1 Humans strategically shift decision bias by flexibly

2 adjusting sensory evidence accumulation in visual cortex

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31 Abstract

32 Decision bias is traditionally conceptualized as a flexible internal reference against 33 which sensory evidence is compared. Here, in contrast, we show that experimental 34 manipulation of decision bias adjusts the rate of evidence accumulation in visual cortex towards one of the choice alternatives. Participants performed a visual detection task 35 36 during EEG recordings. We experimentally manipulated participants' response criterion using different stimulus-response reward contingencies, inducing liberal and 37 38 conservative decision biases in different conditions. Drift diffusion modeling of choice 39 behavior revealed that an experimentally induced liberal decision bias specifically 40 biased the rate of sensory evidence accumulation towards 'yes' choices. In visual cortex, the liberal bias manipulation suppressed prestimulus 8-12 Hz (alpha) power, 41 42 which in turn boosted cortical stimulus-related activity in the 59–100 Hz (gamma) 43 range. Together, these findings show that observers can intentionally control cortical 44 excitability to strategically bias evidence accumulation towards the decision bound that maximizes reward within a given ecological context. 45

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47 Introduction

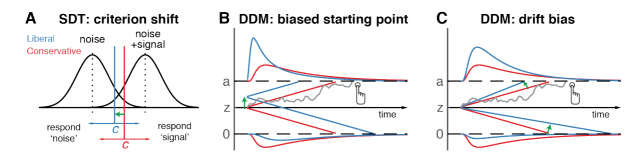
Perceptual decisions arise not only from the evaluation of sensory evidence, but are often biased towards one or the other response alternative by environmental factors, for example as a result of task instructions and/or stimulus-response reward contingencies (White & Poldrack, 2014). The ability to willfully control decision bias enables the behavioral flexibility required to survive in an ever-changing and uncertain environment. But despite its central and important role in decision making, the neural mechanisms underlying decision bias are not fully understood.

55 The traditional account of decision bias comes from signal detection theory 56 (SDT) (Green & Swets, 1966). In SDT, decision bias is quantified by estimating the 57 relative position of a decision point or 'criterion' in between sensory evidence 58 distributions for noise and signal (see Figure 1A). In this framework, a more liberal 59 decision bias arises by moving the criterion closer towards the noise distribution (see 50 green arrow in Figure 1A). Although SDT has been very successful at quantifying 51 decision bias, it has not done much to elucidate the mechanism behind it. One reason

for this lack of insight may be that SDT does not have a temporal component to track
how decisions are reached over time (Fetsch, Kiani, & Shadlen, 2014).

As an alternative to SDT, the drift diffusion model (DDM) conceptualizes 64 perceptual decision making as the accumulation of noisy sensory evidence over time 65 66 into an internal decision variable (Bogacz, Brown, Moehlis, Holmes, & Cohen, 2006; Gold & Shadlen, 2007; Ratcliff & McKoon, 2008). A decision in this model is made 67 when the decision variable crosses one of two decision bounds corresponding to the 68 69 choice alternatives. Within this framework, a strategic decision bias imposed by the 70 environment can be modelled in two different ways: either by moving the starting point 71 of evidence accumulation closer to one of the boundaries (see green arrow in Figure 72 1B), or by biasing the rate of the evidence accumulation process itself towards one of 73 the boundaries (see green arrow in Figure 1C). In both the SDT and DDM frameworks, 74 decision bias shifts have little effect on the sensitivity of the observer when 75 distinguishing signal from noise; they predominantly affect the relative response ratios 76 (and in the case of DDM the speed with which one or the other decision bound is 77 reached). There has been some evidence to suggest that decision bias induced 78 through shifting the response criterion is best characterized by a drift bias in the DDM 79 (Urai, de Gee, & Donner, 2018; White & Poldrack, 2014). However, the drift bias 80 parameter has as yet not been related to a well-described cortical mechanism.





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83 Figure 1 I Theoretical accounts of decision bias. A. The SDT account of decision bias. In this 84 framework, signal and noise+signal distributions are plotted as a function of the strength of internal 85 sensory evidence. Here, the decision point (or criterion) that determines whether to indicate signal 86 presence or absence is plotted as a vertical criterion line c, reflecting the degree of decision bias. c can 87 be shifted left- or rightwards to respectively model a more liberal or conservative bias (green arrow 88 indicates a shift to liberal). In drift diffusion models (DDMs, panels B. and C.), decisions are modelled 89 in terms of a dynamic process of sensory evidence accumulation. When sensory input is presented, 90 evidence starts to accumulate (drift) over time departing from starting point Z. The rate at which

91 evidence accumulates is called the drift rate, and a response is given when it either crosses decision 92 boundary a (signal presence) or decision boundary 0 (no signal). DDMs are fitted to distributions of 93 reaction times obtained over multiple trials. In panels B. and C., reaction time distributions for signal-94 present responses are plotted at the top and reaction time distributions for no-signal responses are 95 plotted in mirror image at the bottom. DDMs can model bias in two different ways. In panel B., bias is 96 modelled in terms of the DDM starting point Z, which is moved closer or further away from the decision 97 bounds a and 0. In panel C., decision bias is modelled in terms of drift bias, where the rate of evidence 98 accumulation for signal and noise move upwards or downwards in tandem (green arrows indicate a 99 shift to liberal). The predicted reaction time distributions under the models in B. and C. are plotted 100 separately for a liberal and conservative bias above and below the graphs. Panels B. and C. are 101 modified and reproduced with permission from Urai, de Gee, & Donner (2018).

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103 Likewise, there have been a number of reports about a correlational 104 relationship between cortical activity and decision bias. For example, spontaneous 105 trial-to-trial variations in prestimulus oscillatory activity in the 8-12 Hz (alpha) band 106 have been shown to correlate with decision bias and confidence (lemi, Chaumon, Crouzet, & Busch, 2017; Limbach & Corballis, 2016; Samaha, Iemi, & Postle, 2017). 107 108 Relatedly, alpha oscillations have been proposed to be involved in the gating of task-109 relevant sensory information (Jensen & Mazaheri, 2010), possibly encoded in high-110 frequency (gamma) oscillations in visual cortex (Ni et al., 2016; Popov, Kastner, & 111 Jensen, 2017). Although these reports suggest a link between alpha suppression and 112 decision bias, they do not uncover whether pre-stimulus alpha plays an instrumental 113 role in decision bias and how exactly this might be achieved. For example, it is 114 unknown whether an experimentally induced shift in decision bias is implemented in the brain by willfully adjusting pre-stimulus alpha in sensory areas. 115

116 Here, we explicitly investigate these potential mechanisms by employing a task 117 paradigm in which shifts in decision bias were experimentally induced within 118 participants through instruction and asymmetries in stimulus-response reward 119 contingencies during a visual detection task. By applying drift diffusion modeling to the 120 participants' choice behavior, we show that strategically adjusting decision bias 121 specifically affects the rate of sensory evidence accumulation towards one of the two 122 decision bounds. Further, we demonstrate that this drift bias is achieved by flexibly up-123 and down-regulating prestimulus alpha to control the response gain of stimulus-related 124 gamma activity in visual cortex. Critically, we also show that gamma activity accurately

predicts the strength of the evidence accumulation bias within subjects, providing a direct link between the proposed mechanism and decision bias. Together, these findings identify the neural mechanism by which intentional control of cortical excitability is applied to strategically bias perceptual decisions in order to maximize reward in a given context.

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131 **Results**

132 Liberal decision bias manipulation shifts sensory evidence accumulation

133 In three EEG recording sessions, human participants (N = 16) viewed a continuous 134 stream of horizontal, vertical and diagonal line textures alternating at a rate of 25 135 textures/second. The participants' task was to detect an orientation-defined square 136 presented in the center of the screen and report it via a button press (Figure 2A). Trials 137 consisted of a fixed-order sequence of textures (total sequence duration 1 second) embedded in the continuous stream. A square appeared in the fifth texture of a trial in 138 139 75% of the presentations (target trials), while in 25% a homogenous diagonal texture 140 appeared in the fifth position (nontarget trials). Although the onset of trials within the 141 continuous stream of textures was not explicitly cued, the similar distribution of 142 reaction times in target and nontarget trials suggests that participants employed the 143 temporal structure of the task even when no target appeared (Figure S1A). Consistent 144 significant EEG power modulations after trial onset even for non-target trials further 145 confirm that subjects registered trial onsets even without an explicit cue, plausibly using the onset of a fixed order texture sequence as an implicit cue (Figure S1B). 146

147 In alternating nine-minute blocks of trials, we actively biased participants' 148 perceptual decisions by instructing them either to report as many targets as possible 149 ("Detect as many targets as possible!"; liberal condition), or to only report high-150 certainty targets ("Press only if you are really certain!"; conservative condition). 151 Participants were free to respond at any time during a block whenever they detected 152 a target. We provided auditory feedback following missed targets (misses) in the liberal 153 condition and falsely detected targets (false alarms) in the conservative condition and 154 applied monetary penalties for these errors (Figure 2A; see Methods for details).

155 Participants reliably adopted the intended criterion shift (see Figure 2B showing 156 that both the hit rate and the false alarm rate went down in tandem as a consequence of a more conservative criterion). The difference between hit rate and false alarm rate 157 158 was not significant between conservative and liberal (p = 0.81, right bars in Figure 2B). 159 However, detection performance (sensitivity) computed using standard SDT d' 160 (reflecting the distance between the noise and signal distributions in Figure 1A) was 161 slightly higher during conservative (liberal: d' = 2.0 (s.d. 0.90), versus conservative: d'162 = 2.31 (s.d. 0.82), p = 0.0002, two-sided permutation test, 10,000 permutations, see 163 Figure 2C, left bars)(Green & Swets, 1966). We also computed the standard SDT 164 criterion measure c reflecting bias in the decision process (see the blue and red 165 vertical lines in Figure 1A), which uncovered a strong experimentally induced criterion 166 shift (liberal: c = -0.13 (s.d. 0.4), versus conservative: c = 0.73 (s.d. 0.36), p = 0.0001, permutation test, see Figure 2C, right bars). 167

168 Because the SDT framework is static, we decided to further investigate how 169 bias affected various components of the dynamic decision process by fitting different 170 drift diffusion models (DDMs) to the behavioral data (Figure 1B, C) (Ratcliff & McKoon, 171 2008). DDMs postulate that perceptual decisions are reached by accumulating noisy 172 sensory evidence towards one of two decision boundaries representing the choice 173 alternatives. Crossing one of these boundaries can either trigger an explicit 174 behavioural report to indicate the decision (for 'yes' responses in our experiment), or 175 remain implicit (i.e. without active response, for 'no' decisions in our experiment) 176 (Ratcliff, Huang-Pollock, & McKoon, 2016). We tested two different DDMs that can 177 potentially account for decision bias: one in which the starting point of evidence accumulation moves closer to one of the decision boundaries ('starting point model', 178 179 Figure 1B) (Mulder, Wagenmakers, Ratcliff, Boekel, & Forstmann, 2012), and one in 180 which the evidence accumulation process (called the drift) itself is biased towards one 181 of the boundaries (de Gee et al., 2017) ('drift bias model', see Figure 1C, referred to as drift criterion by Rattclif and McKoon (2008)). In the two respective models, we 182 183 freed either the drift bias parameter (db, see Figure 2D) for the two conditions while 184 keeping starting point (z) fixed across conditions (for the drift bias model), or vice versa 185 (for the starting point model). The drift bias parameter is determined by estimating the 186 contribution of an evidence-independent constant added to the drift (Figure 2B). These

alternative models make different predictions about the shape of the RT distributions
in combination with the response ratios: a shift in starting point produces large
changes in both the leading edge and tail of the distribution, whereas a shift in drift
bias produces large changes only in the tail (Ratcliff & McKoon, 2008; Urai et al.,
2018), also see the RT distributions above and below the evidence accumulation
graphs in Figure 1B and 1C.

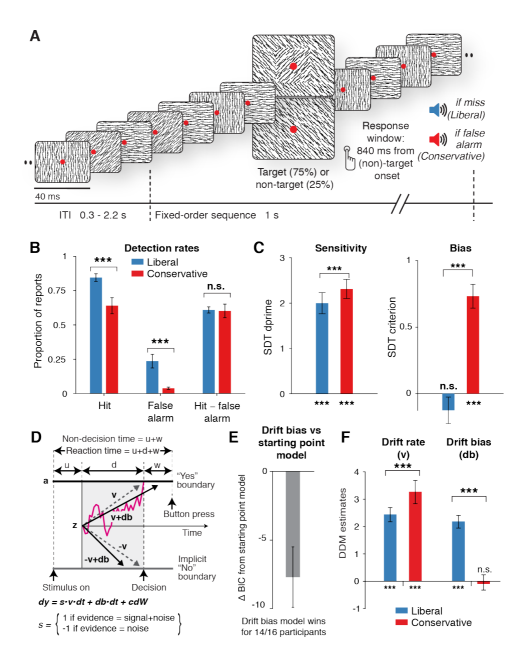
193 We fitted both the starting point and drift bias models to each participant's RT 194 distribution for 'yes' choices and the total number of implicit 'no' choices. In both 195 models, all of the non-bias related parameters (drift rate v, boundary separation a and 196 non-decision time u+w, see Figure 2D) were also allowed to vary by condition. We 197 compared goodness of fit of the models to assess which model best explained the 198 data. We found that the starting point model provided a worse fit to the data, as 199 indicated by higher Bayesian Information Criterion (BIC) estimates than for the drift bias model (Figure 2E, see Methods for details). Specifically, for 14 out of the 16 200 201 participants the drift bias model provided better fits than the starting point model, for 202 ten of which delta BIC was greater than six, indicating strong evidence in favor of the 203 drift bias model. Finally, we compared these models to a model in which both drift bias 204 and starting point were fixed across the conditions, while still allowing the non-bias-205 related parameters to vary per condition. This model provided the lowest goodness of 206 fit (delta BIC greater than six for both models for all participants). See Figure S3 for 207 model fits of the drift bias model for each participant.

208 Given the superior performance of the drift bias model, we further characterized 209 decision making under the bias manipulation using parameter estimates from this 210 model. Drift rate, reflecting the participants' ability to discriminate targets and non-211 targets, was somewhat higher in the conservative compared to the liberal condition 212 (liberal: v = 2.39 (s.d. 1.07), versus conservative: v = 3.06 (s.d. 1.16), p = 0.0001, 213 permutation test, Figure 2F, left bars). An almost perfect correlation across 214 participants between DDM drift rate and SDT d'provided strong evidence that the drift 215 rate parameter captures perceptual sensitivity (liberal, r = 0.97; conservative, r = 0.95, 216 all p-values < 0.005, see Figure S2A).

217 Regarding the DDM bias parameters, the condition-fixed starting point 218 parameter in the drift bias model was smaller than half the boundary separation (i.e. 219 closer to the 'no' boundary: z = 0.24, p < 0.0001, tested against 0.5), indicating an 220 overall conservative starting point across conditions (Figure S2D). Strikingly, however, 221 whereas drift bias was on average not different from zero in the conservative condition 222 (db = -0.04, p = 0.90), drift bias was strongly positive in the liberal condition (db = -0.04, p = 0.90)223 2.08, p = 0.0001; liberal vs conservative: p = 0.0005; Figure 2F, right bars). The overall 224 conservative starting point combined with a condition-specific neutral drift bias 225 explained the conservative decision bias (as quantified by SDT criterion) in the conservative condition (Figure 2C). Likewise, in the liberal condition the overall 226 227 conservative starting point combined with a condition-specific positive drift bias 228 (pushing the drift towards the 'yes' boundary) explained the neutral bias observed with 229 SDT (criterion around zero for liberal, see Figure 2C).

230 Converging with these modelling results, drift bias was strongly anti-correlated 231 across participants with both SDT criterion (liberal, r = -0.83; conservative, r = -0.79, 232 see Figure S2B) and reaction times (liberal, r = -0.66; conservative, r = -0.76, see 233 Figure S2C). The strong correlations between DDM drift rate and SDT d' on the one 234 hand, and DDM drift bias and SDT criterion on the other, provide converging evidence 235 that the SDT and DDM frameworks captured similar underlying mechanisms, while 236 the DDM additionally captured the dynamic nature of perceptual decision making by 237 linking the decision bias manipulation to the evidence accumulation process itself.

238 Finally, the bias manipulation also affected two other parameters in the drift 239 bias model that were not directly related to sensory evidence accumulation: boundary 240 separation was slightly but reliably higher during liberal compared to conservative (p 241 < 0.0001), and non-decision time (comprising time needed for sensory encoding and 242 motor response execution) was shorter during liberal (p < 0.0001)(supplementary 243 Figure S2D). In conclusion, a drift diffusion model of choice behavior implementing a 244 bias in sensory evidence accumulation best explained how participants adjusted to 245 the manipulations of decision bias. In the next sections, we used spectral analysis of the concurrent EEG recordings to identify a plausible neural mechanism that 246 247 implements biased sensory evidence accumulation.



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250 Figure 2 | Biased sensory evidence accumulation underlies experimentally induced liberal 251 decision bias A. Schematic of the visual stimulus and task design. Participants viewed a continuous 252 stream of full-screen diagonally, horizontally and vertically oriented textures at a presentation rate of 40 253 ms (25 Hz). After random inter-trial intervals (range 0.3–2.2 s), a fixed-order sequence (duration 1 s) 254 was presented, embedded in the stream. The fifth texture in each sequence either consisted of a single 255 diagonal orientation (nontarget), or contained an orthogonal orientation-defined square (target of either 256 45° or 135° orientation). Participants decided whether they had just seen a target, reporting detected 257 targets by button press within 840 ms after target onset. Liberal and conservative conditions were

258 administered in alternating nine-minute blocks by penalizing either misses or false alarms, respectively, 259 using aversive tones and monetary deductions. Depicted square and fixation dot sizes are not to scale. 260 B. Average detection rates (hits and false alarms) during both conditions. C. SDT parameters for 261 sensitivity and criterion. D. Schematic and simplified equation of drift diffusion model accounting for 262 reaction time (RT) distributions for explicit 'yes' and implicit 'no' decisions. Decision bias in this model 263 can be implemented by either shifting the starting point of the evidence accumulation (Z), or by adding 264 an evidence-independent constant ('drift bias', db) to the drift rate. See text and Figure 1 for details. 265 Notation: dy, change in decision variable y per unit time dt; v.dt, mean drift (multiplied with 1 for signal 266 + noise (target) trials, and -1 for noise-only (non-target) trials); db·dt, drift bias; and cdW, Gaussian 267 white noise (mean = 0, variance = $c2 \cdot dt$). **E.** The difference in Bayesian Information Criterion (BIC) 268 goodness of fit estimates for the drift bias and the starting point models, A lower delta BIC value 269 indicates a better fit, showing superiority of the drift bias model to account for the observed results. F. 270 Estimated model parameters for drift rate and drift bias in the drift bias model. Error bars, SEM across 271 16 participants. ***p < 0.001; n.s., not significant.

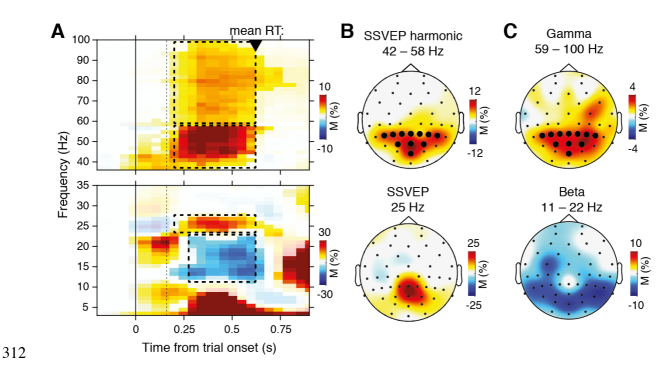
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Task-relevant textures induce stimulus-related responses in visual cortex

274 Sensory evidence accumulation in the visual detection task presumably relies on 275 stimulus-related signals processed in visual cortex. Such stimulus-related signals are typically reflected in cortical population activity exhibiting a rhythmic temporal structure 276 277 (Buzsáki & Draguhn, 2004). Specifically, bottom-up processing of visual information has previously been linked to increased high-frequency (> 40 Hz, i.e. gamma) 278 279 electrophysiological activity over visual cortex (Bastos et al., 2015; Michalareas et al., 2016; Popov et al., 2017; van Kerkoerle et al., 2014). Figure 3A shows time-frequency 280 281 representations of EEG power modulations over posterior cortex for the low and high 282 frequency bands, normalized with respect to the prestimulus baseline period.

283 We observed a total of four distinct stimulus-related power modulations after 284 trial onset: two in the high-frequency range (> 36 Hz, Figure 3A, top panel) and two in 285 the low frequency range (< 36 Hz, Figure 3A, bottom panel). First, a spatially focal 286 modulation in a narrow frequency range around 25 Hz reflecting the steady state visual 287 evoked potential (SSVEP) arising from entrainment by the visual stimulation frequency 288 of our experimental paradigm (Figure 3B, lower panel). A second modulation from 42–58 Hz (Figure 3B, top panel) comprised the first harmonic of the SSVEP, as can 289 290 be seen from their similar topographic distributions (Figure 3B, compare top and lower 291 panel).

292 Third, we observed a 59–100 Hz gamma power modulation (Figure 3C, top 293 panel), after carefully controlling for high-frequency EEG artifacts due to small 294 fixational eye movements (microsaccades) by removing microsaccade-related activity from the data (Hassler, Trujillo-Barreto, & Gruber, 2011; Hipp & Siegel, 2013; Yuval-295 296 Greenberg, Tomer, Keren, Nelken, & Deouell, 2008), and by suppressing non-neural 297 EEG activity through scalp current density transformation (Melloni, Schwiedrzik, 298 Wibral, Rodriguez, & Singer, 2009; Perrin, Pernier, Bertrand, & Echallier, 1989) (see 299 Methods for details). Importantly, the topography of the observed gamma modulation 300 was confined to posterior electrodes (electrodes highlighted in Figures 3B and 3C, top 301 panels), in line with the role of gamma in stimulus-related processing in visual cortex 302 (Ni et al., 2016). Finally, we observed suppression of low-frequency beta (11-22 Hz) 303 activity in posterior cortex, which typically occurs in parallel with enhanced stimulus-304 related gamma activity (Donner & Siegel, 2011; Kloosterman et al., 2015; 305 Meindertsma, Kloosterman, Nolte, Engel, & Donner, 2017; Werkle-Bergner et al., 306 2014)(Figure 3A and 3C, lower panels). In the next section, we used the topographies 307 of the high-frequency post-stimulus effects in visual cortex (Figures 3B and 3C, top 308 panels) to identify a prestimulus neural mechanism that could explain the observed 309 biased evidence accumulation resulting from the experimental decision bias 310 manipulation.

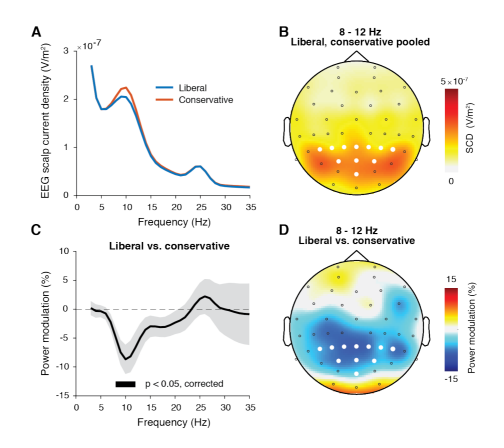


313 Figure 3 | Task-relevant textures induce stimulus-related responses in visual cortex. A. Time-314 frequency representations of high- (top) and low-frequency (bottom) EEG power modulations with 315 respect to the prestimulus period (-0.4 - 0 s), pooled over the two conditions. Saturated colors indicate 316 clusters of significant modulation, cluster threshold p < 0.05, two-sided permutation test across 317 participants, cluster-corrected; N = 15). Solid and dotted vertical lines respectively indicate the onset of 318 the trial and the target stimulus. M, power modulation. B. Scalp maps showing topography of the SSVEP 319 power modulation around 25 Hz (top) and its harmonic from 42 – 58 Hz (bottom), from 0.2 – 0.6 s after 320 trial onset. C. 59-100 Hz gamma power modulation from 0.2 - 0.6 s (top) and concurrent low frequency 321 ('beta') power suppression from 11 - 22 Hz; see dashed outlines on time-frequency representations in 322 A. Thick dots indicate electrodes used for the time-frequency representations in A, and which were 323 selected for further analysis.

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325 Adopting a liberal decision bias suppresses prestimulus alpha power

326 As a first step, we examined prestimulus power between 0.8 and 0.2 s before trial 327 onset, using the same electrodes that showed the strongest post-stimulus effects 328 (Figure 4A). This uncovered a highly specific modulation in the alpha range, which we 329 confirmed to be strongest over the same cortical region that showed strong modulation 330 in the gamma range (Figure 4B, white dots indicate electrodes showing stimulus-331 related gamma modulation). Indeed, when expressing spectral power during the 332 liberal condition as the percentage signal change from the conservative condition, we 333 observed a statistically significant cluster of suppressed frequencies precisely in the 334 8-12 Hz frequency range (p < 0.05, cluster-corrected for multiple comparisons) 335 (Figure 4C), which again showed a posterior topography (Figure 4D). This shows that 336 an experimentally induced liberal decision bias suppresses prestimulus alpha power, 337 suggesting that alpha modulations are a hallmark of strategic bias adjustment rather 338 than a mere correlate of spontaneous shifts in decision bias. Importantly, this finding 339 implies that humans are able to actively control prestimulus alpha power in visual cortex, plausibly acting to bias sensory evidence accumulation towards the response 340 alternative that maximizes rewards. 341



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343 Figure 4 | Adopting a liberal decision bias suppresses prestimulus alpha power. A. Low-344 frequency power spectra of prestimulus neural activity for both conditions based on the electrodes that 345 show large post-stimulus power modulations in Figure 3B and C (top panels). B. Scalp map of raw 346 prestimulus EEG alpha power (8–12 Hz neural activity between 0.8 and 0.2 s before sequence onset), 347 pooled over conditions. White symbols indicate visual cortical electrodes used for the power spectra in 348 A. and C. C. Liberal versus conservative power spectrum. Black horizontal bar indicates statistically 349 significant frequency range (p < 0.05, cluster-corrected for multiple comparisons, two-sided). Error bars, 350 SEM across participants (N = 15). **D.** The corresponding scalp map of power modulation in the liberal 351 condition, expressed as percent signal change from the conservative condition.

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354 Alpha suppression enhances the gain of cortical gamma responses

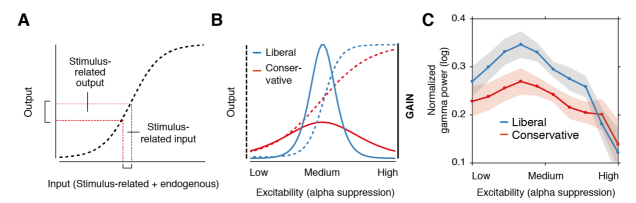
How could suppression of prestimulus alpha activity bias the process of sensory evidence accumulation? One possibility is that alpha suppression influences evidence accumulation by modulating the susceptibility of visual cortex to sensory stimulation, a phenomenon dubbed 'neural excitability' (lemi et al., 2017; Jensen & Mazaheri, 2010). We explored this possibility using a theoretical framework put forward by Rajagovindan and Ding (2011). This framework assumes that the relationship between total synaptic input activity that a neuronal ensemble receives and the total

362 output activity it produces is characterized by a sigmoidal function (Figure 5A); a notion 363 that is biologically plausible (Destexhe, Rudolph, Fellous, & Sejnowski, 2001; Freeman, 1979). Within this framework, both sensory input (i.e. as a result of sensory 364 365 stimulation) and ongoing fluctuations in endogenous neural activity (i.e. levels of 366 neural excitability) contribute to the synaptic input into visual cortex. The isolated effect 367 of sensory input on the total output (i.e. the gain of the output response as caused by 368 an input stimulus; see marked interval in Figure 5A), can then be expressed as the 369 first order derivative (the slope) of the sigmoid in Figure 5A. In our experiment, 370 stimulus-related input activity can be assumed to be more or less constant across trials 371 since the same stimulus sequence was shown in each trial (see Figure 2A). Thus, 372 modulations in the stimulus-related output gain generated in visual cortex are largely 373 determined by the brain's excitability state. This can be seen in Figure 5B, where the 374 stimulus-related output gain (the first order derivative, or slope from Figure 5A) is 375 plotted as a function of neural excitability, yielding an inverted-U shaped function.

376 Figure 5B then shows the *effective range* in which the impact of neural 377 excitability on the stimulus-related output response is largest, while its impact during 378 low and high excitability is lower. When heightened excitability in the liberal condition 379 is observed, this framework predicts enhanced output activity in visual cortex when 380 compared to the conservative condition (Figure 5B), in particular when excitability 381 differences between conditions occur in its effective range (Rajagovindan & Ding, 382 2011) (i.e. steeper slope of the solid blue curve compared to the red curve in Figure 383 5B).

384 We tested this model in our data by following the method put forward by 385 Rajagovindan and Ding (2011), in which we operationalized neural excitability as pre-386 stimulus alpha (Jensen & Mazaheri, 2010), and stimulus-related output gain as post-387 stimulus gamma (Ni et al., 2016). We exploited the large number of trials per 388 participant per condition across the multiple sessions in our study (range 543 to 1391 389 trials) by sorting each participant's trials into ten excitability bins based on equal-sized 390 ranges of descending (log-transformed) prestimulus alpha power (indicating 391 increasing excitability), separately for the conservative and liberal conditions. We 392 subsequently computed and averaged the (log-transformed) gamma power across the 393 trials within each excitability bin. Following Rajagovindan and Ding (2011), we

- removed individual differences in overall gamma power magnitude by subtracting the
 lowest binned gamma observation in the conservative condition from all observations.
 Finally, we plotted normalized gamma power as a function of excitability, separately
- 397 for liberal and conservative (Figure 5C, see Methods for details).



399 Figure 5 | Neural excitability boosts visual cortical responses by enhancing gain. A. Theoretical 400 response gain model, which describes the transformation of input (both stimulus-related and 401 endogenous) to the total output activity in visual cortex as a sigmoidal function. B. Model predictions. 402 Stimulus-related output responses (solid lines) are formalized as the first derivative of the sigmoidal 403 functions (dotted lines), resulting in inverse-U shaped response gain functions. The model predicts that 404 a liberal decision bias increases the steepness of the sigmoidal function (right) compared to a 405 conservative bias (left), resulting in stronger stimulus-related responses due to higher gain 406 (Rajagovindan & Ding, 2011). C. Corresponding empirical test. Log-transformed gamma activity 407 (normalized within participants by subtracting the minimum gamma power during the conservative 408 condition from all bins) plotted as a function of neural excitability. Error bars, within-subject SEM across 409 participants (N = 14).

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411 The resulting plot indeed closely follows an inverted-U shaped relationship 412 between excitability and stimulus-related gamma activity for both conditions, with 413 particularly low gamma responses for the highest excitability bins (Figure 5C). 414 Critically, average gamma power was higher in the liberal than in the conservative 415 condition, except during the highest excitability bins (Figure 5C, rightmost two data 416 points). Indeed, the flanks of the inverted-U curve for the liberal condition were steeper 417 for the liberal condition, suggesting increased response gain. A three-way repeated 418 measures ANOVA with factors condition (conservative, liberal), brain activity type 419 (prestimulus alpha, poststimulus gamma) and bin level (1–10) revealed a significant 420 three-way interaction (F(9,117) = 2.96, p = 0.003, partial η^2 = 0.19, Greenhouse-

421 Geisser corrected p = 0.046). Importantly, the marginally significant quadratic contrast (F(1,13) = 3.47, p = 0.085, partial η^2 = 0.21) fitted this interaction almost as well as a 422 423 linear contrast (F(1,13) = 4.69, p = 0.049, partial η^2 = 0.265). This three-way quadratic interaction effect indeed suggests a more steeply U-shaped curve for gamma 424 425 responses in the liberal condition, in line with enhanced gain. Taken together, these 426 findings indicate that increased excitability during the liberal condition boosted input-427 related activity, which in turn indiscriminately biased sensory evidence accumulation 428 towards 'yes' responses. In the next section, we confirm a direct link between drift bias 429 and cortical stimulus response gain as measured through gamma.

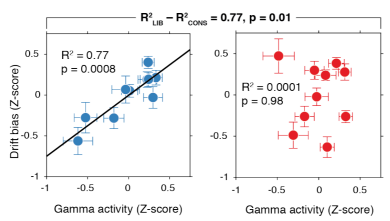
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431 Visual cortical gamma activity predicts strength of evidence accumulation bias

The findings presented so far suggest that behaviorally, a liberal decision bias shifts 432 433 evidence accumulation towards 'yes' responses (drift bias in the DDM), while neurally 434 it results in an increase of prestimulus cortical excitability concomitant with post-435 stimulus response gain expressed in gamma modulation in visual cortex. In a final 436 analysis, we asked whether increases in gamma activity are directly related to a 437 stronger drift bias. We predicted such a direct correspondence during the liberal 438 condition, in which both drift bias and gamma activity were increased (see Figures 2F 439 and 5C), but not during the conservative condition, in which drift bias was around zero 440 and gamma was weaker than during liberal.

441 To test these predictions, we again applied the drift bias DDM to the behavioral 442 data, but now freed the drift bias parameter not only for the two conditions, but also 443 for the ten alpha suppression bins used to show the inverted-U-shaped relationship 444 between excitability and stimulus-related gamma (see Figure 5C). We normalized the 445 bin-resolved drift bias and gamma scalar values by z-scoring within each participant 446 to remove individual differences in their ranges and averaged across participants 447 within each alpha (excitability) bin. Finally, we directly tested the correspondence 448 between drift bias and gamma using a within-subject group regression. Gamma 449 activity indeed accurately predicted drift bias in the liberal condition ($R^2(9) = 0.77$, p = 450 0.0008, Figure 6 left panel). In contrast, drift bias was not well predicted by the 451 corresponding gamma activity in the conservative condition ($R^2(9) = 0.0001$, p = 0.98,

Figure 6 right panel), which is perhaps unsurprising given the fact that drift bias was 452 453 around zero in the conservative condition (see Figure 2F). Accordingly, predictive power was significantly greater in the liberal than in the conservative condition (R²LIB 454 $-R^{2}_{CONS} = 0.77$, p = 0.01). The increase in gamma power in liberal versus conservative 455 456 also predicted the increase in drift bias across the conditions ($R^2(9) = 0.56$, p = 0.0126, 457 Figure S4), further suggesting that the experimental bias manipulation indeed 458 enhanced gamma activity across excitability bins. We obtained qualitatively similar 459 results without averaging across participants, but instead correlating across bins of all 460 participants together using either ten or five bins per participant, suggesting these 461 effects were not driven by single participants (data not shown). Taken together, these 462 results show that enhanced post-stimulus gamma activity during the liberal condition 463 underlies the evidence accumulation bias reflected in the drift bias parameter of the drift diffusion model. 464



465

Figure 6 I Visual cortical gamma activity predicts strength of evidence accumulation bias. Linear
regression of drift bias on gamma activity, separately for the two conditions. Gamma activity accurately
predicts drift bias in the liberal (left), but not the conservative condition (right). Each dot represents an
excitability bin and is obtained after averaging across participants (N = 14, see Methods for details).
Error bars, SEM across participants.

471

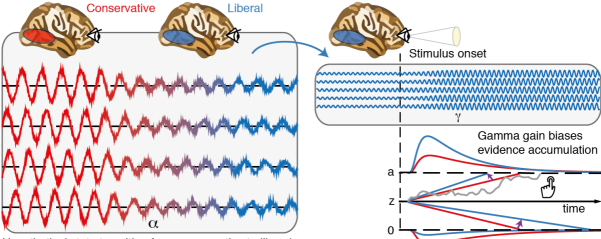
472 **Discussion**

Humans possess a remarkable ability to strategically shift decision biases in order to flexibly adapt to the environment and maximize rewards. Traditionally, bias has been conceptualized in SDT as a criterion threshold that can be shifted towards or away from a noise or signal distribution. To date, however, the neural underpinnings of such bias shifts have remained elusive. Here, in contrast, we use a DDM drift bias model to demonstrate that an experimentally induced bias shift affects the process of sensory
evidence accumulation itself, rather than shifting a threshold entity as SDT implies.
Moreover, we reveal the neural signature of drift bias, by showing that a liberal
decision bias increases alpha suppression (neural excitability) of visual cortex.

482 Although previous studies have shown correlations between suppression of prestimulus alpha (8-12 Hz) power and a liberal decision bias (lemi et al., 2017; 483 484 Limbach & Corballis, 2016), these studies have not established the effect of 485 experimentally induced bias shifts. In the current study, by experimentally 486 manipulating decision bias we show for the first time that prestimulus alpha plays an 487 instrumental, and not merely a correlational role in decision bias. Further, we show 488 that alpha suppression in turn boosts stimulus-related gamma activity through 489 increased cortical response gain. Critically, gamma activity accurately predicted the 490 strength of the drift bias parameter in the DDM drift bias model. Together, these 491 findings show for the first time that humans are able to actively implement decision 492 biases by flexibly adapting neural excitability to strategically shift sensory evidence 493 accumulation towards one of two decision bounds.

494 Based on our results, we propose that decision biases are implemented by 495 flexibly adjusting neural excitability in visual cortex. Figure 7 summarizes this 496 proposed mechanism graphically by visualizing a hypothetical transition in neural 497 excitability following an experimentally induced liberal decision bias, as reflected in 498 visual cortical alpha suppression (left panel). This increased excitability translates into 499 stronger gamma-band responses following stimulus onset (right panel, top). This 500 increased gamma gain finally biases evidence accumulation towards the 'ves' decision 501 boundary during a liberal state, resulting in more 'yes' responses, whereas 'no' 502 responses are decimated (blue RT distributions; right panel, bottom). Our 503 experimental manipulation of decision bias in different blocks of trials suggests that 504 decision makers are able to control this biased evidence accumulation mechanism 505 willfully by adjusting excitability, as reflected in alpha.

506



507 Hypothetical state transition from conservative to liberal

Figure 7 I Illustrative graphical depiction of the excitability state transition from conservative to liberal, and subsequent biased evidence accumulation under a liberal bias. The left panel shows the transition from a conservative to a liberal stimulus block. The experimental induction of a liberal decision bias causes alpha suppression in visual cortex, which increases neural excitability. The right top panel shows increased gamma-gain for incoming sensory evidence under conditions of high excitability. The right bottom panel shows how increased gamma-gain causes a bias in the drift rate, resulting in more 'target present' responses than in the conservative state.

515

516 A neural mechanism that could underlie bias-related alpha suppression may be 517 under control of the catecholaminergic neuromodulatory systems, consisting of the 518 noradrenaline-releasing locus coereleus (LC) and dopamine systems (Aston-Jones & 519 Cohen, 2005). These systems are able to modulate the level of arousal and neural 520 gain, and show tight links with pupil responses (de Gee et al., 2017; de Gee, Knapen, 521 & Donner, 2014; Joshi, Li, Kalwani, & Gold, 2015; McGinley, David, & McCormick, 522 2015). Accordingly, prestimulus alpha power suppression has also recently been 523 linked to pupil dilation (Meindertsma et al., 2017). From this perspective, our results 524 reconcile previous studies showing relationships between a liberal bias, suppression of spontaneous alpha power and increased pupil size. Consistent with this, a recent 525 526 monkey study observed increased neural activity during a liberal bias in the superior 527 colliculus (Crapse, Lau, & Basso, 2018), a mid-brain structure tightly interconnected 528 with the LC (Joshi et al., 2015). Taken together, a more liberal within-person bias 529 (following experimental instruction) might activate neuromodulatory systems that 530 subsequently increase cortical excitability and enhance sensory responses for both stimulus and 'noise' signals in visual cortex, thereby increasing a person's propensity
for 'yes' responses (lemi et al., 2017).

533 Rather than a link between alpha activity and decision bias, several previous 534 studies have reported a link between alpha and task performance, particularly in the 535 phase of alpha oscillations (Busch, Dubois, & VanRullen, 2009; Mathewson, Gratton, Fabiani, Beck, & Ro, 2009). Our findings can be reconciled with those by considering 536 537 that detection sensitivity in many previous studies was often quantified in terms of raw 538 stimulus detection rates, which do not dissociate objective sensitivity from response 539 bias (see Figure 2B) (Green & Swets, 1966). Indeed, our findings are in line with recently reported links between decision bias and spontaneous fluctuations in 540 541 excitability (lemi et al., 2017; lemi & Busch, 2017; Limbach & Corballis, 2016), 542 suggesting an active role of neural excitability in decision bias.

543 Relatedly, a concern regarding our findings could be that the observed change in cortical excitability reflects a change in detection sensitivity (drift rate) rather than 544 545 an intentional bias shift. This is unlikely because that would predict effects opposite to 546 those we observed. We found increased excitability in the liberal condition compared 547 to the conservative condition. If this were related to improved detection performance, 548 one would predict higher sensitivity in the liberal condition, while in fact we found 549 higher sensitivity in the conservative condition (compare drift rate to drift bias in both 550 conditions in Fig. 2C). This finding convincingly ties cortical excitability in our paradigm 551 to a strategically applied bias shift, as opposed to a change in detection sensitivity. 552 Convergently, other studies also report a link between prestimulus low-frequency EEG 553 activity and subjective perception, but not objective task performance (Benwell et al., 554 2017; lemi & Busch, 2017).

555 Summarizing, our results show that stimulus-related responses are boosted 556 during a liberal decision bias due to increased cortical excitability, in line with recent 557 work linking alpha power suppression to response gain (Peterson & Voytek, 2017). 558 Future studies can now establish whether this same mechanism is at play in other 559 subjective aspects of decision-making, such as confidence and meta-cognition 560 (Fleming, Putten, & Daw, 2018; Samaha et al., 2017) as well as in a dynamically 561 changing environment (Norton, Fleming, Daw, & Landy, 2017). Explicit manipulation

562 of cortical response gain during a bias manipulation by pharmacological manipulation of the noradrenergic LC-NE system (Servan-Schreiber, Printz, & Cohen, 1990) or by 563 enhancing occipital alpha power using transcranial stimulation (Zaehle, Rach, & 564 565 Herrmann, 2010) would further establish the underlying mechanisms involved in 566 decision bias. In the end, although one may be unaware, every decision we make is 567 influenced by biases that operate on the noisy evidence accumulation process 568 towards one of the decision bounds. Understanding how these biases affect our 569 decisions is key to becoming aware of these biases (Pleskac, Cesario, & Johnson, 570 2017), allowing us to control or invoke them adaptively. Pinpointing the neural 571 mechanisms underlying bias in an elementary perceptual task (as used here) may 572 paves the way for understanding how more abstract and high-level decisions are 573 modulated by decision bias (Tversky & Kahneman, 1974).

574

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582

583 Author Contributions

584 NAK and JJF designed research, NAK performed research, NAK, JWdG and JJF 585 analyzed data, MWB and DDG provided theoretical background, NAK, JJF, UL, MWB, 586 DDG, JWdG wrote the paper, NAK, JJF, UL, MWB, DDG, JWdG edited and 587 commented on the manuscript.

588

589 **Declaration of Interests**

590 The authors declare no competing interests.

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780

781 Materials and Methods

782 **Participants** Sixteen participants (eight female, mean age 24.1 years, ± 1.64) took 783 part in the experiment, either for financial compensation or in partial fulfillment of first 784 year course requirements. Each participant completed three experimental sessions on 785 different days, each session lasting ca. 2 hours, including preparation and breaks. One 786 participant completed only two sessions, yielding a total number of sessions across 787 subjects of 47. Due to technical issues, for one session only data for the liberal condition was available. One participant was an author. All participants were included 788 789 in the signal-detection-theoretical and drift diffusion modeling analyses (Figure 2). One participant was excluded from the stimulus-related and the prestimulus alpha analysis 790 791 (Figures 3 and 4) due to excessive noise (EEG power spectrum opposite of 792 1/frequency). One further participant was excluded from the alpha-versus-gamma 793 power modulation (Figure 5) and gamma-versus-drift bias analyses (Figure 6) 794 because the liberal-conservative difference in gamma power in this participant was > 795 3 standard deviations away from the other participants. All participants had normal or 796 corrected-to-normal vision and were right handed. Participants provided written 797 informed consent before the start of the experiment. All procedures were approved by 798 the ethics committee of the University of Amsterdam.

799 Stimuli Stimuli consisted of a continuous semi-random rapid serial visual presentation 800 (rsvp) of full screen texture patterns. The texture patterns consisted of line elements 801 approx. 0.07° thick and 0.4° long in visual angle. Each texture in the rsvp was 802 presented for 40 ms (i.e. stimulation frequency 25 Hz), and was oriented in one of four 803 possible directions: 0°, 45°, 90° or 135°. Participants were instructed to fixate a red 804 dot in the center of the screen. At random inter trial intervals (ITI's) sampled from a 805 uniform distribution (ITI range 0.3–2.2 s), the rsvp contained a fixed sequence of 25 806 texture patterns, which in total lasted one second. This fixed sequence consisted of 807 four stimuli preceding a (non-)target stimulus (orientations of 45°, 90°, 0°, 90°

808 respectively) and twenty stimuli following the (non)-target (orientations of 0°, 90°, 0°, 809 90°, 0°, 45°, 0°, 135°, 90°, 45°, 0°, 135°, 0°, 45°, 90°, 45°, 90°, 135°, 0°, 135° 810 respectively) (Figure 2). The fifth texture pattern within the sequence (occurring from 811 0.16 s after sequence onset) was either a target or a nontarget stimulus. Nontargets 812 consisted of either a 45° or a 135° homogenous texture, whereas targets contained a 813 central orientation-defined square of 2.42° visual angle, thereby consisting of both a 814 45° and a 135° texture. 50% of all targets consisted of a 45° square and 50% of a 135° 815 square. Of all trials, 75% contained a target and 25% a nontarget. Target and 816 nontarget trials were presented in random order. To avoid specific influences on target 817 stimulus visibility due to presentation of similarly or orthogonally oriented texture patterns temporally close in the cascade, no 45° and 135° oriented stimuli were 818 819 presented directly before or after presentation of the target stimulus. All stimuli had an 820 isoluminance of 72.2 cd/m². Stimuli were created using MATLAB (The Mathworks, 821 Inc., Natick, MA, USA) and presented using Presentation (Neurobehavioral systems, 822 Inc., Albany, CA, USA).

Experimental design The participants' task was to detect targets and actively report them by pressing a button using their preferred hand. Targets occasionally went unreported, presumably due to constant forward and backward masking by the continuous cascade of stimuli and unpredictability of target timing (Fahrenfort, Scholte, & Lamme, 2007). The onset of the fixed order of texture patterns preceding and following (non-)target stimuli was neither signaled nor apparent.

829 At the beginning of the experiment, participants were informed they could earn 830 a total bonus of EUR 30, on top of their regular pay or course credit. In two separate 831 conditions within each session of testing, we encouraged participants to use either a 832 conservative or a liberal bias for reporting targets using both aversive sounds as well 833 as reducing their bonus after errors. In the conservative condition, participants were 834 instructed to only press the button when they were relatively sure they had seen the 835 target. The instruction on screen before block onset read as follows: "Try to detect as 836 many targets as possible. Only press when you are relatively sure you just saw a 837 target." To maximize effectiveness of this instruction, participants were told the bonus 838 would be diminished by ten cents after a false alarm. During the experiment, a loud 839 aversive sound was played after a false alarm to inform the participant about an error.

840 During the liberal condition, participants were instructed to miss as few targets as 841 possible. The instruction on screen before block onset read as follows: "Try to detect as many targets as possible. If you sometimes press when there was nothing this is 842 843 not so bad". In this condition, the loud aversive sound was played twice in close 844 succession whenever they failed to report a target, and three cents were subsequently 845 deducted from their bonus. The difference in auditory feedback between both 846 conditions was included to inform the participant about the type of error (miss or false 847 alarm), in order to facilitate the desired bias in both conditions. After every block, the 848 participant's score (number of missed targets in the liberal condition and number of 849 false alarms in the conservative condition) was displayed on the screen, as well as the 850 remainder of the bonus. After completing the last session of the experiment, every 851 participant was paid the full bonus as required by the ethical committee.

852 During a block, participants continuously monitored the screen and were free 853 to respond by button press whenever they thought they saw a target. Each block contained 240 trials, of which 180 target and 60 nontarget trials. Participants 854 performed six blocks per session. The task instruction was presented on the screen 855 856 before the block started. The condition of the first block of a session was 857 counterbalanced across participants. Prior to EEG recording in the first session, 858 participants performed a 10-minute practice run of both conditions, in which visual 859 feedback directly after a miss (liberal condition) or false alarm (conservative) informed 860 participants about their mistake, allowing them to adjust their decision bias 861 accordingly.

862 **Behavioral analysis** We calculated participants criterion *c* (Green & Swets, 1966) 863 across the trials in each condition as follows:

864
$$c = -\frac{1}{2} \left[Z(Hit-rate) + Z(FA-rate) \right]$$

where Z(...) is the inverse standard normal distribution. Furthermore, we calculated
objective sensitivity measure d' using:

867

868
$$d' = Z(Hit-rate) - Z(FA-rate)$$

as well as by subtracting hit and false alarm rates. Reaction times (RT's) were
measured as the period between target onset and button press.

872 Drift diffusion modeling of choice behavior We fitted the drift diffusion model to our 873 behavioural data for each subject individually, and separately for the liberal and 874 conservative conditions. We fitted the model using a G square method based on quantile RT's (RT cutoff, 200 ms, for details, see Ratcliff et al. (2016)), using a modified 875 876 version of the HDDM 0.6.0 package (Wiecki, Sofer, & Frank, 2013) (code will be made 877 available). The RT distributions for 'yes' responses were represented by the 0.1, 0.3, 878 0.5, 0.7 and 0.9 quantiles, and, along with the associated response proportions, 879 contributed to G square. In addition, a single bin containing the number of 'no' 880 responses contributed to G square. Fitting the model to RT distributions for 'yes' and 881 'no' choices (termed 'stimulus coding' in Wiecki et al. (2013)), as opposed to the more 882 common fits of correct and incorrect choice RT's (termed 'accuracy coding' in Wiecki 883 et al. (2013)), allowed us to estimate parameters that could have induced biases in 884 subjects' behavior.

885 Parameter recovery simulations showed that letting both the the starting point 886 of the accumulation process and drift bias (an evidence-independent constant added 887 to the drift toward one or the other bound) free to vary with experimental conditions is 888 problematic for data with no explicit "no" responses (data not shown). Thus, to test 889 whether shifts in drift bias or starting point underlied bias we fitted three separate 890 models. In the first model ('fixed model'), we allowed only the following parameters to 891 vary between the liberal and conservative condition: (i) the mean drift rate across trials; 892 (ii) the separation between both decision bounds (i.e., response caution); and (iii) the 893 non-decision time (sum of the latencies for sensory encoding and motor execution of 894 the choice). Additionaly, the bias parameters starting point and drift bias were fixed for 895 the experimental conditions. The second model ('starting point model') was the same 896 as the fixed model, except that we let the starting point of the accumulation process 897 vary with experimental condition, whereas the drift bias was kept fixed for both 898 conditions. The third model ('drift bias model') was the same as the fixed model, except 899 that we let the drift bias vary with experimental condition, while the starting point was 900 kept fixed for both conditions. We used Bayesian Information Criterion (BIC) to select 901 the model which provided the best fit to the data (Neath & Cavanaugh, 2012). The BIC

902 compares models based on their maximized log-likelihood value, while penalizing903 for the number of parameters.

904 **EEG recording** Continuous EEG data were recorded at 256 Hz using a 48-channel 905 BioSemi Active-Two system (Biosemi, Amsterdam, the Netherlands), connected to a 906 standard EEG cap according to the international 10-20 system. Electrooculography 907 (EOG) was recorded using two electrodes at the outer canthi of the left and right eves 908 and two electrodes placed above and below the right eve. Horizontal and vertical EOG 909 electrodes were referenced against each other, two for horizontal and two for vertical 910 eye movements (blinks). We used the Fieldtrip toolbox (Oostenveld, Fries, Maris, & 911 Schoffelen, 2011) and custom software in MATLAB R2016b (The Mathworks Inc., 912 Natick, MA, USA) to process the data (see below). Data were re-referenced to the 913 average voltage of two electrodes attached to the earlobes.

914 **Trial extraction and preprocessing** We extracted trials of variable duration from 1 s 915 before target sequence onset until 1.25 after button press for trials that included a 916 button press (hits and false alarms), and until 1.25 s after stimulus onset for trials 917 without a button press (misses and correct rejects). The following constraints were 918 used to classify (non-)targets as detected (hits and false alarms), while avoiding the 919 occurrence of button presses in close succession to target reports and button presses 920 occurring outside of trials: 1) A trial was marked as detected if a response occurred 921 within 0.84 s after target onset: 2) when the onset of the next target stimulus sequence 922 started before trial end, the trial was terminated at the next trial's onset; 3) when a 923 button press occurred in the 1.5 s before trial onset, the trial was extracted from 1.5 s 924 after this button press; 4) when a button press occurred between 0.5 s before until 0.2 925 s after sequence onset, the trial was discarded. See Kloosterman et al. (2015) and 926 Meindertsma et al. (2017) for similar trial extraction procedures. After trial extraction, 927 channel time courses were linearly detrended and the mean of every channel was 928 removed per trial.

Artifact rejection Trials containing muscle artifacts were rejected from further
 analysis using a standard semi-automatic preprocessing method in Fieldtrip. This
 procedure consists of bandpass-filtering the trials of a condition block in the 110–125
 Hz frequency range, which typically contains most of the muscle artifact activity,

933 followed by a Z-transformation. Trials exceeding a threshold Z-score were removed 934 completely from analysis. We used as the threshold the absolute value of the minimum 935 Z-score within the block, + 1. To remove eye blink artifacts from the time courses, the 936 EEG data from a complete session were transformed using independent component 937 analysis (ICA), and components (typically one or two of the 48) due to blinks was 938 removed from the data. In addition, to remove microsaccade-related artifacts we 939 included two virtual channels in the ICA based on channels Fp1 and Fp2, which 940 included transient spike potentials as identified using the algorithm from Hassler et al. 941 (2011). The two components loading high on these virtual electrodes (typically with a 942 frontal topography) were also removed. Blinks and eye movements were then semi-943 automatically detected from the horizontal and vertical EOG (frequency range 1–15 944 Hz; z-value cut-off 4 for vertical; 6 for horizontal) and trials containing eye artefacts 945 within 0.1 s around target onset were discarded. This step was done to remove trials 946 in which the target was not seen because the eyes were closed. Finally, trials 947 exceeding a threshold voltage range of 200 μ V were discarded. To attenuate volume 948 conduction effects and suppress any remaining microsaccade-related activity, the 949 scalp current density (SCD) was computed using the second-order derivative (the 950 surface Laplacian) of the EEG potential distribution (Perrin et al., 1989).

951 Spectral analysis of EEG power We used a sliding window Fourier transform ((Mitra & Pesaran, 1999); step size, 50 ms; window length, 400 ms; frequency resolution, 2.5 952 953 Hz) to calculate time-frequency representations (spectrograms) of the EEG power for 954 each electrode and each trial. We used a single Hann taper for the frequency range 955 of 3–35 Hz (spectral smoothing, 4.5 Hz, bin size, 1 Hz) and the multitaper technique 956 for the 36 – 100 Hz frequency range (spectral smoothing, 8 Hz; bin size, 2 Hz; five 957 tapers). See Kloosterman et al. (2015) and Meindertsma et al. (2017) for similar 958 settings.

959 Spectrograms were aligned to the onset of the stimulus sequence containing 960 the (non)target. Power modulations (denoted as M in Figure 3) during the trials were 961 quantified as the percentage of power change at a given time point and frequency bin, 962 relative to a baseline power value for each frequency bin. We used as a baseline the 963 mean EEG power in the interval 0.4 to 0 s before trial onset. If this interval was not 964 completely present in the trial due to preceding events (see Trial extraction), this

965 period was shortened accordingly. We subtracted the trial-specific baseline value from 966 each sample in the time course per frequency bin and divided by the mean baseline power across all trials within a session. For the analysis of raw prestimulus power 967 968 modulations no baseline correction was applied. We focused our analysis of EEG 969 power modulations around target onsets on those electrodes that processed the visual 970 stimulus. To this end, we averaged the power modulations or raw power across eleven 971 occipito-parietal electrodes that showed stimulus-induced responses in the gamma-972 band range (59–100 Hz). See Kloosterman et al. (2015) and Meindertsma et al. (2017) 973 for a similar procedure.

974 **Condition-related EEG power modulation** To test at which frequencies raw EEG 975 power differed for the liberal and conservative conditions, we averaged power 976 modulation from 0.8 s up to 0.2 s (i.e. up to half the window size used for spectral 977 analysis, to avoid contamination of post- with pre-stimulus activity (lemi et al., 2017)) 978 from trial onset. Then, we expressed the power at each frequency in units of percent 979 signal change with respect to the conservative condition and statistically tested 980 whether this signal differed from zero (Figure 4C) (see Statistical comparisons).

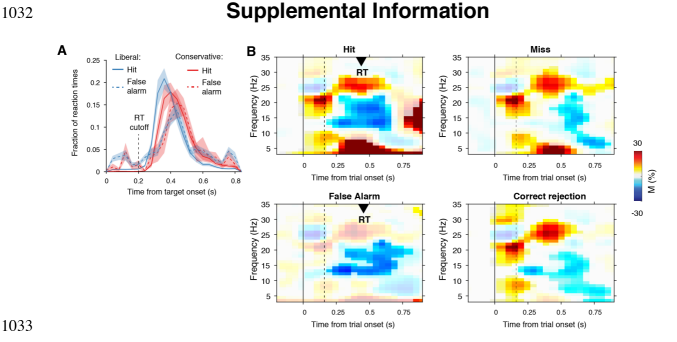
981 **Response gain model test** To test the prediction of increased gain during liberal of 982 the gain model, we first averaged activity in the 8–12 Hz range from 0.8 to 0.2 s before 983 trial onset (staying half our window size from trial onset, to avoid mixing pre- and post-984 stimulus activity, also see lemi et al. (2017)) and took the log transform, yielding a 985 single scalar alpha power value per trial expressing neural excitability. If this interval 986 was not completely present in the trial due to preceding events (see Trial extraction), 987 this period was shortened accordingly. Trials in which the scalar was > 3 standard 988 deviations away from the participant's mean were excluded. We then sorted all single 989 trials for each participant in ascending order of excitability and assigned them to ten 990 equally-spaced bins ranging from the lowest to the highest excitability scalars of that 991 participant. Adjacent bin ranges overlapped for 50% to stabilize estimates. Then we 992 averaged the corresponding log-transformed gamma modulation of these trials 993 (consisting of the average power within 59–100 Hz 0.2 to 0.6 s after trial onset) and 994 normalized each participants response by subtracting the minimum gamma power 995 during the conservative condition from all bins. Finally, we averaged across 996 participants and plotted the excitability bin number against the normalized gamma

power for each condition. See Rajagovindan and Ding (2011) for a similar procedure.
To statistically test the gain prediction, we employed a three-way repeated measures
ANOVA (see Statistical comparisons). For plotting purposes (Figure 5C), we
computed within-subject error bars by removing within each participant the mean
across conditions from the estimates.

1002 **Correlation between gamma power and drift bias** To link DDM drift bias and cortical 1003 gamma power, we re-fitted the DDM drift bias model while freeing the drift bias 1004 parameter within each condition for the ten neural excitability bins as determined by 1005 prestimulus alpha suppression (see section Response gain model test), while freeing 1006 the other parameters (drift rate, boundary separation, non-decision time) for each 1007 condition and fixing starting point across conditions. Then, we normalized the obtained 1008 scalars for gamma power and drift bias separately within participants using a Z-1009 transformation, and averaged across participants. Finally, we used within-subject 1010 group regression of the two measures across the ten bins for both conditions 1011 separately. In a control analysis, we conducted this regression after taking the liberal 1012 - conservative difference for each excitability bin before regressing, and obtained 1013 convergent results (Figure S4).

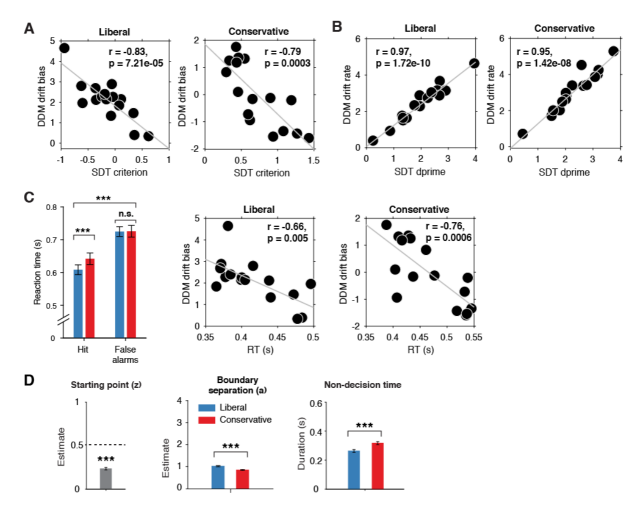
1014 **Statistical comparisons** We used two-sided permutation tests (10,000 permutations) 1015 (Efron & Tibshirani, 1998) to test the significance of behavioral effects and the model 1016 fits. To quantify power modulations after (non-)target onset, we tested the overall 1017 power modulation for significant deviations from zero. For these tests, we used a 1018 cluster-based permutation procedure to correct for multiple comparisons (Maris & 1019 Oostenveld, 2007). For time-frequency representations of power modulation, this 1020 procedure was conducted across all time-frequency bins. For frequency spectra, this 1021 procedure was performed across all frequency bins. To test whether there was 1022 evidence for increased gain in the liberal compared to the conservative condition, we 1023 conducted a three-way repeated measures ANOVA (condition (conservative, liberal) 1024 x brain activity type (prestimulus alpha, poststimulus gamma power) x bin level (1-1025 10)) using SPSS 23 (IBM, Inc.), inspecting linear and quadratic contrasts. As sphericity 1026 was violated in this model (p = 0.0001), we report both the uncorrected and 1027 Greenhouse-Geisser-corrected p-values. We used Pearson correlation to test the link 1028 between gamma power and drift bias. We tested the difference in correlation between

- 1029 the liberal and conservative conditions using the Fisher r-to-Z transformation and
- 1030 obtained the corresponding two-tailed p-value.



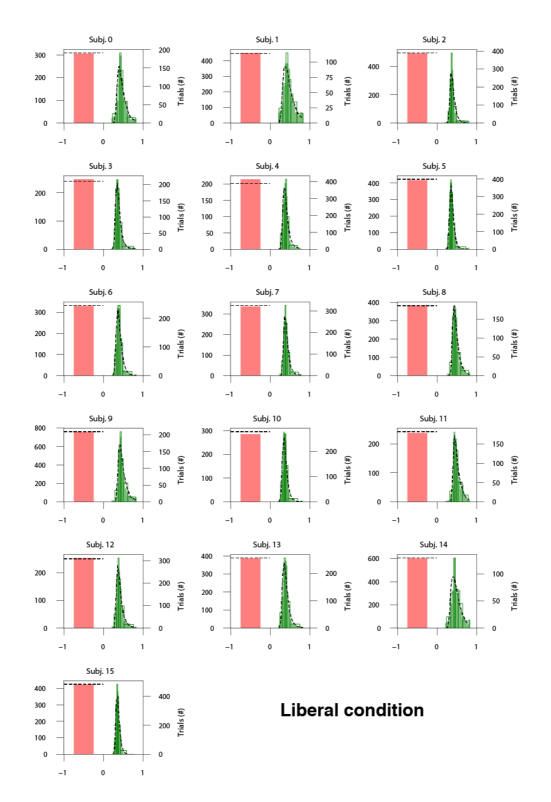
1034 Figure S1 I Behavioral and neurophysiological evidence that participants were sensitive to the 1035 implicit task structure. A. Participant-average RT distributions for hits and false alarms in both 1036 conditions. The presence of similar RT distributions for false alarms and hits indicates that participants 1037 were sensitive to trial onset despite the fact that trial onsets were only implicitly signalled. Error bars, 1038 SEM. B. Time-frequency representations of low-frequency EEG power modulations with respect to the 1039 prestimulus period (-0.4 - 0 s), pooled across the two conditions. Significant low-frequency modulation 1040 occurred even for nontarget trials without overt response (correct rejections), indicating that participants 1041 detected the onset of a trial even when neither a target was presented nor a response was given. 1042 Saturated colors indicate clusters of significant modulation, cluster threshold p < 0.05, two-sided 1043 permutation test across participants, cluster-corrected; N = 15). Solid and dotted vertical lines 1044 respectively indicate the onset of the trial and the target stimulus. M, power modulation.

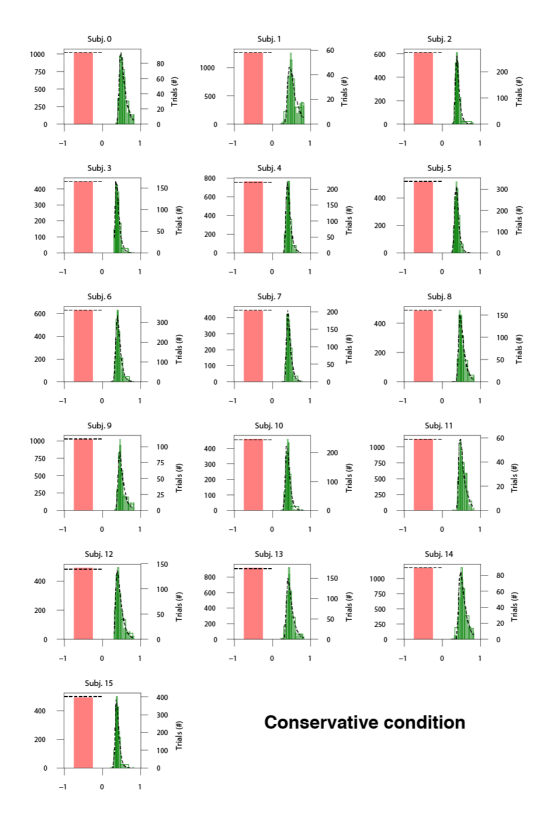
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1047 Figure S2 | Signal-detection-theoretic (SDT) behavioral measures during both conditions 1048 correspond closely to drift diffusion modeling (DDM) parameters. A. Across-participant Pearson 1049 correlation between criterion and DDM drift bias for the two conditions. The correlation is negative due 1050 to a lower criterion reflecting a stronger liberal bias. Each dot represents a participant. B. As A. but for 1051 correlation between dprime and drift rate. C. Left panel, mean reaction times (RT) for hits and false 1052 alarms for the two conditions. Middle and right panels, As A. but for correlation between RT and drift 1053 bias. **D.** Parameter estimates in the drift bias DDM not related to evidence accumulation (drift). ***p < 1054 0.001; n.s., not significant.



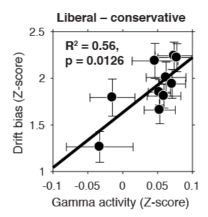


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1057 Figure S3 I Single-participant drift diffusion model fits for the drift bias model for both conditions

1058 Pink bars, number of "No" trials; Green bars, RT quantiles for "Yes" trials; dotted lines, model fits for

1059 the drift bias model.



- 1062 Figure S4 I Liberal conservative gamma activity predicts corresponding drift bias increase,
- 1063 showing that the experimental bias manipulation enhanced gamma activity. A. Linear regression
- 1064 of drift bias on gamma power across excitability bins for the liberal conservative contrast. Gamma
- 1065 and drift bias values were computed within participant within ten alpha suppression bins reflecting
- 1066 neural excitability, then Z-scored, and finally the conservative liberal difference across bins was taken.