392 Gaussian kernel, and high pass filtered (100s). The structural images were segmented

393 (into grey matter, white matter and cerebro-spinal fluid), bias corrected and spatially

394 normalized to the MNI template using 'FAST' <sup>59</sup>. Functional images were registered into

395 MNI space using boundary based registration  $(BBR)^{60}$ .

396

#### 397 *EEG data preprocessing.*

398 We performed standard EEG preprocessing offline using MATLAB (MathWorks) with 399 the following digital Butterworth filters: 0.5 Hz high pass to remove direct current drift, 400 60 and 120 Hz notches to remove electrical line noise and its first harmonic, and 100 Hz 401 low pass to remove high-frequency artifacts not associated with neurophysiological 402 processes. These filters were applied together in the form of a zero-phase finite impulse 403 response filter to avoid distortions caused by phase delays. We extracted stimulus-locked 404 1500 ms epochs (-500:1000) and subtracted the mean baseline - -200 ms to stimulus 405 onset – from the rest of the epoch. Through visual inspection, we discarded trials

406 containing motion and/or blink artifacts, evidenced by sudden high-amplitude

407 deflections.

408 Sliding window logistic regression.

409 We used linear discrimination to associate each trial with the level of stimulus evidence

410 represented in the EEG. We considered high stimulus and low stimulus evidence trials

411 irrespective of behavioral accuracy. Regularized logistic regression was used as a

412 classifier to find an optimal projection for discriminating between high and low stimulus

413 evidence trials over a specific temporal window. A sweep of the regularization

414 parameters was implemented using FaSTGLZ<sup>61</sup>. This approach has been previously

415 applied to identify neural components underlying rapid perceptual decision-making

416 17,18,31,33,34,45,50,62

417 Specifically, we defined 50ms duration training windows centered at time,  $\tau$ ,

418ranging from stimulus onset to 800ms following the stimulus in 25ms steps. We used419logistic regression to estimate a spatial weighting, on N EEG channels, vector ( $w_{\tau}$  which420is N x 1) that maximally discriminated between EEG sensor array signals E for each class

421 (e.g., high vs. low stimulus evidence trials):

 $y_{\tau} = w_{\tau}^T E_{\tau} \tag{1}$ 

In eqn. 1,  $E_{\tau}$  is an N x p vector (N sensors per time window  $\tau$  by p trials). For our experiments, the center of the window ( $\tau$ ) was varied across the trial in 25ms time-steps. We quantified the performance of the linear discriminator by the area under the receiver operator characteristic (ROC) curve, referred to here as AUC, using a leave-one-out procedure. We used the ROC AUC metric to characterize the discrimination performance as a function of sliding our training window (i.e., varying  $\tau$ ). For each subject, this

429	produced a matrix Y where the rows corresponded to trials and the columns to training
430	windows, i.e. Y is the combination of the calculated $y_{\tau}$ for each time window.
431	Traditional fMRI analysis.
432	We first ran a traditional general linear model (GLM) fMRI analysis in FSL, using
433	event-related (high and low stimulus evidence) and response time (RT) variability
434	regressors. The event-related regressors comprised boxcar functions with unit amplitude
435	and onset and offset matching that of the stimuli. RT variability was modeled using the z-
436	scored RT as the amplitude of the boxcars with onset and offset matching that of the
437	stimulus, and these were orthogonalized to the event-related regressors.
438	Orthogonalization was implemented using the Gram-Schmidt procedure <sup>63</sup> to decorrelate
439	the RT regressor from all other event-related regressors. All regressors were convolved
440	with the canonical hemodynamic response function (HRF), and temporal derivatives
441	were included as confounds of no interest. An event-related high versus low stimulus
442	evidence contrast was also constructed. A fixed-effects model was used to model
443	activations across runs, and a mixed-effects approach was used to compute the contrasts
444	across subjects. Activated regions that passed a family-wise error (FWE) <sup>64</sup> corrected
445	cluster threshold of $p < 0.01$ at a z-score threshold of 2.57 were considered significant.
446	fMRI deconvolution.
447	Associating fMRI data to each trial is challenging for two main reasons: (a) the temporal
448	dynamics of the hemodynamic response function (HRF) evolve over a longer time-scale
449	than the mean ITI of the event-related design, resulting in overlapping responses between
450	adjacent trials; and (b) the ITI was random for each trial so that the fMRI data was not
451	acquired at a common lag relative to stimulus onset. To overcome these issues, we

452	employed the `least squares - separate' (LS-S) deconvolution <sup>65</sup> method to estimate the
453	voxel activations for each trial. For every trial, the time series of each voxel was
454	regressed against a "signal" regressor and a "noise" regressor. The "signal" regressor was
455	the modeled HRF response due to that trial (a delta function centered at stimulus onset
456	convolved with a canonical HRF), while the "noise" regressor was the modeled HRF
457	response due to all other trials (superimposed linearly). The resulting regression
458	coefficients of the "signal" regressor represented the estimated voxel activations due to
459	that trial. These voxel activations were then organized into a single brain volume per trial.
460	We extracted 58697 voxels from a common gray matter group mask at 3 mm <sup>3</sup> spatial
461	resolution that excluded white matter and CSF and assembled the resulting voxel
462	activations into rows of the data matrix F.
463	Single subject encoding model.
464	All encoding model analyses were performed in MATLAB. To relate the EEG data with
465	the fMRI, we devised a subject-wise spatio-temporal decomposition using singular value
466	decomposition (SVD). Let F be an m x p matrix denoting m-voxels and p-trials that is the
467	deconvolved high and low stimulus evidence fMRI data for each trial. Let Y be the r x p
468	matrix denoting r-windows (33 $\text{EEG}_{\tau}$ windows and response time (RT)) and p-trials. For
469	each trial, the first row of Y is the response times while subsequent rows are the y values
470	at each window time. Let W be an m x r matrix that is the weights on Y that solve for F.
471	$F = WY \tag{2}$
472	Normally, if we solve for W using the least squares approach, we get:

473 
$$W=(FY^{T})(YY^{T})^{-1}$$
 (3)

474 However, each time point might be highly correlated with its neighbors, which reduces the stability of the least-squares regression. We can use SVD to reduce the feature space 475 476 and improve our estimation of W (the weights on each window). Then for a leave-one-477 out cross validation, we hold out a single trial from the EEG Y matrix and the 478 corresponding volume from the fMRI data F and train on the remaining trials. We 479 repeated this for all trials.  $\mathbf{Y}^{\text{Train}} = \mathbf{U} \boldsymbol{\Sigma} \mathbf{V}^{\text{T}}$ 480 (4) 481 Where U is an r x r orthonormal matrix,  $\Sigma$  is a r x p diagonal matrix and V is a p x p orthonormal matrix. After SVD on Y<sup>Train</sup>, we reduced the feature dimensions on Y<sup>Train</sup> to 482 483 retain 75% of the variance by only keeping v components. To do this, we selected the first v rows of  $\Sigma$  and zeroed the other rows. We now have  $\tilde{\Sigma}$  as our reduced spaced 484 485 matrix. If we now recalculate our least squares solution where we have replaced Y by its reduced form  $U\tilde{\Sigma}V^{T}$  in equation 3: 486  $\hat{W} = (F^{Train}V\tilde{\Sigma}^T)(\Sigma\Sigma^T)^{-1}U^T$ 487 (5) 488 So for each leave one out fold, we first calculated the SVD of the training set. We then calculated the number of components to keep and then solve for  $\hat{W}$ , the weight estimate 489 per fold. To test, we then applied the weights to the left-out test data Y<sup>Test</sup> to estimate the 490 encoded fMRI data  $\hat{F}$  for the encoding part: 491  $\hat{F} = \hat{W}Y^{Test}$ 492 (6)

493 While for the decoding model using the left out test data  $F^{Test}$ :

494 
$$\hat{Y} = \hat{W}^T F^{Test} (\hat{W}^T \hat{W})^+$$
(7)

495 Here,  $\hat{W}^T \hat{W}$  is not invertible, and so we used the pseudo-inverse.

496	At this point, we have $\hat{F}$ , a m x p matrix with m voxels by p trials. For each voxel
497	j, we calculated the correlation of $\hat{F}_j$ with F <sub>j</sub> , resulting in the matrices R <sup>fMRI</sup> (Pearson
498	Correlation Map) and $P^{fMRI}$ (p-value map of the Pearson Correlation) that are m x 1. The
499	$P^{fMRI}$ was then converted to a z-score map. We constructed the m x r weight matrix W by
500	taking the average of all the trained $\hat{W}$ matrices. To test which time windows were
501	significant, we also calculated, $R_{\tau}^{EEG}$ , the correlation between $\hat{Y}_{\tau}$ and $Y_{\tau}$ .
502	Group level spatio-temporal analysis.
503	For group level statistics, we first analyzed the $R_{\tau}^{EEG}$ vectors across all subjects. The $R_{\tau}^{EEG}$
504	vectors were converted into their p-values, and for each time window ( $\tau$ ), used to
505	compute combined Stouffer p-values <sup>66</sup> . These group level results were then false
506	discovery rate corrected (FDR) for multiple comparisons <sup>67</sup> . To identify group level
507	voxels where our model predictions were significant, each subject's p-value maps for the
508	leave-one-out correlation were converted into their respective z-values, and voxel-wise
509	significance was calculated using threshold-free cluster enhancement (TFCE) using a
510	non-parametric randomization procedure implemented in FSL <sup>68</sup> . Voxels were considered
511	significant if they passed a conservative false discovery rate threshold of $p < 0.01$ .
512	These significant voxels were then used as a mask to temporally localize
513	activations by computing the voxels that were consistent in their direction ( positive (high
514	stimulus evidence) or negative (low stimulus evidence) ) and timing ( $\tau$ window). To this
515	end, we implemented a spatio-temporal TFCE (stTFCE) in both space (neighboring
516	voxels) and time (neighboring time windows - response time window not included) and
517	computed significance through a randomization procedure. 33000 permutations (1000
518	permutations per window) were run by randomly altering the sign of each subject and the

519	temporal ordering of the windows, as we were testing whether the weights were
520	consistent in sign, voxel space, and temporal window. P-values were calculated by
521	comparing the true stTFCE value with the distribution of permuted values. Again, voxels
522	were considered significant if they passed FDR correction at p<0.05 (high stimulus
523	evidence: FDR-Corrected p<0.0019, low stimulus evidence: FDR-Corrected p<0.00036).
524	Note, that now our number of multiple comparisons was the number of voxels in the
525	FDR-mask (20256) times the number of time windows (33). We analyzed the response
526	time separately with a standard TFCE randomization procedure implemented in FSL
527	(Fig. S2D).
528	Dynamic causal modeling.
529	To validate the encoding model timing, we implemented single-state linear
530	dynamic causal modeling (DCM) using DCM10 in SPM8 <sup>69</sup> , and applied this to the
531	BOLD data to test the hypothesis that the temporal sequence of BOLD activations we
532	found in our EEG-fMRI encoding method was most likely, relative to other possible
533	sequences of these same activations, given only the BOLD data. We used the results of
534	the encoding model to select seven regions of interest that spanned the entire trial. For the
535	first region (labeled 175 in our figures), we computed the union of activations during the
536	175ms and 200ms windows. Activations of the 225ms (225) and 250ms combined with
537	275ms (250) windows become the second and third regions. We computed the union of
538	activations during the 325ms and 350ms windows to create the fourth (325). For the fifth
539	region (400), we computed the union of the activations during the 400ms-450ms
540	windows. For the sixth region (650), we computed the union of the activations during the
541	650ms and 675ms windows. Finally, the union of the activations during the 725-800ms

542 windows was computed to create the seventh region (725). We removed any overlapping 543 voxels between any of the regions and then extracted time series from individual 544 subjects' preprocessed functional data in MNI space by estimation of the first principal 545 component within each region. 546 We constructed 11 models (Figure S6) to investigate the directed connectivity of 547 these regions and validate the temporal ordering found by the encoding model. Each 548 model was feed-forward with first node in each model as the input region. The first 549 model was the temporal ordering of the regions inferred from our EEG-fMRI encoding 550 model analysis. For five of the models, we randomized the temporal ordering of the early 551 regions (175, 225, 250) and the late regions (325, 400, 650, 725) separately. For the other 552 five models, we fully randomized the temporal ordering of all the regions. 553 We used fixed-effects Bayesian model selection (BMS) to compare these 11

models both on a single-subject level and at the group level. BMS balances model fit and complexity, thereby selecting the most generalizable model. It estimates the relative model evidence and provides a distribution of posterior probabilities for all of the models considered. We also compared families of similar models<sup>70</sup>; the model space was divided into two families (early/late or fully randomized).

559

560 Drift Diffusion Model (DDM) and Confidence Proxy.

The DDM models decision-making in two-choice tasks. Here, we treated the decision (correct vs. incorrect) as our two choices. A drift-process accumulates evidence over time until it crosses one of two boundaries (upper or lower) and initiates the corresponding response<sup>68</sup>. The speed with which the accumulation process approaches one of the two

boundaries (a) is called drift-rate (v) and represents the relative evidence for or against a particular response. Recently, Philiastides, et al. <sup>45</sup> showed that for conditions in which the boundary (a) does not change, a proxy for decision confidence for each trial (i) can be computed by  $1/\sqrt{RT_i - T_{non}}$ .

569 We used Hierarchical Bayesian estimation of the Drift-Diffusion Model in Python (HDDM) to calculate the drift rate (v), decision boundary (a) and non-decision time  $T_{non}$ 570 for each subject <sup>43</sup>. Specifically, we modeled high and low stimulus evidence response 571 572 time data separately. This was to ensure our confidence proxies were consistent within 573 trial types. We included the response time and whether the subject got the trial correct. 574 HDDM obtains a sequence of samples (i.e., a Markov chain Monte Carlo; MCMC) from 575 the posterior of each parameter in the DDM. In our model, we generated 5000 samples 576 from the posteriors, the first 1000 (burn-in) samples were discarded, and the remaining 577 samples were thinned by 5%.

After modeling the DDM process, each trial's (i) confidence proxy (CP) for each subject (j) was computed by  $CP_{i,j} = 1/\sqrt{RT_i - T_{non,j}}$  and then z-scored across trials where  $T_{non,j}$  was varied for high or low stimulus evidence trials, separately. Therefore, CP was a measure of relative trial confidence within difficulty levels.

582

## 583 Confidence Proxy and Decision Replay.

584 Trial to trial reactivation amplitude was defined as  $Y_{j,i}^{R} = W_{j,PostACC}^{T}F_{j,i}$  for each 585 subject (j) and trial (i), where  $W_{postACC}$  is the weight matrix from the encoding model 586 thresholded by voxels that were significant in the group results from the 600-800ms

587	windows. The mean of the $Y_{j,i}^{R}$ across time becomes a measure of "decision replay"
588	strength for that trial (more negative y's indicate more replay activation, more positive y's
589	indicate less replay activation). $Y_{j,i}^R$ was quintiled for high and low stimulus evidence
590	and the average confidence proxy was calculated within each quintile (Fig. 7). A linear
591	mixed effects model <sup>71</sup> was used to test if the slope of confidences across quintile
592	grouping, $Y_{j,i}^{R}$ , were significantly different from 0 while including stimulus evidence as a
593	condition. Separate similar analyses with non-replay windows (175-250ms) and testing
594	for behavioral accuracy were also performed (Fig. S7-8). To test the contribution of each
595	cluster to the correlation with confidence, we implemented recursive feature elimination,
596	where our features were clusters of significant voxels (> 48 voxels) during the 600-
597	800ms time window. This procedure removed clusters from the 'replay' network before
598	calculating trial-to-trial reactivation. We then calculated the percent change in slope
599	(reactivation x confidence proxy) when the cluster was removed compared to the total
600	network. This procedure ranks cluster importance by sorting which clusters, when
601	removed, had the strongest negative effect on slope height.
602	

- -

# 603 Author Contributions

#### 604 Conceptualization, J.M. and P.S.; Methodology, J.M., T.R.B., J.W. B.C., R.I.G. and P.S.;

- 605 Investigation, J.M.; Software, J.M., B.C.; Writing Original Draft, J.M. and P.S.; Writing
- 606 Review & Editing, J.M., T.R.B., R.I.G., J.W., and P.S. ; Funding Acquisition, P.S. ;
- 607 Resources, J.M., T.R.B., J.W. B.C., R.I.G. and P.S.; Supervision, T.R.B and P.S.
- 608

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- 838 839

# 840841 *Figure Captions*

#### 842 Figure 1. Paradigm and traditional EEG and fMRI results

- 843 A, 3-AFC task where stimulus evidence for each category is modulated by varying the
- 844 phase coherence in the images. **B**, Example of face images with high stimulus evidence
- 845 (high coherence: 50%) and low stimulus evidence (low coherence: 35%). C, Behavioral
- 846 performance shows significant differences, as a function of stimulus evidence, in
- accuracy (p<  $10^{-12}$ , paired t-test) and **D**, response time (p<  $10^{-8}$ , paired t-test) across the
- group. E, Grand average stimulus-locked event related potentials (ERPs) for electrode Pz
- show that differences in stimulus evidence span the time from stimulus to response. F,
- 850 fMRI analysis showing cortical areas correlated with high (red) vs. low (blue) stimulus
- evidence across the entire trial (Z > 2.57 with p < 0.01 Family-Wise Error cluster
- 852 corrected).

# 853 Figure 2. Temporally precise trial-to-trial EEG variability tags brain regions during 854 decision-making

855 A, Illustration of how trial-to-trial variability of neural activity in spatially distinct 856 cortical areas can be used to tag brain regions. In this hypothetical example Region 1 is 857 involved in sensory encoding while Region 2 integrates sensory evidence to form a 858 decision (in NHP literature, Region 1 might represent MT, while Region 2 LIP). Neural 859 activity across the trial is shown for two stimulus types, one with high sensory evidence 860 for the choice (red curves) and one with low sensory evidence (blue curves). Also 861 shown are two temporal windows ( $\tau_1$  and  $\tau_2$ ) that represent different times during the 862 trial. **B**, Linear classifiers are trained to separate trials based on the two levels of stimulus 863 evidence at specific temporal windows. Shown are classifiers (parameterized by weight

864	vectors $w_1$ and $w_2$ ) for two temporal windows ( $\tau_1$ and $\tau_2$ ) with respect to two EEG sensors
865	(for simplicity only two dimensions of the full N=43 sensor space are shown. Though
866	the component hyperplane is optimal for the full 43 dimensions, when projected to a line
867	in two dimensions for illustration, it may appear that the separation is sub-optimal). This
868	yields an EEG discriminant component for each temporal window. Variability along
869	these components serves as a unique feature vector for temporally tagging the BOLD
870	data—e.g. variability along an EEG component trained with data from $\tau_1$ tags BOLD
871	voxels with time $\tau_1$ while variability along an EEG component trained with data from $\tau_2$
872	tags them with $\tau_2$ .
873	
874	Figure 3. Encoding models based on stimulus derived features versus EEG
875	variability
875 876	A, A traditional encoding model used in fMRI analysis extracts a set of features from the
	·
876	A, A traditional encoding model used in fMRI analysis extracts a set of features from the
876 877	A, A traditional encoding model used in fMRI analysis extracts a set of features from the stimulus that are potentially representative of low level structure and high level semantics
876 877 878	A, A traditional encoding model used in fMRI analysis extracts a set of features from the stimulus that are potentially representative of low level structure and high level semantics (green box). Weights are learned to model how these stimulus features are encoded in
876 877 878 879	A, A traditional encoding model used in fMRI analysis extracts a set of features from the stimulus that are potentially representative of low level structure and high level semantics (green box). Weights are learned to model how these stimulus features are encoded in the fMRI BOLD signal. The resulting encoding model is used to make predictions based
876 877 878 879 880	A, A traditional encoding model used in fMRI analysis extracts a set of features from the stimulus that are potentially representative of low level structure and high level semantics (green box). Weights are learned to model how these stimulus features are encoded in the fMRI BOLD signal. The resulting encoding model is used to make predictions based on how well different voxels predict the features from novel stimuli. For example, one
876 877 878 879 880 881	A, A traditional encoding model used in fMRI analysis extracts a set of features from the stimulus that are potentially representative of low level structure and high level semantics (green box). Weights are learned to model how these stimulus features are encoded in the fMRI BOLD signal. The resulting encoding model is used to make predictions based on how well different voxels predict the features from novel stimuli. For example, one can create maps of the brain that are labeled based on the stimulus features that each
876 877 878 879 880 881 882	A, A traditional encoding model used in fMRI analysis extracts a set of features from the stimulus that are potentially representative of low level structure and high level semantics (green box). Weights are learned to model how these stimulus features are encoded in the fMRI BOLD signal. The resulting encoding model is used to make predictions based on how well different voxels predict the features from novel stimuli. For example, one can create maps of the brain that are labeled based on the stimulus features that each voxel represents. <b>B</b> , The same encoding model concept applied to EEG variability (EEG
876 877 878 879 880 881 881 882 883	<b>A</b> , A traditional encoding model used in fMRI analysis extracts a set of features from the stimulus that are potentially representative of low level structure and high level semantics (green box). Weights are learned to model how these stimulus features are encoded in the fMRI BOLD signal. The resulting encoding model is used to make predictions based on how well different voxels predict the features from novel stimuli. For example, one can create maps of the brain that are labeled based on the stimulus features that each voxel represents. <b>B</b> , The same encoding model concept applied to EEG variability (EEG encoding model). Instead of features being estimated from the stimulus, they are derived

traditional encoding model, predictions on novel stimuli can be done to test the model

and results can be used to construct a map —in this case a map of the brain that shows

- the timing of the EEG component variability that each voxels represents.
- 890

#### 891 Figure 4. EEG discrimination and encoding model results

892 A, Group average area under the receiver operating curve (AUC) for the sliding window

893 logistic regression EEG discrimination analysis, comparing high versus low stimulus

894 evidence trials; standard error across subjects is shown with shading. B, A single subject's

discriminating y-value distributions for high (red) and low stimulus evidence (blue) trials

for two EEG time points (225ms and 600ms). C, Significant fMRI voxels resulting from

the group level analysis for the encoding model (p < 0.01 TFCE-False Discovery Rate

898 (FDR) corrected). Activity is seen encompassing early visual processing regions,

attention networks, and the task positive network. **D**, A random subset of 100 (50 for

900 each stimulus evidence condition) from 700 total trials of the actual (circle) and predicted

901 (diamond) BOLD responses from the encoding model, for an example subject at a single

902 voxel (MNI X/Y/Zmm: -27/-54/-15, r=0.206,  $p<10^{-6}$ ). High and low stimulus evidence

903 trials are shown separately for clarity. E, The averaged correlation of the predicted y-

values with the true y-values across the trial duration. Blue shading represents the

905 standard error across subjects. Grey shading indicates significant time windows (p < 0.05

906 FDR-corrected).

907

908 Figure 5. Group-level encoding model weights results show neural activation cascade

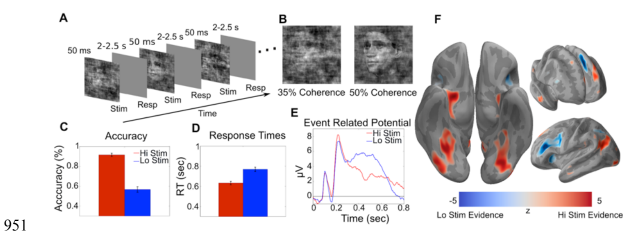
909	Subset of thresholded (p< 0.05 FDR-Corrected, k=10) group level statistical parametric
910	maps created by stTFCE randomization procedure on the encoding model weight
911	matrices show the progression of spatial activity across the trial. Activation can be seen
912	early in the trial in the occipital regions while progressing more anteriorly later in the trial
913	to executive control areas. Activations in red indicate areas where high stimulus evidence
914	trials had larger activations than low stimulus evidence trials, and blue the inverse.
915	
916	Figure 6. Spatial-temporal event-related activations show coordinated reactivations.
917	A, Union across time windows of significant voxels for high (red) and low (blue)
918	stimulus evidence activations. Voxels with activations for both high and low conditions
919	(at different time windows) are displayed in green. Also shown are the encoding model
920	weights for specific voxels, including fusiform gyrus (FG-R):36/-51/-18, (FG-L):-42/-
921	42/-18, superior lateral occipital cortex (sLOC):24/-63/36, superior parietal lobule
922	(SPL):27/-51/54, anterior cingulate cortex (ACC):-6/24/30, intraparietal sulcus (IPS):-
923	30/-60/39, middle frontal gyrus (MFG):-45/27/30, middle temporal gyrus (MT):-57/-
924	60/0. Asterisks indicate significant windows. <b>B</b> , Sequence of significant weights showing
925	a "replay" of the network after the onset of ACC activation (shaded ellipse). "Replay" is
926	faster than the initial stimulus driven sequence and strongest for low evidence trials.
927	
928	Figure 7. Trial-to-trial reactivation correlates with decision confidence.
929	Trial-to-trial reactivation amplitude ( $Y_{j,i}^{R}$ – see Methods) of "replay" correlates with
930	confidence proxy for both high (A) and low (B) stimulus evidence conditions. Error bars

931 represent standard errors across subjects. (C), Stimulus-locked replay activation clusters

932	and feature importance. (inset) Regions of interest used in computing the reactivation
933	values for computing confidence proxy correlations. These regions were taken from
934	significant group activations from 600-800ms post stimulus. Regions were then clustered
935	(>48 voxels) and a secondary analysis for feature importance was performed. Here, we
936	removed each cluster before computing trial-to-trial reactivations and compared the slope
937	of reactivation x confidence proxy when all clusters were present. Panel C shows the
938	ranking of feature importance for each cluster (more negative % change = more
939	importance). Negative changes in slopes show that by removing that cluster the slope of
940	the correlation between reactivation and confidence decreases, indicating the importance
941	of that cluster. Increases in slope indicate that the correlation is higher with that region
0.40	1

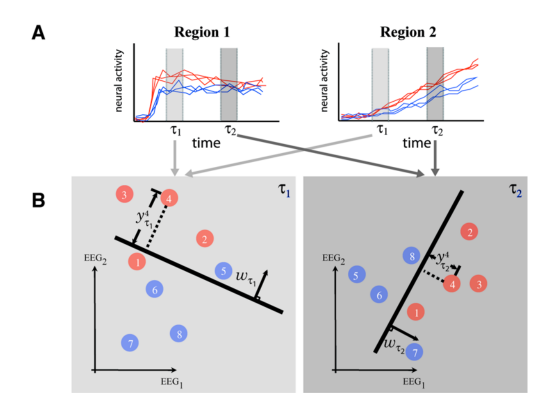
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- **Figures:**



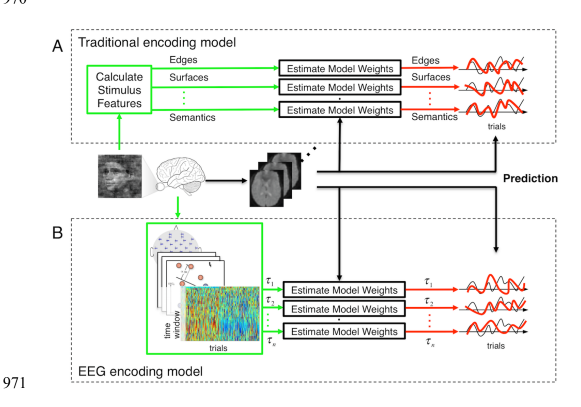
*Figure 1.* 

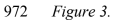


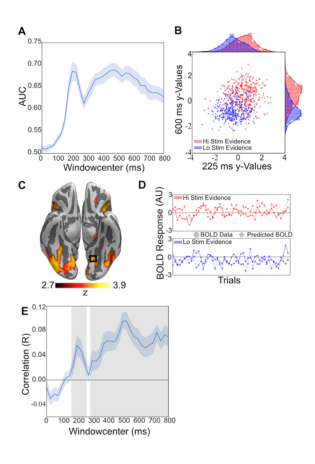


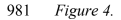
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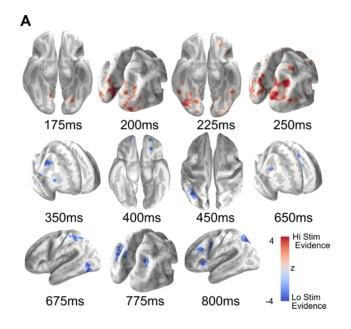






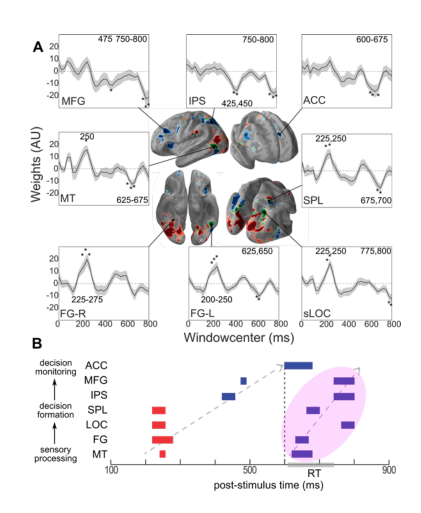




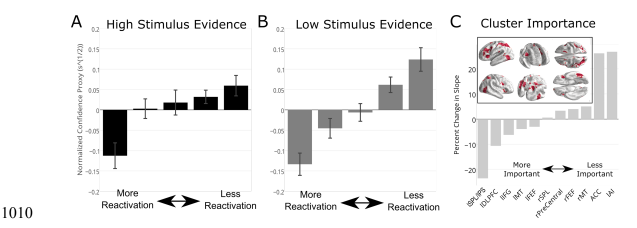


- *Figure 5.*





1000 Figure 6.



1011 Figure 7.