Adaptive stimulus selection for multi-alternative psychometric functions with lapses

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(Dated: June 21, 2018)

Psychometric functions (PFs) quantify how external stimuli affect behavior and play an important role in building models of sensory and cognitive processes. Adaptive stimulus selection methods seek to select stimuli that are maximally informative about the PF given data observed so far in an experiment and thereby reduce the number of trials required to estimate the PF. Here we develop new adaptive stimulus selection methods for flexible PF models in tasks with two or more alternatives. We model the PF with a multinomial logistic regression mixture model that incorporates realistic aspects of psychophysical behavior, including lapses and multiple alternatives for the response. We propose an information-theoretic criterion for stimulus selection and develop computationally efficient methods for inference and stimulus selection based on semi-adaptive Markov Chain Monte Carlo (MCMC) sampling. We apply these methods to data from macaque monkeys performing a multi-alternative motion discrimination task, and show in simulated experiments that our method can achieve a substantial speed-up over random designs. These advances will reduce the data needed to build accurate models of multi-alternative PFs and can be extended to high-dimensional PFs that would be infeasible to characterize with standard methods.

Keywords: adaptive stimulus selection, sequential optimal design, Bayesian adaptive design, psychometric function, closed-loop experiments

Introduction

² Understanding the factors governing psychophysical behavior is ³ a central problem in neuroscience and psychology. Although ac-⁴ curate quantification of the behavior is an important goal in it-⁵ self, psychophysics provides an important tool for interrogating ⁶ the mechanisms governing sensory and cognitive processing in the ⁷ brain. As new technologies allow direct manipulations of neural ⁸ activity in the brain, there is a growing need for methods that can ⁹ characterize rapid changes in psychophysical behavior.

In a typical psychophysical experiment, an observer is trained to report judgements about a sensory stimulus by selecting a response from among two or more alternatives. The observer is assumed to have an internal probabilistic rule governing these decisions; this probabilistic map from stimulus to response is called the observer's

psychometric function. Because the psychometric function is not 15 directly observable, it must be inferred from multiple observations 16 of stimulus-response pairs. However, such experiments are costly 17 due to the large numbers of trials typically required to obtain good 18 estimates of psychometric functions. Therefore, a problem of ma-19 jor practical importance is to develop efficient experimental de-20 signs that can minimize the amount of data required to accurately 21 infer an observer's psychometric function. 22

Bayesian adaptive stimulus selection. A powerful approach for improving the efficiency of psychophysical experiments is to design the data collection process so that the stimulus is adaptively selected on each trial by maximizing a suitably defined objective function (MacKay, 1992). Such methods are known by a variety of names, including "active learning", "adaptive or sequential optimal experimental design", and "closed-loop experiments." 29

Bayesian approaches to adaptive stimulus selection define optimality of a stimulus in terms of its expected ability to improve the posterior distribution over the psychometric function, e.g., by reducing its variance or entropy. The three key ingredients of Bayesian adaptive stimulus selection method are (Chaloner & Verdinelli, 1995; Pillow & Park, 2016):

• **model** - parametrizes the psychometric function of interest;

• **prior** - captures initial beliefs about model parameters;

utility function - quantifies the usefulness of a hypothetical
 stimulus-response pair for improving the posterior.

Sequential algorithms for adaptive Bayesian experiments rely on 40 repeated application of three basic steps: (i) data collection (stim-41 ulus presentation and response measurement); (ii) inference (pos-42 terior updating using data from the most recent trial); and (iii) se-43 lection of an optimal stimulus for the next trial by maximizing ex-44 pected utility (see Fig. 1A). The inference step involves updating 45 the posterior distribution over the model parameters according to 46 Bayes rule with data from the most recent trial. Stimulus selection 47 involves calculating the expected utility (i.e., the expected improve-48 ment in the posterior) for a set of candidate stimuli, averaging over 49 the responses that might be elicited for each stimulus, and select-50 ing the stimulus for which the expected utility is highest. Example 51 utility functions include the negative trace of the posterior covari-52 ance (corresponding to the sum of the posterior variances for each 53 parameter) and the mutual information or information gain (which 54 corresponds to minimizing the entropy of the posterior). 55

Methods for Bayesian adaptive stimulus selection have been de-56 veloped over several decades in a variety of different disciplines. 57 If we focus on the specific application of estimating psychomet-58 ric functions, the field goes back to the QUEST (A. B. Watson & 59 Pelli, 1983) and ZEST (King-Smith, Grigsby, Vingrys, Benes, & 60 Supowit, 1994) algorithms, which were focused on the estimation 61 of discrimination thresholds, and to the simple case of 1-dimension 62 stimulus and binary responses (Treutwein, 1995). The Ψ method 63 (Kontsevich & Tyler, 1999) used Bayesian inference for estimat-64

ing both threshold and slope of a psychometric function, which 65 were extended to two-dimensional stimuli by Kujala and Lukka (2006). Further development of the method allowed for adap-67 tive estimation of more complex psychometric functions, where 68 the parameters were no longer limited to a threshold and a slope 69 (Barthelmé & Mamassian, 2008; Kujala & Lukka, 2006; Lesmes, 70 Lu, Baek, & Albright, 2010; Prins, 2013); and possibly related to 71 each other (Vul, Bergsma, & MacLeod, 2010). Models with multi-72 dimensional stimuli were also considered (DiMattina, 2015; Kujala 73 & Lukka, 2006; A. B. Watson, 2017). 74

Different models have been used to describe the psychometric 75 function. Standard choices include the logistic regression model 76 (Chaloner & Larntz, 1989; DiMattina, 2015; Zocchi & Atkinson, 77 1999), the Weibull distribution function (A. B. Watson & Pelli, 78 1983), and the cumulative function of Gaussian distribution (Kont-79 sevich & Tyler, 1999). More recent works also considered Gaussian Process models (Gardner, Song, Weinberger, Barbour, & Cun-81 ningham, 2015). Most of the previous works, however, were lim-82 ited to the case of binary responses. 83

In parallel, the development of Bayesian methods for inferring 84 psychometric functions (Kuss, Jäkel, & Wichmann, 2005; Prins, 85 2012; Wichmann & Hill, 2001a, 2001b) have enlarged the space of statistical models that could be employed to describepsychophys-87 ical phenomena based on (often limited) data. A variety of re-88 cent advances also arose in sensory neuroscience or neurophysiol-89 ogy, driven by the development of efficient inference techniques for neural encoding models (Lewi, Butera, & Paninski, 2009; M. Park, 91 Horwitz, & Pillow, 2011) or model comparison and discrimina-92 tion methods (Cavagnaro, Myung, Pitt, & Kujala, 2010; DiMattina 93 & Zhang, 2011; Kim, Pitt, Lu, Steyvers, & Myung, 2014). These 94 advances can in many cases be equally well applied to psychophys-95 ical experiments. 96

Our contributions. In this paper, we develop methods for adaptive stimulus selection in psychophysical experiments that are applicable to realistic models of human and animal psychophysical behavior. First, we describe a psychophysical model that incor-

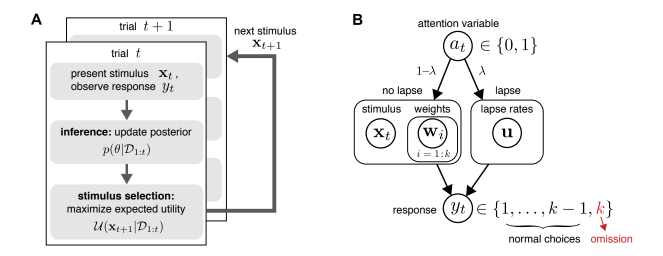


Figure 1: (A) Schematic of Bayesian adaptive stimulus selection. On each trial: (i) a stimulus is presented and response is observed; (ii) the posterior over the parameters θ is updated using all data collected so far in the experiment \mathcal{D}_t ; and (iii) the stimulus that maximizes the expected utility (in our case, information gain) is selected for the next trial. (B) A graphical model illustrating a hierarchical psychophysical observer model that incorporates lapses as well as the possibility of omissions. On each trial, a latent attention or lapse variable a_t is drawn from a Bernoulli distribution with parameter λ , to determine whether the observer attends to the stimulus \mathbf{x}_t on that trial or lapses. With probability $1 - \lambda$, and the observer attends to the stimulus $(a_t = 0)$, and the response y_t is drawn from a multinomial logistic regression model, where the probability of choosing option *i* is proportional to $\exp(\mathbf{w}_i^\top \mathbf{x}_t)$. With probability λ , the observer lapses ($a_t = 1$) and selects a choice from a (stimulus-independent) response distribution governed by parameter vector \mathbf{u} . So-called "omission" trials, in which the observer does not select one of the valid response options, are modeled with an additional response category $y_t = k$.

porates multiple response alternatives and "lapses", in which the 101 observer makes a response that does not depend on the stimulus. 102 This model can also incorporate "omission" trials, where the ob-103 server does not make a valid response (e.g., breaking fixation be-104 fore the go cue), by considering them as an additional response 105 category. Second, we describe efficient methods for updating the 106 posterior over the model parameters after every trial. Third, we in-107 troduce two algorithms for adaptive stimulus selection, one based 108 on a Gaussian approximation to the posterior and a second based 109 on Markov Chain Monte Carlo (MCMC) sampling. We apply these 110 algorithms to simulated data and to real data analyzed with simu-111 lated closed-loop experiments, and show that they can substantially 112 reduce in the number of trials required to estimate multi-alternative 113 psychophysical functions. 114

Psychophysical observer model

Here we describe a flexible model of psychometric functions (PFs) 116 based on the multinomial logistic (MNL) response model (Glonek 117 & McCullagh, 1995). We show how omission trials can be naturally incorporated into a model as one of the multiple responses 119 alternatives. We then develop a hierarchical extension of the model 120 that incorporates lapses (see Fig. 1B). 121

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Multinomial logistic response model. We consider the setting where the observer is presented with a stimulus $\mathbf{x} \in \mathbb{R}^d$ and selects a response $y \in \{1, \dots, k\}$ from one of k discrete choices on each trial. We will assume the stimulus is represented internally by some (possibly non-linear) feature vector $\phi(\mathbf{x})$, which we will write simply as ϕ for notational simplicity.

In the multinomial logistic model, the probability p_i of each possible outcome $i \in \{1, \dots, k\}$ is determined by the dot product

between the feature ϕ and a vector of weights \mathbf{w}_i according to:

$$p_i = \frac{\exp(\mathbf{w}_i^{\top} \boldsymbol{\phi})}{\sum_{j=1}^k \exp(\mathbf{w}_j^{\top} \boldsymbol{\phi})},$$
 (1)

where the denominator ensures that these probabilities sum to 1, 132 $\sum_{i=1}^{k} p_i = 1$. The function from stimulus to a probability vector 133 over choices, $\mathbf{x} \mapsto (p_1, \dots, p_k)$, is the psychometric function, and 134 the set of weights $\{\mathbf{w}_i\}_{i=1}^k$ are its parameters. Note that the model 135 is over-parameterized when written this way, since the requirement 136 that probabilities sum to 1 removes one degree of freedom from 137 the probability vector. Thus, we can without loss of generality fix 138 one of the weight vectors to zero, for example $\mathbf{w}_k = \mathbf{0}$, so that 139 the denominator in (eq. 1) becomes $z = 1 + \sum_{i=1}^{k} \exp(\mathbf{w}_{i}^{\top} \boldsymbol{\phi})$ and 140 $p_k = 1/z.$ 141

We consider the feature vector ϕ to be a known function of the 142 stimulus x, even when the dependence is not written explicitly. 143 For example, we can consider a simple form of feature embedding, 144 $\phi(\mathbf{x}) = [1, \mathbf{x}^{\top}]^{\top}$, corresponding to a linear function of the stim-145 ulus plus an offset. In this case, the weights for the *i*'th choice 146 would correpond to $\mathbf{w}_i = [b_i, \mathbf{a}_i^\top]^\top$, where b_i is the offset or bias 147 for the *i*'th choice, and \mathbf{a}_i are the linear weights governing sensi-148 tivity to x. The resulting choice probability has the familiar form, 149 $p_i \propto \exp(b_i + \mathbf{a}_i^{\top} \mathbf{x})$. Nonlinear stimulus dependencies can be 150 incorporated by including nonlinear functions of x in the feature 151 vector $\phi(\mathbf{x})$ (Knoblauch & Maloney, 2008; Murray, 2011; Neri & 152 Heeger, 2002). Dependencies on the trial history, such as the pre-153 vious stimulus or reward, may also be included as additional fea-154 tures in ϕ (see for example Bak, Choi, Akrami, Witten, and Pillow 155 (2016)).156

It is useful to always work with a normalized stimulus space, 157 in which the mean of each stimulus component x_{α} over the stim-158 ulus space is $\langle x_{\alpha} \rangle = 0$, and the standard deviation $\operatorname{std}(x_{\alpha}) = 1$. 159 This normalization ensures that the values of the weight parameters 160 are defined in more interpretable ways. The zero-mean condition 161 ensures that the bias b is the expectation value of log probability 162 over all possible stimuli. The unit-variance condition means that 163 the effect of moving a certain distance along one dimension of the 164

weight space is comparable to the moving the same distance in another dimension, again averaged over all possible stimuli. In other words, we are justified to use the same unit along all dimensions of the weight space.

Including omission trials. Even in binary tasks with only two 169 possible choices per trial, there is often an implicit third choice, 170 which is to make no response, make an illegal response, or to inter-171 rupt the trial at some point before the allowed response period. For 172 example, animals are often required to maintain an eye position 173 or a nose poke, or wait for a "go" cue before reporting a choice. 174 Trials on which the animal fails to obey these instructions are com-175 monly referred to as "omissions" or "violations", and are typically 176 discarded from analysis. However, failure to take these trials into 177 account may bias the estimates of the PF if they are more common 178 for some stimuli than others (see Fig. 2B). 179

The multinomial response model provides a natural framework 180 for incorporating omission trials because it accommodates an arbi-181 trary number of response categories. Thus we can model omissions 182 explicitly as one of the possible choices the observer can choose 183 from, or as the (k + 1)'st response category in addition to the k 184 valid responses. One could even consider different kinds of omis-185 sions separately, e.g., allowing choice k+1 to reflect fixation period 186 violations and choice k + 2 to reflect failure to report a choice dur-187 ing the response window. Henceforth, we will let k reflect the total 188 number of choices, including omission, as illustrated in Fig. 1B. 189

This formulation can also be useful for the rated Yes/No task in 190 human psychophysics, where a "Not Sure" response is explicitly 191 presented (C. S. Watson, Kellogg, Kawanishi, & Lucas, 1973). Al-192 though such model was considered for adaptive stimulus selection 193 (Lesmes et al., 2015), the third alternative was not handled as a 194 fully independent choice, as the goal was only to estimate the two 195 detection thresholds separately: one for a strict Yes, another for 196 a collapsed response of either Yes or Not Sure. Our model treats 197 each of the multiple alternatives equivalently. 198

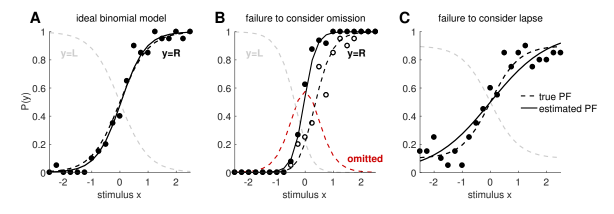


Figure 2: **Effects of omission and lapse.** Here we illustrate the undesirable effects of failing to take into account of omission and lapse. (A) If the PF follows an ideal binomial logistic model, it can be estimated very well from data. The black dashed line shows the true PF for one of the two responses (say y = R), and the gray dashed line shows the true PF for the other response (say y = L), such that the two dashed curves always add up to 1. The black dots indicate the mean probability to observe this response y = R at each stimulus point x. We drew 20 observations per stimulus point, at each of the 21 stimulus points along the 1-dimensional axis. The resulting estimate for P(y = 1) is shown in the solid black line. The inference method is not important for the current purpose, but we used the MAP estimate, discussed in a later section. (B) Now suppose that some trials fell into the implicit third choice which is the omission (red dashed line shows omission probability). The observed probability of y = R at each stimulus point (open black circles) follows the true PF (black dashed line). But if the omitted trials are systematically excluded from analysis, as in common practice, the estimated PF (solid black line) reflects a biased set of observations (filled black circles), and fail to recover the true PF. (C) When there is a finite lapse rate (we used a total lapse of $\lambda = 0.2$, uniformly distributed to the two outcomes), the true PF (dashed black line) asymptotes to a finite offset from 0 or 1. If the resulting observations (black dots) are fitted to a plain binomial model without lapse, the slope of the estimated PF (solid black line) is systematically biased.

Modeling lapse with a mixture model. Another important 199 feature of real psychophysical observers is the tendency to occa-200 sionally make errors that are independent of the stimulus. Such 201 choices, commonly known as "lapses" or "button press errors", 202 may reflect lapses in concentration or memory of the response 203 mapping (Kuss et al., 2005; Wichmann & Hill, 2001a). Lapses 204 are most easily identified by errors on "easy" trials, that is, trials 205 that should be performed perfectly if the observer were paying at-206 tention. 207

Although lapse rates can be negligible in highly trained ob-208 servers (Carandini & Churchland, 2013), they can be substantially 209 greater than zero in settings involving non-primates or complicated 210 psychophysical tasks. Lapses affect the psychometric function by 211 causing it to saturate above 0 and below 1, so that "perfect" per-212 formance is never achieved even for the easiest trials. Failure to 213 incorporate lapses into the PF model may therefore bias estimates 214 of sensitivity, as quantified by PF slope or threshold (illustrated 215

in Fig. 2C; also see Wichmann and Hill (2001a, 2001b) or Prins (2012)).

To model lapses, we use a mixture model that treats the observer's choice on each trial as coming from one of two probability distributions: a stimulus-dependent distribution (governed by the multinomial logistic model) and stimulus-independent distribution (reflecting a fixed probability of choosing any option when lapsing). Simpler versions of such mixture model have been proposed previously (Kuss et al., 2005).

Fig. 1B shows a schematic of the resulting model. On each trial, 225 a Bernoulli random variable $a \sim Ber(\lambda)$ governs whether the ob-226 server lapses: with probability λ and the observer lapses (i.e., ig-227 nores the stimulus), and with probability $1 - \lambda$, and the observer at-228 tends to the stimulus. If the observer lapses (a = 1), the response is 229 drawn according to fixed probability distribution (c_1, \ldots, c_k) gov-230 erning the probability of selecting options 1 to k, where $\sum c_i = 1$. 231 If the observer does not lapse (a = 0), the observer selects a re-232

sponse according to the multinomial logistic model. Under this model, the conditional probability of choosing option i given the stimulus can be written:

$$p_i = (1 - \lambda)q_i + \lambda c_i, \qquad q_i = \frac{\exp(\mathbf{w}_i^{\top} \phi)}{\sum_j \exp(\mathbf{w}_j^{\top} \phi)}$$
(2)

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where q_i is the lapse-free probability probability under the classical MNL model (eq. 1).

It is convenient to re-parameterize this model so that λc_i , the conditional probability of choosing the *i*-th option due to a lapse, is written

$$\lambda c_i = \frac{\exp(u_i)}{1 + \sum_j \exp(u_j)},\tag{3}$$

where each auxiliary lapse parameter u_i is proportional to the log probability of choosing option *i* due to lapse. The lapse-conditional probabilities of each choice, c_i , and the total lapse probability, λ , are respectively

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$$c_i = \frac{\exp(u_i)}{\sum_j \exp(u_j)}, \qquad \lambda = \sum_i \frac{\exp(u_i)}{1 + \sum_j \exp(u_j)}.$$
 (4)

Because each u_i lives on the entire real line, fitting can be carried 248 out with unconstrained optimization methods, although adding rea-249 sonable constraints may improve performance in some cases. The 250 full parameter vector of the resulting model is $\boldsymbol{\theta} = [\mathbf{w}^{\top}, \mathbf{u}^{\top}]^{\top}$, 251 which includes k additional lapse parameters $\mathbf{u} = \{u_1, \cdots, u_k\}$. 252 Note that in some cases it might be desirable to assume lapse 253 choices obey a uniform distribution, where the probability of each 254 option is $c_i = 1/k$. For this simplified "uniform-lapse" model we 255 need only a single lapse parameter u. Note that we have unified 256 the parameterizations of the "lapse rate" (deviation of the upper 257 asymptote of the PF from 1; in this case $\lambda - \lambda c_i$) and the "guess 258 rate" (deviation of the lower asymptote from 0; in this case λc_i), 259 which are often modeled separately in previous works with two-260 alternatives responses (Schütt, Harmeling, Macke, & Wichmann, 261 2016; Wichmann & Hill, 2001a, 2001b). Here they are written in 262 terms of a single family of parameters $\{u_i\}$, and extended naturally 263 to multi-alternative responses. 264

Our model provides a general and practical parametrization of psychometric functions with lapses. Although previous work has considered the problem of modeling lapses in psychophysical data, 267 much of it assumed a uniform-lapse model, where all options are 268 equally likely during lapses. Earlier approaches have often as-269 sumed either that the lapse probability was known a priori (Kontse-270 vich & Tyler, 1999), or was fit by a grid search over a small set of 271 candidate values (Wichmann & Hill, 2001a). Here we instead in-272 fer individual lapse probabilities for each response option, similar 273 to recent approaches described in Kuss et al. (2005); Prins (2012, 274 2013); Schütt et al. (2016). Importantly, our method infers the full 275 parameter θ that includes both the weight and the lapse parameters, 276 rather than treating the lapse separately. In particular, our parame-277 terization (eq. 3) has the advantage that there is no need to constrain 278 the support of the lapse parameters u_i . These parameters' relation-279 ship to lapse probabilities c_i takes the same ("softmax") functional 280 form as the multinomial logistic model, placing both sets of param-281 eters on an equal footing. 282

Before closing this section, we would like to reflect briefly on 283 the key differences between omissions and lapses. First, although 284 omissions and lapses both reflect errors in decision making, omis-285 sions are defined as invalid responses and are thus easily identi-286 fiable from the data; lapses, on the other hand, are indistinguish-287 able from normal responses, and are identifiable only from the fact 288 that the psychometric function does not saturate at 0 or 1. Second, 289 modeling omissions as a response category under the multinomial 290 logistic model means that the probability of omission is stimulus-291 dependent (e.g., more likely to arise on trials with high difficulty, or 292 generally when the evidence for other options is low). Even if the 293 omissions are not stimulus-dependent, and governed entirely by a 294 "bias" parameter, the probability of omission will still be higher 295 when the evidence for other choices is low, or lower when the ev-296 idence for other choices is high. Omissions that arise in a purely 297 stimulus-independent fashion, on the other hand, will be modeled 298 as arising from the lapse parameter associated with the omission 299 response category. Omissions can thus arise in two ways under the 300 model: as categories selected under the multinomial model or as 301 lapses arising independent of the stimulus and other covariates. 302

Posterior inference

Bayesian methods for adaptive stimulus selection require the pos-304 terior distribution over model parameters given the data observed 305 so far in an experiment. The posterior distribution results from the 306 combination of two ingredients: a prior distribution $p(\theta)$, which 307 captures prior uncertainty about the model parameters θ , and a 308 likelihood function $p(\{y_s\}|\{\mathbf{x}_s\}, \boldsymbol{\theta})$, which captures information 309 about the parameters from the data $\{(\mathbf{x}_s, y_s)\}, s = 1, \dots, t, \text{ con-}$ 310 sisting of stimulus-response pairs observed up to the current time 311 bin t. 312

Unfortunately, the posterior distribution for our model has no analytic form. We therefore describe two methods for approximate posterior inference: one relying on a Gaussian approximation to the posterior, known as the Laplace approximation, and a second one based on MCMC sampling.

Prior. The prior distribution specifies our beliefs about model 318 parameters before we have collected any data, and serves to reg-319 ularize estimates obtained from small amounts of data, e.g., by 320 shrinking estimated weights toward zero. Typically we want the 321 prior to be weak enough that the likelihood dominates the posterior 322 for reasonable-sized datasets. However, the choice of prior is es-323 pecially important in adaptive stimulus selection settings because 324 it determines the effective volume of the search space (M. Park 325 & Pillow, 2012; M. Park, Weller, Horwitz, & Pillow, 2014). For 326 example, if the weights are known to exhibit smoothness, then a 327 correlated or smoothness-inducing prior can improve the perfor-328 mance of adaptive stimulus selection because the effective size (or 329 entropy) of the parameter space is much smaller than under an in-330 dependent prior (M. Park & Pillow, 2012). 331

In this study, we use a generic independent, zero-mean Gaussian prior over the weight vectors

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$$p(\mathbf{w}_i) = \mathcal{N}(\mathbf{0}, \sigma^2 I), \tag{5}$$

for all $i \in (1, ..., k)$, with a fixed standard deviation σ . This choice of prior is appropriate when the regressors $\{x\}$ are standardized,

since any single weight can take values that allow for a range of psychometric function shapes along that axis, from flat (w = 0) to 338 steeply decreasing $(w = -2\sigma)$ or increasing $(w = +2\sigma)$. We 339 used $\sigma = 3$ in the simulated experiments in Results. For the 340 lapse parameters $\{u_i\}$, we used a uniform prior over the range 341 $[\log(0.001), 0]$ with the natural log, so that each lapse probabil-342 ity λc_i is bounded between 0.001 and 1/2. We set the lower range 343 constraint below 1/N, where N = 100 is the number of observed 344 trials in our simulations, since we cannot reasonably infer lapse 345 probabilities with precision finer than 1/N. The upper range con-346 straint gives maximal lapse probabilities of 1/(k+1) if all u_i take 347 on the maximal value of 0. Note that our prior is uniform with 348 respect to the rescaled lapse parameters $\{u_i\}$, rather than to the ac-349 tual lapse rates; projected to the space of the lapse probabilities, 350 given the bounds, the prior increases towards smaller lapse. For a 351 comprehensive study of the effect of different priors on lapse, see 352 for example Schütt et al. (2016). 353

Psychometric function likelihood. The likelihood is the con-354 ditional probability of the data as a function of the model param-355 eters. Although we have thus far considered the response variable 356 y to be a scalar taking values in the set $\{1, \ldots, k\}$, it is more con-357 venient to use a "one-hot" or "1-of-k" representation, in which the 358 response variable y for each trial is a length-k vector with one 1 359 and (k-1) zeros; the position of the 1 in this vector indicates the 360 category selected. For example, in a task with four possible options 361 per trial, a response vector $\mathbf{y} = \begin{bmatrix} 0 & 0 & 1 & 0 \end{bmatrix}$ indicates a trial on which 362 the observer selected the third option. 363

With this parametrization, the log-likelihood function for a single trial can be written 365

$$\log p(\mathbf{y}|\mathbf{x}, \boldsymbol{\theta}) = \sum_{i} y_i \log p_i(\mathbf{x}, \boldsymbol{\theta}) = \mathbf{y}^\top \log \mathbf{p}(\mathbf{x}, \boldsymbol{\theta}), \quad (6) \quad {}_{366}$$

where $p_i(\mathbf{x}, \boldsymbol{\theta})$ denotes the probability $p(y_i = 1 | \mathbf{x}, \boldsymbol{\theta})$ under the model (eq. 1), and $\mathbf{p}(\mathbf{x}, \boldsymbol{\theta}) \equiv [p_1(\mathbf{x}, \boldsymbol{\theta}), \dots, p_k(\mathbf{x}, \boldsymbol{\theta})]^{\top}$ denotes the vector of probabilities for a single trial.

In the classical (lapse-free) multinomial logistic model, where ${}_{370} \theta = {\mathbf{w}_i}$, the log likelihood is a concave function of θ , which ${}_{371}$

guarantees that numerical optimization of the log-likelihood will
find a global optimum. With a finite lapse rate, however, the log
likelihood is no longer concave. (See Appendix A).

Posterior distribution. The log-posterior can be written as the
sum of log-prior and log-likelihood summed over trials, plus a constant:

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$$\log p(\boldsymbol{\theta}|\mathcal{D}_t) = \log p(\boldsymbol{\theta}) + \sum_{s=1}^t \log p(\mathbf{y}_s|\mathbf{x}_s, \boldsymbol{\theta}) + c, \quad (7)$$

where $\mathcal{D}_t \equiv {\{\mathbf{x}_s, y_s\}_{s=1}^t}$ denotes the accumulated data up to trial t and $c = -\log \left(\int p(\theta) \prod_s p(\mathbf{y}_s | \mathbf{x}_s) d\theta\right)$ is a normalization constant that does not depend on the parameters θ . Because this constant has no tractable analytic form, we rely on two alternate methods for obtaining a normalized posterior distribution.

Inference via Laplace approximation. The Laplace approximation is a well-known Gaussian approximation to the posterior
distribution, which can be derived from a second-order Tayler series approximation to the log-posterior around its mode (Bishop,
2006).

³⁸⁹ Computing the Laplace approximation involves a two-step pro-³⁹⁰ cedure. The first step is to perform a numerical optimization of ³⁹¹ $\log p(\theta | D_t)$ to find the posterior mode, or maximum a posteriori ³⁹² (MAP) estimate of θ . This vector, given by

$$\hat{\boldsymbol{\theta}}_{t} = \operatorname*{argmax}_{\boldsymbol{\theta}} \log p(\boldsymbol{\theta}) + \sum_{s=1}^{t} \log p(\mathbf{y}_{s} | \mathbf{x}_{s}, \boldsymbol{\theta}), \tag{8}$$

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provides the mean of the Laplace approximation. Because we can
explicitly provide the gradient and Hessian of the log likelihood
(see Appendix A) and log-prior, this optimization can be carried
efficiently via Newton-Raphson or trust region methods.

The second step is to compute the second derivative (the Hessian matrix) of the log-posterior at the mode, which provides the inverse covariance of the Gaussian. This gives us a local Gaussian approximation of the posterior, centered at the posterior mode:

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$$p(\boldsymbol{\theta}|\mathcal{D}_t) \approx \mathcal{N}(\hat{\boldsymbol{\theta}}_t, C_t),$$
 (9)

where covariance $C_t = -H_t^{-1}$ is the inverse Hessian of the log posterior, $H_t(i, j) = \partial^2 (\log p(\theta | \mathcal{D}_t) / (\partial \theta_i \partial \theta_j))$, evaluated at $\hat{\theta}_t$.

Note that when the log-posterior is concave (i.e., when the 405 model does not contain lapse), numerical optimization is guaran-406 teed to find a global maximum of the posterior. Log-concavity 407 also strengthens the rationale for using the Laplace approximation, 408 since the true and approximate posterior are both log-concave den-409 sities centered on the true mode (Paninski et al., 2010; Pillow, Ah-410 madian, & Paninski, 2011). When the model incorporates lapses, 411 these guarantees no longer apply globally. 412

Inference via MCMC sampling. A second approach to in-413 ference is to generate samples from the posterior distribution over 414 the parameters via Markov Chain Monte Carlo (MCMC) sampling. 415 Sampling-based methods are typically more computationally in-416 tensive than the Laplace approximation, but may be warranted 417 when the posterior is not provably log-concave (as is the case when 418 lapse rates are non-zero) and therefore not well approximated by a 419 single Gaussian. 420

The basic idea in MCMC sampling is to set up an easy-to-sample 421 Markov Chain that has the posterior as its stationary distribution. 422 Sampling from this chain produces a dependent sequence of posterior samples: $\{\boldsymbol{\theta}_m\} \sim p(\boldsymbol{\theta}|\mathcal{D}_t)$, which can be used to evaluate 424 posterior expectations via Monte Carlo integrals: 425

$$\mathbb{E}[f(\boldsymbol{\theta})] \approx \frac{1}{M} \sum_{m=1}^{M} f(\boldsymbol{\theta}_m), \qquad (10) \quad {}_{426}$$

for any function $f(\theta)$. The mean of the posterior is obtained from 427 setting $f(\theta) = \theta$, although for adaptive stimulus selection we will 428 be interested in the full shape of the posterior. 429

The Metropolis-Hastings (MH) algorithm is perhaps the sim-430 plest and most widely-used MCMC sampling method (Metropo-431 lis, Rosenbluth, Rosenbluth, Teller, & Teller, 1953). It generates 432 samples via a proposal distribution centered on the current sample 433 (see Appendix B). The choice of proposal distribution is critical to 434 the efficiency of the MH algorithm, since this governs the rate of 435 "mixing", or the the number of Markov Chain samples required to 436 obtain independent samples from the posterior distribution (Rosen-437 thal, 2011). Faster mixing implies that fewer samples M are re-438 quired to obtain an accurate approximation to the posterior. 439

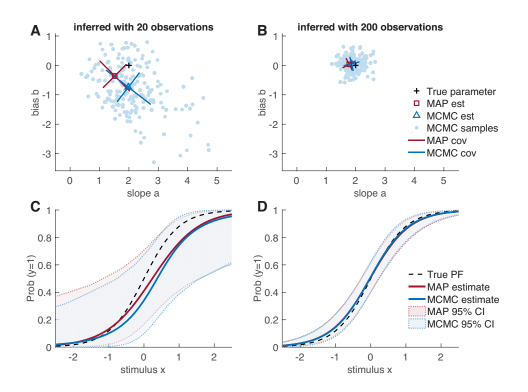


Figure 3: Inferring the psychometric function. Example of a psychometric problem, with a lapse-free binomial logistic model $f(v) = e^v/(1 + e^v)$. Given a 1D stimulus, a response were drawn from a "true" model P(y = 1) = f(b + ax) with two parameters, slope a = 2 and bias b = 0. (A-B) Viewing on the parameter space, the posterior distributions become sharper (and closer to the true parameter values) as the dataset size N increases. Shown at a small (A) N = 20 and a large (B) N = 200. For the MAP estimate, the mode of the distribution is marked with a square, and the two standard deviations ("widths") of its Gaussian approximation are shown with bars. For the MCMC sampling method, all M = 500 samples of the chain are shown in dots, the sample mean with a triangle, and the widths with the bars. The widths are the standard deviations along the principal directions of the sampled posterior (eigenvectors of the covariance matrix; not necessary aligned with the a - b axes). (C-D) The accuracy of the estimated PF improves with the number of observations N, using either of the two posterior inference methods (MAP-based and sampling-based). Shown at a small (C) N = 20 and a large (D) N = 200. The two methods are highly consistent in this simple case, especially when N is large enough.

Here we propose a semi-adaptive MH algorithm, developed 440 specifically for the current context of sequential learning. Our 441 approach is based on an established observation that the optimal 442 width of the proposal distribution should be proportional to the 443 typical length scale of the distribution being sampled (Gelman, 444 Roberts, & Gilks, 1996; Roberts, Gelman, & Gilks, 1997). Our al-445 gorithm is motivated by the adaptive Metropolis algorithm (Haario, 446 Saksman, & Tamminen, 2001), where the proposal distribution is 447 updated at each proposal within a single chain; here we do not 448 adapt the proposal within chains, but rather after each trial. Specif-449

ically, we set the covariance of a Gaussian proposal distribution to
be proportional to the covariance of the samples from the previous
trial, using the scaling factor of Haario et al. (2001). See Appendix
B for details. The adaptive algorithm takes advantage of the fact
that the posterior cannot change too much between trials, since it
changes only by a single-trial likelihood term on each trial.

Adaptive stimulus selection methods

As data are collected during the experiment, the posterior distribu-457 tion becomes narrower due to the fact that each trial carries some 458 additional information about the model parameters (see Fig. 3). 459 This narrowing of the posterior is directly related to information 460 gain. A stimulus that produces no expected narrowing of the pos-461 terior is, by definition, uninformative about the parameters. On the 462 other hand, a stimulus that (on average) produces a large change 463 in the current posterior is an informative stimulus. Selecting infor-464 mative stimuli will reduce the number of stimuli required to obtain 465 narrow posterior, which is the essence of adaptive stimulus se-466 lection methods. In this section, we introduce a precise measure 467 of information gain between a stimulus and the model parameters, 468 and propose an algorithm for selecting stimuli to maximize it. 469

⁴⁷⁰ Infomax criterion for stimulus selection. At each trial, we ⁴⁷¹ present a stimulus x and observe the outcome y. After t trials, the ⁴⁷² expected gain in information from a stimulus x is equal to the mu-⁴⁷³ tual information between y and the model parameters θ , given the ⁴⁷⁴ data D_t observed so far in the experiment. We denote this condi-⁴⁷⁵ tional mutual information:

$$I_{t}(\boldsymbol{\theta}; \mathbf{y} | \mathbf{x}) = \int \int d\boldsymbol{\theta} \, d\mathbf{y} \, p(\boldsymbol{\theta}, \mathbf{y} | \mathbf{x}, \mathcal{D}_{t}) \log \frac{p(\boldsymbol{\theta}, \mathbf{y} | \mathbf{x}, \mathcal{D}_{t})}{p(\boldsymbol{\theta} | \mathcal{D}_{t}) p(\mathbf{y} | \mathbf{x}, \mathcal{D}_{t})}, \quad (11)$$

where $p(\theta, \mathbf{y} | \mathbf{x}, \mathcal{D}_t)$ is the joint distribution of θ and \mathbf{y} given a stimulus \mathbf{x} and dataset \mathcal{D}_t , the term $p(\theta | \mathcal{D}_t)$ is the current posterior distribution over the parameters from previous trials, and $p(\mathbf{y} | \mathbf{x}, \mathcal{D}_t) = \int d\theta \, p(\mathbf{y} | \mathbf{x}, \theta) p(\theta | \mathcal{D}_t)$ is known as the posteriorpredictive distribution of \mathbf{y} given \mathbf{x} .

It is useful to note that the mutual information can equivalently be written in two other ways involving Shannon entropy. The first is given by:

488
$$I_t(\boldsymbol{\theta}; \mathbf{y} | \mathbf{x}) = H_t(\boldsymbol{\theta}) - H_t(\boldsymbol{\theta} | \mathbf{y}; \mathbf{x})$$
(12)

where the first term is the entropy of the posterior at time t,

490
$$H_t(\boldsymbol{\theta}) = -\int d\boldsymbol{\theta} \, p(\boldsymbol{\theta}|\mathcal{D}_t) \log p(\boldsymbol{\theta}|\mathcal{D}_t), \quad (13)$$

and the second is the conditional entropy of θ given y,

$$H_t(\boldsymbol{\theta}|\mathbf{y}; \mathbf{x}) = -\mathbb{E}_{\boldsymbol{\theta}, \mathbf{y}} \Big[\log p(\boldsymbol{\theta}|\mathbf{y}, \mathbf{x}, \mathcal{D}_t) \Big]$$

$$= -\int \int d\boldsymbol{\theta} \, d\mathbf{y} \, p(\boldsymbol{\theta}, \mathbf{y}|\mathbf{x}, \mathcal{D}_t) \log p(\boldsymbol{\theta}|\mathbf{y}, \mathbf{x}, \mathcal{D}_t) \quad (14) \quad (14)$$

$$= -\iint d\boldsymbol{\theta} \, d\mathbf{y} \, p(\boldsymbol{\theta}, \mathbf{y} | \mathbf{x}, \mathcal{D}_t) \log p(\boldsymbol{\theta} | \mathbf{y}, \mathbf{x}, \mathcal{D}_t), \quad (14) \quad {}_{494}$$

492

which is the entropy of the updated posterior *after* having observed 496 x and y, averaged over draws of y from the posterior predictive 497 distribution. Written this way, the mutual information can be seen 498 as the expected reduction in posterior entropy from a new stimulusresponse pair. Moreover, the first term, $H_t(\theta)$, is independent of 500 the stimulus and response on the current trial, so infomax stimulus 501 selection is equivalent to picking the stimulus that minimizes the 502 expected posterior entropy $H_t(\theta|\mathbf{y}; \mathbf{x})$. 503

A second equivalent expression for the mutual information, 504 which will prove useful for our sampling-based method, is: 505

$$I_t(\boldsymbol{\theta}; \mathbf{y} | \mathbf{x}) = H_t(\mathbf{y}; \mathbf{x}) - H_t(\mathbf{y} | \boldsymbol{\theta}; \mathbf{x}),$$
 (15) 500

which is the difference between the marginal entropy of the response distribution conditioned on \mathbf{x} , 508

$$H_t(\mathbf{y}; \mathbf{x}) = -\int d\mathbf{y} \, p(\mathbf{y} | \mathbf{x}, \mathcal{D}_t) \log p(\mathbf{y} | \mathbf{x}, \mathcal{D}_t) \qquad (16) \quad {}_{509}$$

and the conditional entropy of the response y given θ , conditioned on the stimulus:

$$H_t(\mathbf{y}|\boldsymbol{\theta}; \mathbf{x}) = -\iint d\mathbf{y} \, d\boldsymbol{\theta} \, p(\boldsymbol{\theta}, \mathbf{y}|\mathbf{x}, \mathcal{D}_t) \log p(\mathbf{y}|\mathbf{x}, \boldsymbol{\theta}). \quad (17) \quad {}_{512}$$

This formulation shows the mutual information to be equal to the difference between the entropy of the marginal distribution of y conditioned on x (with θ integrated out) and the average entropy of y given x and θ , averaged over the posterior distribution of θ . The dual expansion of the mutual information was also used by Kujala and Lukka (2006).

In a sequential setting where t is the latest trial and t + 1 is the upcoming one, the optimal stimulus is the information-maximizing ("infomax") solution: 521

$$\mathbf{x}_{t+1} = \arg\max_{\mathbf{x}} I_t(\boldsymbol{\theta}; \mathbf{y} | \mathbf{x}). \tag{18}$$

Fig. 4 shows an example of a simulated experiment where the stimulus was selected adaptively following the infomax criterion. Note

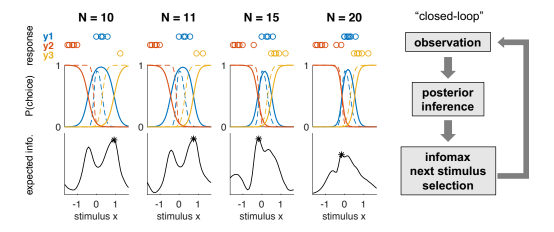


Figure 4: Example of infomax adaptive stimulus selection, simulated with a three-alternatives lapse-free model on 1D stimulus. The figure shows how given a small set of data (the stimulus-response pairs shown in top row), the PFs are estimated based on the accumulated data (middle row), and the next stimulus is chosen to maximize the expected information gain (bottom row). Each column shows the instance after the *N* observations in a single adaptive stimulus selection sequence, for N = 10, 11, 15 and 20 respectively. In the middle row, the estimated PFs (solid lines) quickly approach the true PFs (dashed lines) through the adaptive and optimal selection of stimuli. This example was generated using the Laplace approximation based algorithm, with an independent Gaussian prior over the weights with mean zero and standard deviation $\sigma = 10$.

that our algorithm takes a "greedy" approach of optimizing one trial at a time. For work on optimizing beyond the next trial, see for example Kim, Pitt, Lu, and Myung (2017).

Selecting the optimal stimulus thus requires maximizing the mu-528 tual information over the set of all possible stimuli $\{x\}$. Since each 529 evaluation of the mutual information involves a high-dimensional 530 integral over parameter space and response space, this is a highly 531 computationally demanding task. In the next sections, we present 532 two algorithms for efficient infomax stimulus selection based on 533 each of the two approximate inference methods described previ-534 ously. 535

Infomax with Laplace approximation. Calculation of the mutual information is greatly simplified by a Gaussian approximation of the posterior. The entropy of a Gaussian distribution with covariance C is equal to $\frac{1}{2} \log |C|$ up to a constant factor. If we expand the mutual information as in (eq. 12), and recall that we need only minimize the expected posterior entropy after observing the response, the optimal stimulus for time-step t + 1 is given by:

543
$$\mathbf{x}_{t+1}^* = \underset{\mathbf{x}}{\operatorname{argmin}} \int d\mathbf{y} \, p(\mathbf{y}|\mathbf{x}, \mathcal{D}_t) \log |\tilde{C}(\mathbf{x}, \mathbf{y})|, \quad (19)$$

where $\tilde{C}(\mathbf{x}, \mathbf{y})$ is the covariance of the updated (Gaussian) poste-544 rior after observing stimulus-response pair (x, y). To evaluate the 545 updated covariance $\tilde{C}(\mathbf{x}, \mathbf{y})$ under the Laplace approximation, we 546 would need to numerically optimize the posterior for θ for each 547 possible resonse y, for any candidate stimulus x, which would be 548 computationally infeasible. We therefore use a fast approximate 549 method for obtaining a closed-form update for $\tilde{C}(\mathbf{x}, \mathbf{y})$ from the 550 current posterior covariance C_t , following an approach developed 551 in Lewi et al. (2009). See Appendix C for details. Note that this 552 approximate sequential update is only used for calculating the ex-553 pected utility of each candidate stimulus by approximating the pos-554 terior distribution at the next trial. For obtaining the MAP estimate 555 of the current model parameter, θ_t , numerical optimization needs 556 to be performed using the full accumulated data \mathcal{D}_t each time. 557

Once we have $\log |\tilde{C}(\mathbf{x}, \mathbf{y})|$ for each given stimulus-observation pair, we numerically sum this over a set of discrete counts \mathbf{y} that are likely under the posterior-predictive distribution. This is done

in two steps, by separating the integral in (eq. 19) as:

562
$$\int d\mathbf{y} \, p(\mathbf{y}|\mathbf{x}, \mathcal{D}_t) \, \log |\tilde{C}(\mathbf{x}, \mathbf{y})|$$
563
$$= \int d\boldsymbol{\theta}_t \, p(\boldsymbol{\theta}_t | \mathcal{D}_t) \int d\mathbf{y} \, p(\mathbf{y}|\mathbf{x}, \boldsymbol{\theta}_t) \, \log |\tilde{C}(\mathbf{x}, \mathbf{y})|. \quad (20)$$

Note that the outer integral is over the current posterior $p(\theta_t | D_t) \approx$ 565 $\mathcal{N}(\hat{\theta}_t, C_t)$, which is to be distinguished from the future posterior 566 $p(\boldsymbol{\theta}|\mathbf{y}, \mathbf{x}, \mathcal{D}_t) \approx \mathcal{N}(\tilde{\boldsymbol{\theta}}(\mathbf{x}, \mathbf{y}), \tilde{C}(\mathbf{x}, \mathbf{y}))$ whose entropy we are trying 567 to minimize. Whereas the inner integral is simply a weighted sum 568 over the set of outcomes y, the outer integral over the parameter θ 569 is in general challenging, especially when the parameter space is 570 high-dimensional. In the case of the standard multinomial logistic 571 model that does not include lapse, we can exploit the linear struc-572 ture of model to reduce this to a lower-dimensional integral over 573 the space of the linear predictor, which we evaluate numerically 574 using Gauss-Hermite quadrature (Heiss & Winschel, 2008). (This 575 integral is 1D for classic logistic regression, and (k-1)-dimensional 576 for multinomial logistic regression with k classes; see Appendix 577 C for details.) When the model incorporates lapses, the full pa-578 rameter vector $\boldsymbol{\theta} = [\mathbf{w}^{\top}, \mathbf{u}^{\top}]^{\top}$ includes the lapse parameters in 579 addition to the weights w. In this case, our method with Laplace 580 approximation may suffer from reduced accuracy due to the fact 581 that the posterior may be less closely approximated by a Gaussian. 582 In order to exploit the convenient structure of reduced integral 583 over the weight space, we choose to maximize the partial infor-584 mation between the observation and the psychophysical weights, 585 $I(\mathbf{w};\mathbf{v}|\mathbf{x})$, instead of the full information $I(\boldsymbol{\theta};\mathbf{v}|\mathbf{x})$. This is 586 a reasonable approximation in many cases where the stimulus-587 dependent behavior is the primary focus of the psychometric ex-588 periment (also see Prins (2013) for a similar approach). How-589 ever, we note that this is the only piece in this work where we 590 treat the weights separately from the lapse parameters; posterior 591 inference is still performed for the full parameter θ . Thus for 592 Laplace-based infomax exclusively, the partial covariance $C_{ww} =$ 593 $-(\partial^2(\log \mathcal{P})/\partial \mathbf{w}^2)^{-1}$ is used in place of the full covariance 594 $C = -(\partial^2 (\log \mathcal{P})/\partial \theta^2)^{-1}$, where $\mathcal{P}(\theta)$ is the posterior distribution over the full parameter space. Because the positive semidefiniteness of the partial covariance is still not guaranteed, it needs to be approximated to the nearest symmetric positive semi-definite matrix when necessary (Higham, 1988). We can show, however, that the partial covariance is asymptotically positive semi-definite in the small-lapse limit (Appendix A).

Infomax with MCMC. Sampling-based inference provides an 602 attractive alternative to Laplace's method when the model includes 603 non-zero lapse rates, where the posterior may be less well approx-604 imated by a Gaussian. To compute mutual information from sam-605 ples, it is more convenient to use the expansion given in (eq. 15), so 606 that it is expressed as the expected uncertainty reduction in entropy 607 of the response y, instead of a reduction in the posterior entropy. 608 This will make it straightforward to approximate integrals needed 609 for mutual information by Monte Carlo integrals involving sums 610 over samples. Also note that we are back in the full parameter 611 space; we no longer treat the lapse parameters separately, as we 612 did for the Laplace-based infomax. 613

Given a set of posterior samples $\{\theta_m\}$ from $p(\theta|\mathcal{D}_t)$, the posterior distribution at time t, we can evaluate the mutual information using sums over "potential" terms that we denote by

$$L_{jm}(\mathbf{x}) \equiv p(y_j = 1 | \mathbf{x}, \boldsymbol{\theta}_m).$$
(21) 617

618

620

622

This allows us to evaluate the conditional response entropy as

$$H_t(\mathbf{y}|\boldsymbol{\theta}; \mathbf{x}) \approx -\frac{1}{M} \sum_{j,m} L_{jm}(\mathbf{x}) \log L_{jm}(\mathbf{x}), \qquad (22) \quad {}_{619}$$

and the marginal response entropy as

$$H_t(\mathbf{y}; \mathbf{x}) \approx -\sum_j \left(\frac{1}{M} \sum_m L_{jm}(\mathbf{x})\right) \log\left(\frac{1}{M} \sum_m L_{jm}(\mathbf{x})\right), \quad (23) \quad \text{621}$$

where we have evaluated the posterior-predictive distribution as

$$p(y_j = 1 | \mathbf{x}, \mathcal{D}_t) \approx \frac{1}{M} \sum_m L_{jm}(\mathbf{x}).$$
 (24) 623

Putting together these terms, the mutual information can be evaluated as

$$I_t(\boldsymbol{\theta}; \mathbf{y} | \mathbf{x}) = -\frac{1}{M} \sum_{j,m} L_{jm}(\mathbf{x}) \log \frac{L_{jm}(\mathbf{x})}{\sum_{m'} L_{jm'}(\mathbf{x})/M}, \quad (25) \quad \text{626}$$

which is straightforward to evaluate for a set of candidate stimuli $\{x\}$. The computational cost of this approach is therefore linear in the number of samples, and the primary concern is the cost of obtaining a representative sample from the posterior.

631 Results

We consider two approaches for testing the performance of our proposed stimulus-selection algorithms, one using simulated data, and a second using an offline analysis of data from real psychophysical experiments.

Simulated experiments. We first tested the performance of our algorithms using simulated data from a fixed psychophysical observer model. In these simulations, a stimulus x was selected on each trial and the observer's response y was sampled from a "true" psychometric function, $p_{true}(\mathbf{y}|\mathbf{x}) = p(\mathbf{y}|\mathbf{x}, \boldsymbol{\theta}_{true})$.

We considered psychophysical models defined on a continuous 641 2-dimensional stimulus space with 4 discrete response alternatives 642 for every trial, corresponding to the problem of estimating the di-643 rection of 2D stimulus moving along one of the four cardinal di-644 rections (up, down, left, right). We computed expected informa-645 tion gain over a set of discrete stimulus values corresponding to 646 21×21 square grid (Fig. 5A). The stimulus plane is colored in 647 Fig. 5A, to indicate the most likely response (one of the four alter-648 natives) in each stimulus region. Lapse probabilities λc_i were set 649 to either zero (the "lapse-free" case), or a constant value of 0.05, 650 resulting in a total lapse probability of $\lambda = 0.2$ across the four 651 choices (Fig. 5B). We compared performance of our adaptive algo-652 rithms with a method that selected a stimulus uniformly at random 653 from the grid on each trial. We observed that the adaptive methods 654 tended to sample more stimuli near the boundaries between colored 655 regions on the stimulus space (Fig. 5C), which led to more efficient 656 estimates of the PF compared to the uniform stimulus selection ap-657 proach (Fig. 5D). We also confirmed that the posterior entropy of 658 the inferred parameters decrease more rapidly with our adaptive 659 stimulus sampling algorithms, in all cases (Fig. 5E-F). This was 660

expected because our algorithms explicitly attempt to minimize the posterior entropy, by maximizing the mutual information. 662

For each true model, we compared the performances of four dif-663 ferent adaptive methods (Fig. 6A-B), defined by performing infer-664 ence with MAP or MCMC, and assuming lapse rate to be fixed 665 at zero or including a non-zero lapse parameters. Each of these 666 inference methods was also applied to data selected according to 667 a uniform stimulus selection algorithm. We quantified perfor-668 mance using the mean-squared error (MSE) between the true re-669 sponse probabilities $p_{ij} = p(y = j | \mathbf{x}_i, \boldsymbol{\theta}_{true})$ and the estimated 670 probabilities \hat{p}_{ij} over the 21 × 21 grid of stimulus locations $\{\mathbf{x}_i\}$ 671 and the 4 possible responses $\{j\}$. For MAP-based inference, es-672 timated probabilities were given by $\hat{p}_{ij} = p(y = j | \mathbf{x}_i, \boldsymbol{\theta}_{MAP}).$ 673 For the MCMC-based inference, probabilities were given by the 674 predictive distribution, evaluated using an average over samples: 675 $\hat{p}_{ij} = \frac{1}{M} \sum_{m} p(y = j | \mathbf{x}_i, \boldsymbol{\theta}_m)$, where $\{\boldsymbol{\theta}_m\}$ represent samples 676 from the posterior. 677

When the true model was lapse-free (Fig. 6A), lapse-free and 678 lapse-aware inference methods performed similarly, indicating that 679 there was minimal cost to incorporating parameters governing 680 lapse when lapses were absent. Under all inference methods, in-681 fomax stimulus selection outperformed uniform stimulus selec-682 tion by a substantial margin. For example, infomax algorithms 683 achieved in 50 - 60 trials the error levels that their uniform-684 stimulus-selection counterparts required 100 trials to achieve. 685

By contrast, when the true model had a non-zero lapse rate 686 (Fig. 6B), adaptive stimulus selection algorithms based on the 687 lapse-free model failed to select optimal stimuli, performing even 688 worse than uniform stimulus selection algorithms. This empha-689 sizes the impact of model mismatch in adaptive methods, and the 690 importance of a realistic psychometric model. When lapse-aware 691 models were used for inference, on the other hand, both Laplace-692 based and MCMC-based adaptive stimulus selection algorithms 693 achieved a significant speedup compared to uniform stimulus se-694 lection, while MCMC-based adaptive algorithm performed bet-695 ter. This shows that the MCMC-based infomax stimulus selec-696

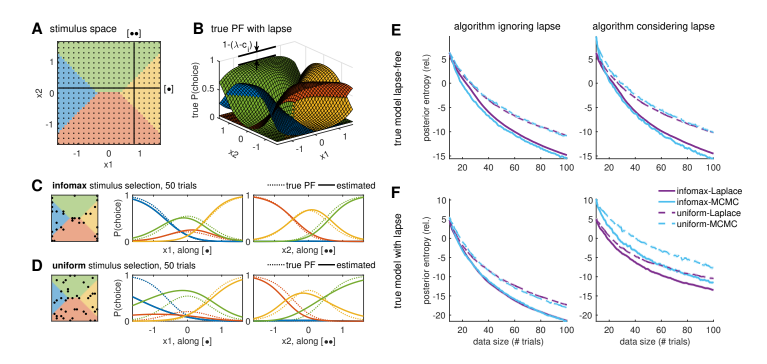


Figure 5: The simulated experiment. (A) At each trial, a stimulus was selected from a 2D stimulus plane with a 21×21 grid. The two lines, running along x_1 and x_2 respectively, indicate the cross-sections used in **C** and **D** below. Colors indicate the most likely response in the respective stimulus regime, according to the true PF shown in **B**, with a consistent color code. (**B**) Given each stimulus, a simulated response was drawn from a true model with 4 alternatives. Shown here is the model with lapse, characterized by a non-deterministic choice (i.e., the choice probability does not approach 0 or 1) even at an easy stimulus, far from the choice boundaries. (**C-D**) Examples of Laplace-approximation-based inference results after 50 trials, where stimuli was selected either using our adaptive infomax method (**C**) or uniformly (**D**), as shown on left. In both cases, the true model was lapse-free, and the algorithm assumed that lapse was fixed at zero. The two sets of curves show the cross-sections of the true PF (dotted lines) and the estimated PF (solid lines), along the two lines marked in **A**, after sampling these stimuli. (**E-F**) Traces of posterior entropy from simulated experiments, averaged over 100 runs each. The true model for simulation was either (**E**) lapse-free, or (**F**) with a finite lapse rate of $\lambda = 0.2$, with a uniform lapse scenario $c_i = 1/4$ for each outcome i = 1, 2, 3, 4. In algorithms considering lapse (panels on the right), the shift in posterior entropy is due to the use of partial covariance (with respect to weight) in the case of Laplace approximation. The algorithm either used the classical MNL model that assumes zero lapse (left column), or our extended model that considers lapse (right column). Average performances of adaptive and uniform stimulus selection algorithms are plotted in solid and dashed lines, respectively; Laplace-based and MCMC-based algorithms are plotted in purple and cyan. The lighter lines show standard error intervals over 100 runs, which are very

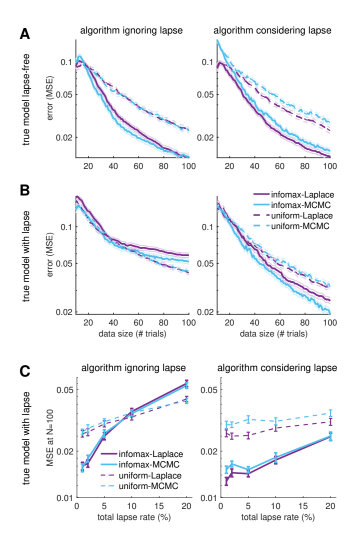


Figure 6: The simulated experiment, continued. We show results from the same set of simulated experiments as in Fig. 5. (A-B) Traces of the mean-square error (MSE), where the true model was either (A) lapse-free, or (B) with a total lapse rate of $\lambda = 0.2$, uniformly distributed to each outcome. Standard error intervals are plotted in lighter lines as in Fig. 5E-F. (C) Effect of lapse, tested by adding varying total lapse rates λ . Shown are the MSE after N = 100 trials of each stimulus selection algorithm, equivalent to the endpoints in **B**. Error bars indicate the standard error over 100 runs, equivalent to the lighter-line intervals in the above panels.

tion method can provide an efficient and robust platform for adaptive experiments with realistic models. When the true behavior
had lapses, the MCMC-based adaptive stimulus selection algorithm with the lapse-aware model automatically included "easy"
trials, which provide maximal information about lapse probabilities. These easy trials are typically in the periphery of the stimulus

space (strong-stimulus regimes, referred to as "asymptotic performance intensity" in Prins (2012)). 704

However, that the effect of model mismatch due to non-zero 705 lapse only becomes problematic at high enough lapse rate; in the 706 simulation shown in Fig. 5F and Fig. 6B, we used a high lapse rate 707 of $\lambda = 0.2$ which is more typical in the case of less sophisti-708 cated animals such as rodents (see for example Scott, Constantino-709 ple, Erlich, Tank, and Brody (2015)). With lapse rates more typ-710 ical in well-designed human psychophysics tasks ($\lambda \lesssim 0.05$; see 711 for example Wichmann and Hill (2001a, 2001b)), infomax algo-712 rithms still tend to perform better than uniform sampling algo-713 rithms (Fig. 6C). 714

Finally, we measured the computation time per trial required by 715 our adaptive stimulus selection algorithms on a personal desktop 716 with an Intel i7 processor. With the Laplace-based algorithm, the 717 major computational bottleneck is the parameter space integration 718 in the infomax calculation, which scales directly with the model 719 complexity. We could easily achieve tens-of-milliseconds trials 720 in the case of the simple 2AFC task, and sub-second trials with 721 2-dimensional stimuli and 4-alternative responses, as used in the 722 current set of simulations (Fig. 7A-B). With the MCMC-based al-723 gorithm, the time-per-trial in the sampling-based method is lim-724 ited by the number of samples in each MCMC chain, M, rather 725 than by the model complexity. Using the standard implementation 726 for the Metropolis-Hastings sampler in Matlab, a time-per-trial of 727 ~ 0.1 seconds was achieved with chains shorter than $M \lesssim 200$ 728 (Fig. 7C-D, top panels). This length of $M \approx 200$ was good enough 729 to represent the posterior distributions for our simulated examples 730 (Fig. 7C-D, bottom panels), although we note that longer chains 731 are required to sample a more complex posterior distribution, and 732 this particular length M should not be taken as the benchmark in 733 general. 734

Optimal re-ordering of real dataset. A second approach for 735 testing the performance of our methods is to perform an off-line 736 analysis of data from real psychophysical experiments. Here we 737 take an existing dataset and use our methods to re-order the trials so 738

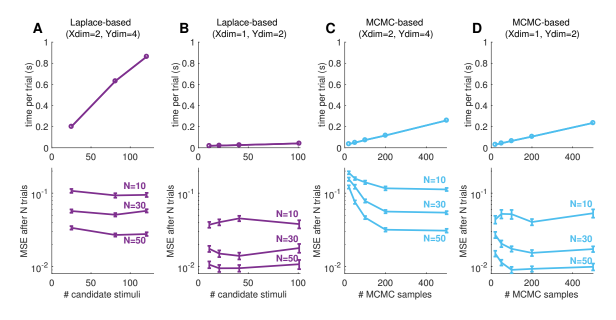


Figure 7: **Computation time and accuracy**. **(A-B)** The computation times for the Laplace-based algorithms grow linearly with the number of candidate stimulus points, as shown on the top panels, because one needs to perform a numerical integration to compute the expected utility of each stimulus. In general, there is a tradeoff between cost (computation time) and accuracy (inversely related to the estimation error). The bottom panels show the mean-square error of the estimated PF, calculated after completing a sequence of *N* trials, where the 10 initial trials were selected at regular intervals, and the following trials were selected under our adaptive algorithm. Error estimates were averaged over 100 independent sequences. Error bars indicate the standard errors. The true model used were the same as either **(A)** in Fig. 5, with 2-dimensional stimulus and 4-alternative response, described by 9 parameters; or **(B)** in Fig. 3, with 1-dimensional stimulus and binary response, with only 2 parameters (slope and threshold). Different rate at which the computation time increases under the two model reflects the different complexity of numerical quadrature involved. We used lapse-free algorithms in all cases in this example. **(C-D)** We similarly tested the MCMC-based algorithms using the two models as in panels **A-B**. In this case, the computation times (top panels) grow linearly the number of samples in each MCMC chain, and are not sensitive to the dimensionality of the parameter space. On the other hand, the estimation error plots (bottom panels) suggest that a high-dimensional model requires more samples for accurate inference.

that the most-informative stimuli are selected first (also see Lewi, 739 Schneider, Woolley, and Paninski (2011) for a similar approach). 740 To obtain a re-ordering, we iteratively apply our algorithm to the 741 stimuli shown during the experiment. On each trial, we use our 742 adaptive algorithm to select the optimal stimulus from the set of 743 stimuli $\{\mathbf{x}_i\}$ not yet incorporated into the model. This selection 744 takes place without access to the actual responses $\{y_i\}$. We then 745 update the posterior using the stimulus x_i and the response y_i it ac-746 tually elicited during the experiment, then proceed to the next trial. 747 We can then ask whether adding the data according to the proposed 748 re-ordering would have led to faster narrowing of the posterior dis-749 tribution than other orderings. 750

monkeys performing a four-alternative motion discrimination task 752 (Churchland, Kiani, & Shadlen, 2008). Monkeys were trained to 753 observe a motion stimulus with dots moving in one of the four car-754 dinal directions, and report this direction of motion with an eye 755 movement. The difficulty of the task was controlled by varying the 756 fraction of coherently moving dots on each trial, with the remain-757 ing dots appearing randomly (Fig. 8A). Each moving-dot stimulus 758 in this experiment could be represented as a two-dimensional vec-759 tor, where the direction of the vector is the direction of the mean 760 movement of the dots, and the amplitude of the vector is given by 761 the fraction of coherently moving dots (a number between 0 and 762 1). Each stimulus presented in the the experiment was aligned with 763 either one of the two cardinal axes of the stimulus plane (Fig. 8B). 764

To perform this analysis, we used a dataset from macaque

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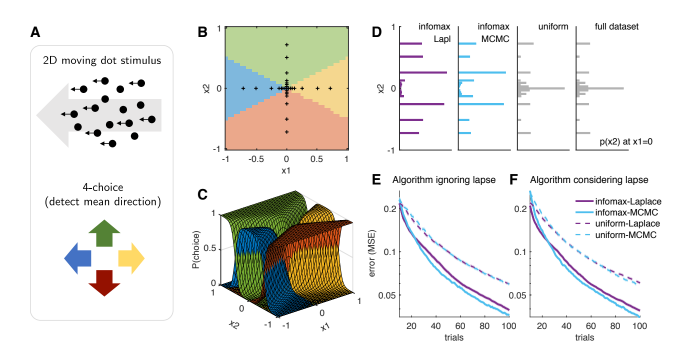


Figure 8: Optimal re-ordering of a real monkey dataset. (A) The psychometric task consisted of a 2D stimulus presented as moving dots, characterized by a coherence and a mean direction of movement, and a 4-alternative response. The four choices are color coded consistently in A-C in this figure. (B) The axes-only stimulus space of the original dataset, with 15 fixed stimuli along each axis. Colors indicate the most likely response in the respective stimulus regime according to the best estimate of the PF. (C) The best estimate of the PF of monkeys in this task, inferred from all observations in the dataset. (D) Stimuli selection in the first N = 100 trials during the re-ordering experiment, under the inference method that ignores lapse. Shown are histograms of x_2 along one of the axes, $x_1 = 0$, averaged over 100 independent runs in each case. (E-F) Error traces under different algorithms, averaged over 100 runs. Both Laplace-based (purple) and MCMC-based (cyan; with M = 1000) algorithms achieve significant speedups over uniform sampling. Because the monkeys were almost lapse-free in this task, inference methods that ignore lapse (E) and consider lapse (F) performed similarly. Standard error intervals over 100 runs are shown in lighter lines, but are very narrow.

The PF for this dataset consists of a set of four 2D curves, where each curve specifies the probability of choosing a particular direction as a function of location in the 2D stimulus plane (Fig. 8C).

This monkey dataset contained more than 10,000 total observations at 29 distinct stimulus conditions, accumulating more than 300 observations per stimulus. This multiplicity of observations per stimulus ensured that the posterior distribution given the full dataset was narrow enough that it could be considered to provide a "ground truth" psychometric function against which the inferences based on the re-ordering experiment could be compared.

The first 100 stimuli selected by the infomax algorithms had noticeably different statistics than the full dataset or its uniform sub-sampling (the first N = 100 trials under uniform sampling). On the other hand, the sets of stimuli selected by both MAPbased and MCMC-based infomax algorithms were similar. Fig. 8D 779 shows the histogram of stimulus component along one of the axes, 780 $p(x_2 | x_1 = 0)$, from the first N = 100 trials, averaged over 100 781 independent runs under each stimulus selection algorithm using the 782 lapse-free model. 783

Because the true PF was unknown, we compared the performance of each algorithm to an estimate of the PF from the entire dataset. When using the MAP algorithm, the full-dataset PF was given by $p_{ij} = p(y = j | \mathbf{x}_i, \hat{\theta}_{full})$, evaluated at the MAP estimate of the log posterior, $\hat{\theta}_{full} = \operatorname{argmax}_{\theta} \log p(\theta | \mathcal{D}_{full})$, given the full dataset \mathcal{D}_{full} . For the MCMC algorithm, the full-dataset PF was computed by $p_{ij} \approx \frac{1}{M} \sum_m p(y = j | \mathbf{x}_i, \theta_m)$, where the MCMC chain $\{\theta_m\} \sim \log p(\theta | \mathcal{D}_{\text{full}})$ sampled the log posterior given the full dataset. The re-ordering test on the monkey dataset showed that our adaptive stimulus sampling algorithms were able to infer the PF to a given accuracy in a smaller number of observations, compared to a uniform sampling algorithm (Fig. 8E-F). In other words, data collection could have been faster with an optimal reordering of the experimental procedure.

Exploiting the full stimulus space. In the experimental 798 dataset considered in the previous section, the motion stimuli were 799 restricted to points along the cardinal axes of the 2D stimulus plane 800 (Fig. 8B) (Churchland et al., 2008). In some experimental settings, 801 however, the psychometric functions of interest may lack identi-802 fiable axes of alignment or may exhibit asymmetries in shape or 803 orientation. Here we show that in such cases, adaptive stimulus 804 selection methods can benefit from the ability to select points from 805 the full space of possible stimuli. 806

We performed experiments with a simulated observer gov-807 erned by the lapse-free psychometric function estimated from the 808 macaque monkey dataset (Fig. 8C). This psychometric function 809 was either aligned to the original stimulus axes (Fig. 9A-B) or ro-810 tated counter-clockwise by 45 degrees (Fig. 9C). We tested the per-811 formance of adaptive stimulus selection using the Laplace infomax 812 algorithm, with stimuli restricted to points along the cardinal axes 813 814 (Fig. 9A), or allowed to a grid of points in the full 2D stimulus plane (Fig. 9B-C). 815

The simulated experiment indeed closely resembled the results 816 of our dataset re-ordering test in terms of the statistics of adap-817 tively selected stimuli (compare Fig. 9A to the purple histogram in 818 Fig. 8D). With the full 2D stimulus space aligned to the cardinal 819 axes, on the other hand, our adaptive infomax algorithm detected 820 and sampled more stimuli near the boundaries between colored re-821 gions in the stimulus plane, which were usually not on the cardi-822 nal axes (Fig. 9B). Finally, we also observed that this automatic 823 exploitation of the stimulus space was not limited by the lack of 824 alignment between the PF and the stimulus axes; our adaptive in-825 fomax algorithm was just as effective in detecting and sampling the 826

boundaries between stimulus regions in the case of the unaligned 827 PF (Fig. 9C). 828

The error traces in Fig. 9D show that we can infer the PF at a 829 given accuracy in an even fewer number of observations using our 830 adaptive algorithm on the full 2D stimulus plane (orange curves), 831 compared to the cardinal-axes design (black curves). It also con-832 firms that we can infer the PF accurately and effectively with an 833 unaligned stimulus space (red curves), as well as with an aligned 834 stimulus space. For comparison purposes, all errors were calcu-835 lated over the same 2D stimulus grid, even when the stimulus se-836 lection was from the cardinal axes. (This had negligible effects on 837 the resulting error values: compare the black curves in Fig. 9D and 838 the purple curves in Fig. 8E.) 839

Discussion

We developed effective Bayesian adaptive stimulus selection algorithms for inferring psychometric functions, with an objective of maximizing the expected informativeness of each stimulus. The algorithms select an optimal stimulus adaptively in each trial, based on the posterior distribution of model parameters inferred from the accumulating set of past observations.

840

We emphasized that in psychometric experiments, especially 847 with animals, it is crucial to use models that can account for the 848 non-ideal yet common behaviors, such as omission (no response; 849 an additional possibility for the outcome) or lapse (resulting in 850 a random, stimulus-independent response). Specifically, we con-851 structed a hierarchical extension of a multinomial logistic (MNL) 852 model that incorporates both omission and lapse. Although we 853 did not apply these additional features to real data, we performed 854 simulated experiments to investigate their impacts on the accurate 855 inference of psychometric functions. To ensure applicability of the 856 extended model in real-time closed-loop adaptive stimulus selec-857 tion algorithms, we also developed efficient methods for inferring 858 the posterior distribution of the model parameters, with approxi-859 mations specifically suited for sequential experiments. 860

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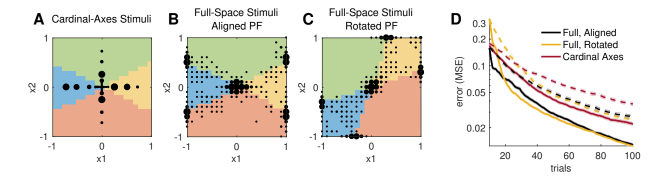


Figure 9: Design of multi-dimensional stimulus space. (**A-C**) Three different stimulus space designs were used in a simulated psychometric experiment. Responses were simulated according to fixed lapse-free PFs, matched to our best estimate of the monkey PF (Fig. 8C). Stimuli were selected within the respective stimulus spaces, (**A**) the cardinal-axes design, as in the original experiment; (**B**) full stimulus plane, with the PF aligned to the cardinal axes of the original stimulus space; (**C**) full stimulus plane, with rotated PF. The black dots in **A-C** indicate which stimuli were sampled by the Laplace-based infomax algorithm during the first N = 100 trials of simulation, where the dot size is proportional to the number of trials in which each stimulus was selected (averaged over 20 independent runs, and excluding the 10 fixed initial stimuli). (**D**) The corresponding error traces, under infomax (solid lines) or uniform (dashed lines) stimulus selection, averaged over 100 runs respectively. Colors indicate the three stimulus space designs, as shown in **A-C**. Standard error intervals over 100 runs are shown in lighter lines.

Advantages of adaptive stimulus selection. We observed 861 two important advantages of using Bayesian adaptive stimulus se-862 lection methods in psychometric experiments. First, we showed 863 that our adaptive stimulus selection algorithms achieved signifi-864 cant speed-ups in learning time (number of measurements), both 865 on simulated data and in re-ordering test of a real experimental 866 dataset, with and without lapse in the underlying behavior. Impor-867 tantly, the success of the algorithm depends heavily on the use of 868 the correct model family; for example, adaptive stimulus selection 869 fails when a classical (lapse-ignorant) model was used to measure 870 behavior with a finite lapse rate. Based on the simulation results, 871 it seems good practice to always use the lapse-aware model unless 872 the behavior under study is known to be completely lapse-free, al-873 though it should be checked that the addition of the lapse param-874 eters does not make the inference problem intractable, given the 875 constraints of the specific experiments. (One way to check this is 876 using a simulated experiment, where lapse is added to the psycho-877 metric function inferred by lapse-free model; similarly to what we 878 did in this paper.) The computational cost for incorporating lapses 879 amounts to having k additional parameters to sample, one per each 880 available choice, which is independent from the dimensionality of 881

the stimulus space.

Second, we demonstrated that our adaptive stimulus selection 883 study has implications on the optimization of the experimental de-884 signs more generally. Contrary to the conventional practice of ac-885 cumulating repeated observations at a small set of fixed stimuli, we 886 suggest that the (potentially high-dimensional) stimulus space can 887 be exploited more efficiently using our Bayesian adaptive stimulus 888 selection algorithm. Specifically, the adaptive stimulus selection 889 algorithm can automatically detect the structure of the stimulus 890 space (with respect to the psychometric function) as part of the 891 process. We also showed that there are benefits of using the full 892 stimulus space even when the PF is aligned to the cardinal axes of 893 the stimulus space. 894

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Comparison of the two algorithms. Our adaptive stimulus selection algorithms were developed based on two methods for effective posterior inference: one based on local Gaussian approximation (Laplace approximation) of the posterior, and another based on MCMC sampling. The well-studied analytical method based on the Laplace approximation is fast and effective in simple cases, but becomes heavier in the case of more complicated PFs, because

the computational bottleneck is the numerical integration over the 902 parameter space that needs to be performed separately for each 903 candidate stimulus. In the case of sampling-based methods, on 904 the other hand, the computational speed is constrained by the num-905 ber of MCMC samples used to approximate the posterior distribu-906 tion, but not directly by the number of parameters or the number 907 of candidate stimuli. In general, however, accurately inferring a 908 higher-dimensional posterior distribution requires more samples, 909 and therefore a longer computation time. We note that our semi-910 adaptive turning algorithm helps with the cost-accuracy tradeoff 911 by optimizing the sampling accuracy in a given number of sam-912 ples, without human intervention. although it does not reduce the 913 computation time itself. 914

To summarize, when the PF under study is low-dimensional and 915 well-described by the multinomial logistic model, for example in 916 a 2AFC study with human subjects, Laplace-based approach pro-917 vides a lightweight and elegant approach. But if the PF is higher-918 dimensional or deviates significantly from the ideal model (e.g., 919 large lapse), MCMC sampling provides a flexible and affordable 920 solution. Results suggest that our MCMC-based algorithm will be 921 applicable to most animal psychometric experiments, as the model 922 complexities are not expected to significantly exceed our simulated 923 example. However, one should always make sure that the number 924 of MCMC samples being used is sufficient to sample the posterior 925 distribution under study. 926

Limitations and Open Problems. One potential drawback 927 of adaptive experiments is the undesired possibility that the psy-928 chometric function of the observer might adapt to the distribution 929 of stimuli presented during the experiments. If this is the case, 930 the system under measurement would no longer be stationary, nor 931 independent of the experimental design, profoundly altering the 932 problem one should try to solve. The usual assumption in psycho-933 metric experiments is that well trained observers exhibit stationary 934 behavior on the timescale of an experiment; under this assumption, 935 the order of data collection cannot bias inference MacKay (1992). 936 However, the empirical validity of this claim remains a topic for 937

future research.

One approach for mitigating non-stationarity is to add regressors 939 to account for the history dependence of psychophysical behavior. 940 Recent work has shown that extending a psychophysical model to 941 incorporate past rewards (Bak et al., 2016; Busse et al., 2011; Cor-942 rado, Sugrue, Seung, & Newsome, 2005; Lau & Glimcher, 2005), 943 past stimuli (Akrami, Kopec, Diamond, & Brody, 2018) or the full 944 stimulus-response history (Fründ, Wichmann, & Macke, 2014) can 945 provide a more accurate description of the factors influencing re-946 sponses on a trial-by-trial basis. 947

938

Our work leaves open a variety of directions for future research. 948 One simple idea is to re-analyze old datasets under the multinomial 949 response model with omissions included as a separate response cat-950 egory; this will reveal whether omissions exhibit stimulus depen-951 dence (e.g., occurring more often on difficult trials), and provide 952 greater insight into the factors influencing psychophysical behavior 953 on single trials. Another set of directions is to extend the multino-954 mial logistic observer model to obtain a more accurate or more 955 flexible model of psychophysical behavior; particular directions 956 include models with nonlinear stimulus dependencies or interac-957 tion terms (Cowley, Williamson, Clemens, Smith, & Byron, 2017; 958 DiMattina & Zhang, 2011; Hyafil & Moreno-Bote, 2017; Neri & 959 Heeger, 2002), models with output nonlinearities other than the 960 logistic (Kontsevich & Tyler, 1999; Schütt et al., 2016; A. B. Wat-961 son, 2017; A. B. Watson & Pelli, 1983), or models that capture 962 overdispersion, e.g., due to non-stationarities of the observer, via a 963 hierarchical prior (Schütt et al., 2016). In general, such extensions 964 will be much easier to implement with the MCMC-based inference 965 method, due to the fact that it does not rely on gradients or Hessians 966 of a particular parametrization of log-likelihood. Finally, it may be useful to consider the same observer model under optimality cri-968 teria other than mutual information - recent work has shown that 969 infomax methods do not necessarily attain optimal performance 970 according to alternate metrics (e.g., mean squared error, I. M. Park 971 and Pillow (2017); M. Park et al. (2014)) — or using non-greedy 972 selection criteria that optimize stimulus selection based on a time 973

⁹⁷⁴ horizon longer than the next trial (Kim et al., 2017; King-Smith et⁹⁷⁵ al., 1994).

976 Code availability

A Matlab implementation of our methods is available online at
https://github.com/pillowlab/adaptivePsychophysicsToolbox.

979 Acknowledgements

We thank Anne Churchland for sharing the monkey data. JHB was supported by the Samsung Scholarship for the study at Princeton. JWP was supported by grants from the McKnight Foundation, Simons Collaboration on the Global Brain (SCGB AWD1004351) and the NSF CAREER Award (IIS-1150186). Computational work was performed using resources at Princeton University and the KIAS Center for Advanced Computing.

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Appendix A

Log likelihood for the classical MNL. Here we provide more the multinodetails about the log likelihood $L = \mathbf{y}^{\top} \log \mathbf{p}$ under the multinomial logistic model (6), first in the lapse-free case.

1199

A convenient property of the multinomial logistic model (a prop-1203 erty common to all generalized linear models) is that the parameter 1204 vector p_i governing y depends only on a 1-dimensional projection 1205 of the input, $V_i = \phi^{\top} \mathbf{w}_i$, which is known as the *linear predictor*. 1206 Recall that $\phi = \phi(\mathbf{x})$ is the input feature vector. In the multinomial 1207 case, it is useful to consider the column vector of linear predictors 1208 for a single trial, $\mathbf{V} = [V_1, \cdots, V_k]^{\top}$, and the concatenated weight 1209 vector $\mathbf{w} = [\mathbf{w}_1^{\top}, \cdots, \mathbf{w}_k^{\top}]^{\top}$, consisting of all weights stacked 1210 into a single vector. We can summarize their linear relationship 1211 as $\mathbf{V} = X\mathbf{w}$, where X is a block diagonal matrix containing k 1212 blocks of ϕ^{\top} along the diagonal. In other words, 1213

$$X = \begin{bmatrix} \boldsymbol{\phi}^{\top} & \boldsymbol{0}^{\top} & \cdots & \boldsymbol{0}^{\top} \\ \boldsymbol{0}^{\top} & \boldsymbol{\phi}^{\top} & \cdots & \boldsymbol{0}^{\top} \\ \vdots & \vdots & \ddots & \vdots \\ \boldsymbol{0}^{\top} & \boldsymbol{0}^{\top} & \cdots & \boldsymbol{\phi}^{\top} \end{bmatrix}, \quad \mathbf{w} = \begin{bmatrix} \mathbf{w}_1 \\ \mathbf{w}_2 \\ \vdots \\ \mathbf{w}_k \end{bmatrix}. \quad (26) \quad {}_{1214}$$

Derivatives. It is convenient to work in terms of the linear predictor $\mathbf{V} = \{V_i\}$ first. If $N_y \equiv \sum_i y_i = 1$ is the total number of 1216 responses per trial, the first and second derivatives of L with respect 1217 to \mathbf{V} are $\partial L/\partial V_j = y_j - N_y p_j$ and $\partial^2 L/\partial V_i \partial V_j = N_y p_i (\delta_{ij} - p_j)$, 1218 respectively. Rewriting in vector forms, we have 1219

$$\frac{\partial L}{\partial \mathbf{V}} = (\mathbf{y} - N_y \mathbf{p})^\top, \tag{27}$$

$$\frac{\partial \mathbf{V}}{\partial \mathbf{V}^2} = -N_y \left(\operatorname{diag}(\mathbf{p}) - \mathbf{p} \mathbf{p}^\top \right) \equiv -N_y \Gamma(\mathbf{p}), \quad (28) \quad \text{(28)} \quad \text{(28)}$$

where $\operatorname{diag}(\mathbf{p}) = [p_i \delta_{ij}]$ is a square matrix with the elements of \mathbf{p} 1223 on the diagonal, and zeros otherwise. 1224

Putting back in terms of the weight vector \mathbf{w} is easy, thanks to the linear relationship $\mathbf{V} = X\mathbf{w}$:

$$\frac{\partial L}{\partial \mathbf{w}} = \frac{\partial L}{\partial \mathbf{V}} X = (\mathbf{y} - \mathbf{p})^{\top} X \equiv \mathbf{\Delta}^{\top}, \qquad (29) \quad {}_{1227}$$

$$\frac{\partial^2 L}{\partial \mathbf{w}^2} = X^\top \frac{\partial^2 L}{\partial \mathbf{V}^2} X = -X^\top \Gamma X \equiv -\Lambda. \tag{30}$$
¹²²⁸
¹²²⁹

)

Concavity. Importantly, L is concave with respect to V (and 1230 therefore with respect to \mathbf{w}). To prove the concavity of L, we show 123 that the Hessian $H = -\text{diag}(\mathbf{p}) + \mathbf{p}\mathbf{p}^{\top} \equiv -\Gamma$ is negative semi-1232 definite, which is equivalent to showing that $\mathbf{z}^{\top} \Gamma \mathbf{z} \geq 0$: 1233

$$\mathbf{z}^{\top} \Gamma \mathbf{z} = \mathbf{z}^{\top} \operatorname{diag}(\mathbf{p}) \mathbf{z} - (\mathbf{z}^{\top} \mathbf{p})^2$$

1236

1237

$$= \sum_{i} z_{i}^{2} p_{i} - \left(\sum_{j} z_{j} p_{j}\right)^{2}$$
$$= \sum_{i} p_{i} \left[\left(z_{i} - \sum_{j} z_{j} p_{j} \right)^{2} \right] \ge 0$$
(31)

for an arbitrary vector z. 1238

Log likelihood with lapse. With a finite lapse rate λ , to recap, 1239 the multinomial logistic model is modified as $p_i = (1 - \lambda)q_i + \lambda c_i$ 1240 where 1241

1242
$$q_i = \frac{\exp(V_i)}{\sum_j \exp(V_j)}, \quad \lambda c_i = \frac{\exp(u_i)}{1 + \sum_j \exp(u_j)}.$$
 (32)

Let us introduce the following abbreviations, 1243

1244
$$r_i \equiv \frac{\lambda c_i}{p_i}, \quad t_i \equiv y_i(1-r_i), \quad s_i \equiv y_i r_i(1-r_i),$$
 (33)

where the dimensionless ratio $r \in [0, 1]$ can be considered as the 1245 order parameter for the effect of lapse. 1246

Derivatives with respect to the weights. Differentiating with the 1247

linear predictor V, we get 1248

1249
$$\frac{\partial q_i}{\partial V_l} = (\delta_{il} - q_l)q_i,$$
1250
$$\frac{\partial^2 q_i}{\partial V_j \partial V_l} = \left[(\delta_{ij} - q_j)(\delta_{il} - q_l) - (\delta_{jl}q_l - q_jq_l) \right]q_i.$$

which leads to 1252

$$\frac{\partial p_i}{\partial V_l} = (1-\lambda)\frac{\partial q_i}{\partial V_l}, \quad \frac{\partial^2 p_i}{\partial V_j \partial V_l} = (1-\lambda)\frac{\partial^2 q_i}{\partial V_j \partial V_l}.$$

We are interested in the derivatives of the log likelihood L = $\mathbf{y}^{\top} \log \mathbf{p}$ with respect to V. The partial gradient: 1255

$$\frac{\partial L}{\partial V_l} = \sum_i y_i \frac{1}{p_i} \frac{\partial p_i}{\partial V_l} = (1 - \lambda) \sum_i y_i \frac{q_i}{p_i} (\delta_{il} - q_l)$$

$$= t_l - q_l \sum_i t_i.$$
1258

1258

Similarly, the partial Hessian is written as

$$\frac{\partial^2 L}{\partial V_j \partial V_l} = \sum_i y_i \left(\frac{1}{p_i} \frac{\partial^2 p_i}{\partial V_j \partial V_l} - \frac{1}{p_i^2} \frac{\partial p_i}{\partial V_j} \frac{\partial p_i}{\partial V_l} \right)$$
 1260

$$= \delta_{jl} \left(s_l - q_l \sum_i t_i \right) - \left(q_j s_l + q_l s_j \right) + q_j q_l \left(\sum_i s_i + \sum_i t_i \right).$$

In vector forms, and with $\tau \equiv \sum_i t_i$ and $\sigma \equiv \sum_i s_i$,

$$\frac{\partial L}{\partial \mathbf{V}} = (\mathbf{t} - \tau \mathbf{q})^{\top}; \qquad (34) \quad {}_{1264}$$

1259

1263

1285

1289

$$\frac{\partial^2 L}{\partial \mathbf{V}^2} = \operatorname{diag}(\mathbf{s} - \tau \mathbf{q}) - (\mathbf{q} \mathbf{s}^\top + \mathbf{s} \mathbf{q}^\top) + (\tau + \sigma) \mathbf{q} \mathbf{q}^\top$$
¹²⁶⁶

$$= -\tau \left[\operatorname{diag}(\mathbf{q}) - \mathbf{q} \mathbf{q}^{\top} \right]$$
 1267

+
$$\left[\operatorname{diag}(\mathbf{s}) - (\mathbf{q}\mathbf{s}^{\top} + \mathbf{s}\mathbf{q}^{\top}) + \sigma \mathbf{q}\mathbf{q}^{\top}\right].$$
 (35) ¹²⁶

Note that we recover $t_i \rightarrow y_i$ and $s_i \rightarrow 0$ in the lapse-free limit 1270 $\lambda \rightarrow 0$. Hence the first square bracket in (35) reduces back to 1271 the lapse-free Hessian, while the second square bracket vanishes 1272 as $\lambda \to 0$. 1273

In the presence of lapse, one might still be interested in the 1274 partial Hessian with respect to the weight parameters, $H \equiv$ 1275 $\partial^2 L/\partial \mathbf{V}^2$, which should be evaluated as in (35). To test the nega-1276 tive semi-definiteness of this partial Hessian, again for an arbitrary 1277 vector z, we end up with 1278

$$\mathbf{z}^{\top}H\mathbf{z} = -\sum_{j} t_{j} \left\langle (z - \langle z \rangle_{q})^{2} \right\rangle_{q} + \sum_{j} s_{j} \left(z_{j} - \langle z \rangle_{q} \right)^{2} \quad (36) \quad {}_{1276}$$

where $\langle x \rangle_q = \sum_j x_j q_j$. The partial Hessian is asymptotically neg-1281 ative semi-definite (which is equivalent to the log likelihood being 1282 concave) in the lapse-free limit, where $t_i \rightarrow y_i$ and $s_i \rightarrow 0$. 1283

Derivatives with respect to lapse parameters. From (2) and (3), 1284

we have $p_i = (1 - \lambda)q_i + \lambda c_i$ where

$$c_i = \frac{\exp(u_i)}{\sum_j \exp(u_j)}; \qquad \lambda = \frac{\sum_j \exp(u_j)}{1 + \sum_j \exp(u_j)}. \tag{37}$$

Differentiating with respect to the auxiliary lapse parameter u_i , 1287

$$\frac{\partial c_i}{\partial u_j} = (\delta_{ij} - c_i)c_j; \qquad \frac{\partial \lambda}{\partial u_j} = (1 - \lambda)\lambda c_j. \tag{38}$$

The gradient is then

$$\frac{\partial p_i}{\partial u_j} = (\delta_{ij} - p_i) \lambda c_j; \tag{39}$$

 $_{1292}$ using the abbreviations in (33), the gradient of the log likelihood is

$$\frac{\partial L}{\partial u_j} = \sum_i y_i \frac{1}{p_i} \frac{\partial p_i}{\partial u_j} = r_j \left(y_j - N_y \cdot p_j \right). \tag{40}$$

¹²⁹⁵ Second derivative with respect to lapse:

$$\frac{\partial^2 p_i}{\partial u_j \partial u_l} = \delta_{jl} \frac{\partial p_i}{\partial u_l} - (\delta_{ij} + \delta_{il} - 2p_i)\lambda c_l \lambda c_j; \qquad (41)$$

1298 it is useful to notice that

$$\frac{\partial p_i}{\partial u_j} \frac{\partial p_i}{\partial u_l} = \delta_{jl} \frac{\partial p_i}{\partial u_l} \lambda c_l - p_i (\delta_{ij} + \delta_{il} - 2p_i) \lambda c_l \lambda c_j.$$
(42)

¹³⁰¹ The corresponding part of the Hessian:

$$\frac{\partial^2 L}{\partial u_j \partial u_l} = \sum_i y_i \left(\frac{1}{p_i} \frac{\partial^2 p_i}{\partial u_j \partial u_l} - \frac{1}{p_i^2} \frac{\partial p_i}{\partial u_j} \frac{\partial p_i}{\partial u_l} \right)$$

$$= \delta_{jl} \sum_i y_i \frac{1}{p_i} \left(1 - \frac{\lambda c_l}{p_i} \right) \frac{\partial p_i}{\partial u_l}$$

$$= \delta_{jl} \left(s_l - r_l p_l N_y + r_l^2 p_l^2 \sum_i \frac{y_i}{p_i} \right). \quad (43)$$

¹³⁰⁶ Finally, the mixed derivative:

$$\frac{\partial^2 p_i}{\partial u_j \partial V_l} = -(1-\lambda)\lambda c_j \cdot (\delta_{il} - q_l)q_l.$$
(44)

1309 again it is useful to notice that

$$\frac{\partial p_i}{\partial u_j} \frac{\partial p_i}{\partial V_l} = -(\delta_{ij} - p_i) \frac{\partial^2 p_i}{\partial u_j \partial V_l}.$$
(45)

1312 Hence

$$\frac{\partial^2 L}{\partial u_j \partial V_l} = \sum_i y_i \left(\frac{1}{p_i} \frac{\partial^2 p_i}{\partial u_j \partial V_l} - \frac{1}{p_i^2} \frac{\partial p_i}{\partial u_j} \frac{\partial p_i}{\partial V_l} \right)$$

$$= -s_j \left(\delta_{jl} + \frac{q_l^2}{q_j} \right). \tag{46}$$

From (40), (43) and (46), we see that all derivatives involving the lapse parameter scale with at least one order of r, therefore vanishing in the lapse-free limit $\lambda \to 0$.

Appendix B

The Metropolis-Hastings algorithm. The Metropolis-Hastings algorithm (Metropolis et al., 1953) generates a chain of samples, using a proposal density and a method to accept or reject the proposed moves.

A proposal is made at each iteration, where the algorithm randomly chooses a candidate for the next sample value \mathbf{x}' based on 1325 the current sample value \mathbf{x}_t . The choice follows the proposal den-1326 sity function, $\mathbf{x}' \sim Q(\mathbf{x}' | \mathbf{x}_t)$. When the proposal density Q is 1327 symmetric, for example a Gaussian, the sequence of samples is a 1328 random walk. In general the width of Q should match with the 1329 statistics of the distribution being sampled, and individual dimen-1330 sions in the sampling space may behave differently in the multi-1331 variate case; finding the appropriate Q can be difficult. 1332

The proposed move is either accepted or rejected with some 1333 probability; if rejected, the current sample value is reused in the 1334 next iteration, $\mathbf{x}' = \mathbf{x}_t$. The probability of acceptance is deter-1335 mined by comparing the values of $P(\mathbf{x}_t)$ and $P(\mathbf{x}')$, where $P(\mathbf{x})$ is 1336 the distribution being sampled. Because the algorithm only consid-1337 ers the acceptance ratio $\rho = P(\mathbf{x}')/P(\mathbf{x}_t) = f(\mathbf{x}')/f(\mathbf{x}_t)$ where 1338 $f(\mathbf{x})$ can be any function proportional to the desired distribution 1339 $P(\mathbf{x})$, there is no need to worry about the proper normalization 1340 of the probability distribution. If $\rho \geq$ 1, the move is always ac-1341 cepted; if $\rho < 1$, it is accepted with a probability ρ . Consequently 1342 the samples tend to stay in the high-density regions, visiting the 1343 low-density regions only occasionally. 1344

Optimizing the sampler. One of the major difficulties in using 1345 the MCMC method is to make an appropriate choice of the pro-1346 posal distribution, which may significantly affect the performance 1347 of the sampler. If the proposal distribution is too narrow, it will 1348 take a long time for the chain to diffuse away from the starting 1349 point, producing a chain with highly correlated samples, requiring 1350 a long time to achieve independent samples. On the other hand if 1351 the proposal distribution is too wide, most of the proposed moves 1352 would be rejected, once again resulting in the chain stuck at the ini-1353 tial point. In either case the chain would "mix" poorly (Rosenthal, 1354 2011). In this paper we restrict our consideration to the Metropolis-1355 Hastings algorithm (Metropolis et al., 1953), although the issue of 1356 proposal distribution optimization is universal in most variants of 1357 MCMC algorithms, only with implementation-level differences. 1358

The basic idea is that the optimal width of the proposal distribu-

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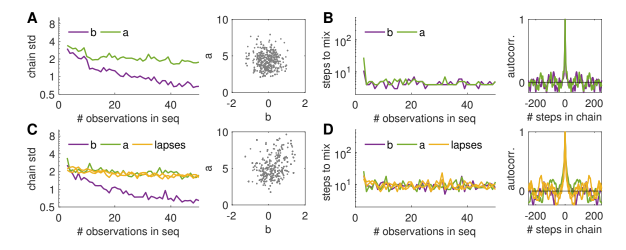


Figure 10: Statistics of the semi-adaptive MCMC in a simulated experiment, with M = 1000 samples per chain. We used the same binomial model as in Fig. 3, and the uniform stimulus selection algorithm. (**A-B**) In a lapse-free model: (**A**) The standard deviation of the samples, along each dimension of the parameter space, decreases as the learning progresses, as expected because the posterior distribution should narrow down as more observations are collected. Also shown is the scatter plot of all 1000 samples at the last trial N = 50, where the true parameter values are (a, b) = (5, 0). (**B**) The mixing time of the chain (number of steps before the autocorrelation falls to 1/e) quickly converges to some small value, meaning that the sampler is quickly optimized. Autocorrelation function at the last trial N = 50 is shown. (**C-D**) Same information as (A) and (B), but with a lapse rate of $\lambda = 0.1$, with uniform lapse $(c_1 = c_2 = 1/2)$.

tion would be determined in proportion to the typical length scale 1360 of the distribution being sampled. This idea was made precise in 1361 the case of a stationary random-walk Metropolis algorithm with 1362 Gaussian proposal distributions, by comparing the covariance ma-1363 trix Σ_p of the proposal distribution to the covariance matrix Σ of 1364 the sampled chain. Once a linear scaling relation $\Sigma_p = s_d \Sigma$ is 1365 fixed, it was observed that it is optimal to have $s_d = (2.38)^2/d$ 1366 where d is the dimensionality of the sampling space (Gelman et 1367 al., 1996; Roberts et al., 1997). An adaptive Metropolis algo-1368 rithm (Haario et al., 2001) followed this observation, where the 1369 Gaussian proposal distribution adapts continuously as the sampling 1370 progresses. Their adaptive algorithm used the same scaling rule 1371 $\Sigma_p = s_d \Sigma$, but updates Σ_p at each proposal where Σ is covariance 1372 of the samples accumulated so far. Additionally, a small diagonal 1373 component was added for stability, as $\Sigma_p = s_d(\Sigma + \epsilon I)$. We used 1374 $\epsilon = 0.0001$ in this work. 1375

Here we propose and use the semi-adaptive Metropolis-Hastings
algorithm, which is a coarse-grained version of the original adaptive algorithm by Haario et al. (2001). The major difference in

our algorithm is that the adjustment of the proposal distribution is 1379 made only at the end of each (sequential) chain, rather than at each 1380 proposal within the chain. This coarse-graining is a reasonable ap-1381 proximation because we will be sampling the posterior distribution 1382 many times as it refines over the course of data collection, once 1383 after each trial. Assuming that the change in posterior distribu-1384 tion after each new observation is small enough, we can justify our 1385 use of the statistics of the previous chain to adjust the properties 1386 of the current chain. Unlike in the fully adaptive algorithm where 1387 the proposal distribution needs to stabilize quickly within a single 1388 chain, we can allow multiple chains until stabilization, usually a 1389 few initial observations - leaving some room for the coarse-grained 1390 approximation. This is because, for our purpose, it is not impera-1391 tive that we have a good sampling of the distribution at the very 1392 early stage of the learning sequence where the accuracy is already 1393 limited by the smallness of the dataset. 1394

When applied to the sequential learning algorithm, our semiadaptive Metropolis sampler shows a consistent well-mixing property after a few initial adjustments, with the standard deviation

of each sampling dimension decreasing stably as data accumulate 1398 (Fig. 10). Whereas Kujala and Lukka (2006) also had the idea of 1399 adjusting the proposal density between trials, their scaling factor 1400 was fixed and independent of the sampling dimension. Building on 1401 more precise statistical observations, our method generalize well to 1402 high-dimensional parameter spaces, typical for multiple-alternative 1403 models. Our semi-adaptive sampler provides an efficient and ro-1404 bust alternative to the particle filter implementations (Kujala & 1405 Lukka, 2006), which has the known problem of weight degener-1406 ation (DiMattina, 2015) as the posterior distribution narrows down 1407 with the accumulation of data. 1408

Appendix C

Fast sequential update of the posterior, with Laplace ap-1410 **proximation.** Use of Laplace approximation was shown to be 1411 particularly useful in a sequential experiment (Lewi et al., 2009), 1412 where it can be assumed that the posterior distribution after the 1413 next trial in sequence, \mathcal{P}_{t+1} , would not be very different from the 1414 current posterior \mathcal{P}_t . Let us consider the lapse-free case $\theta = \mathbf{w}$ for 1415 the moment, where the use of Laplace approximation is valid. Re-1416 arranging from (7) and (9), the sequential update for the posterior 1417 distribution is 1418

¹⁴²⁰ or with Laplace approximation,

$$\log \mathcal{N}(\mathbf{w}|\boldsymbol{\theta}_{t+1}, C_{t+1}) \approx \log \mathcal{N}(\mathbf{w}|\boldsymbol{\theta}_t, C_t) + L_{t+1}(\mathbf{w})$$
(48)

 $\log \mathcal{P}_{t+1}(\mathbf{w}) = \log \mathcal{P}_t(\mathbf{w}) + L_{t+1}(\mathbf{w});$

where $L_i(\mathbf{w}) = \log p(\mathbf{y}_i | \mathbf{x}_i, \mathbf{w})$ is a shorthand for the log likelihood of the *i*-th observation.

¹⁴²⁴ With this, we can achieve a fast sequential update of the posterior ¹⁴²⁵ without performing the full numerical optimization each time. Be-¹⁴²⁶ cause the new posterior mode θ_{t+1} is where the gradient vanishes, ¹⁴²⁷ it can be approximated from the previous mode θ_t by taking the ¹⁴²⁸ first derivative of (48). The posterior covariance C_{t+1} is similarly approximated by taking the second derivate.

$$\boldsymbol{\theta}_{t+1} = \boldsymbol{\theta}_t + C_t \boldsymbol{\Delta}_{t+1}, \qquad \boldsymbol{\Delta}_{t+1} = \left. \frac{\partial L_{t+1}}{\partial \mathbf{w}} \right|_{\mathbf{w} = \boldsymbol{\theta}_t}$$
(49) 143

$$C_{t+1} = \left(C_t^{-1} + \Lambda_{t+1}\right)^{-1}, \quad \Lambda_{t+1} = -\left.\frac{\partial^2 L_{t+1}}{\partial \mathbf{w}^2}\right|_{\mathbf{w}=\boldsymbol{\theta}_{t+1}} \tag{50} \quad \text{(50)} \quad \text{(51)}$$

Using the matrix inversion lemma (Henderson & Searle, 1981), we 1433 can rewrite the posterior covariance update as 1434

$$C_{t+1} = C_t \left[I - (I + \Lambda_{t+1}C_t)^{-1} \Lambda_{t+1}C_t \right].$$
 (51) 1438

Unlike in the earlier application of this trick (Lewi et al., 2009), the 1436 covariance matrix update (50) is not a rank-one update, because of 1437 the multinomial nature of our model (our linear predictor y is a 1438 vector, not a scalar as in a binary model). 1439

Integration over the parameter space: reducing the inte-1440 gration space. The evaluation of expected utility function usu-1441 ally involves a potentially high-dimensional integral over the pa-1442 rameter space. With the Gaussian approximation of the posterior, 1443 we can reduce and standardize the integration space. The process 1444 consists of three steps: diagonalization, marginalization, and stan-1445 dardization. First we choose a new "coordinate system" of the (say 1446 q-dimensional) weight space, such that the first k elements of the 1447 extended weight vector w are coupled one-to-one to the elements 1448 of k-vector y. Then we marginalize to integrate out the remaining 1449 (q - k) dimensions, effectively changing the integration variable 1450 from w to y. Finally, we use Cholesky decomposition to stan-1451 dardize the normal distribution which is the posterior on y. The 1452 resulting integral is still multi-dimensional, due to the multinomial 1453 nature of our model. But once the distribution is standardized, there 1454 are a number of efficient numerical integration methods that can be 1455 applied. For example, in this work, we use the Sparse Grid method 1456 (Heiss & Winschel, 2008) based on Gauss-Hermite quadrature. 1457

Diagonalization. It is clear from (19-20) and (29-30) that all 1458 parameter-dependence in our integrand is in terms of the linear 1459 predictor $\mathbf{y} = X\mathbf{w}$. That is, we are dealing with the integral of 1460 the form 1461

$$F = \int d\mathbf{w}' \,\mathcal{N}(\mathbf{w}'|\hat{\mathbf{w}}', C) \cdot f(X\mathbf{w}'), \qquad (52) \quad {}_{1462}$$

(47)

1429

where *C* is the covariance matrix, and $X = \bigoplus_{j=1}^{k} \mathbf{g'}_{j}^{\top}$ is a fixed matrix constructed from direct sum of *k* vectors. It helps to work in a diagonalized coordinate system, so that we can separate out the relevant dimensions of \mathbf{w} . We use the singular value decomposition of the design matrix ($X = UGV^{\top}$ with U = I and $V = Q^{\top}$). Because of the direct-sum construction, XX^{\top} is already diagonal, and the left singular matrix is always *I* in this case. Then

$$G = XQ^{\top} = \begin{bmatrix} G_k & G_q \end{bmatrix},$$
(53)

where G_k is a $k \times k$ diagonal matrix and G_q is a $k \times (q - k)$ matrix of zeros. We can now denote $\mathbf{w}_k = (w_1, \dots, w_k)$ and $\mathbf{w}_q = (w_{k+1}, \dots, w_q)$ in the diagonalized variable $\mathbf{w} = Q\mathbf{w}'$, such that

¹⁴⁷⁵
$$\mathbf{w} = [\mathbf{w}_k, \mathbf{w}_q]^\top, \quad G\mathbf{w} = G_k \mathbf{w}_k = (g_1 w_1, g_2 w_2, \cdots , g_k w_k).$$

1476 *Marginalization*. Now we have

1477
$$F = \int d\mathbf{w} \, \mathcal{N}(\mathbf{w} | \hat{\mathbf{w}}, B^{-1}) \cdot f(G\mathbf{w}), \qquad B^{-1} = QCQ^{\top}$$
(54)

where *B* is the inverse of the *new* covariance matrix after diagonal-ization. If we block-decompose this matrix,

$$B = \begin{bmatrix} B_{kk} & B_{kq} \\ B_{qk} & B_{qq} \end{bmatrix}, \qquad B_{kq} = (B_{qk})^{\top}, \qquad (55)$$

the Gaussian distribution is also decomposed as

1482
$$\mathcal{N}(\mathbf{w}|\hat{\mathbf{w}}, B^{-1}) = \mathcal{N}(\mathbf{w}_k|\hat{\mathbf{w}}_k, B_*^{-1}) \cdot \mathcal{N}(\mathbf{w}_q|(\hat{\mathbf{w}}_q - \mathbf{b}), B_{qq}^{-1})$$

where $\mathbf{b} = B_{qq}^{-1}B_{qk}\mathbf{w}_k$ and $B_* = B_{kk} - B_{kq}B_{qq}^{-1}B_{qk}$. As the non-parallel part \mathbf{w}_q is integrated out, we have marginalized the integral. It is useful to recall that if a variable $\mathbf{w} \sim \mathcal{N}(\hat{\mathbf{w}}, C)$ is Gaussian distributed, its linear transform $\mathbf{y} = X\mathbf{w}$ is also Gaussian distributed as $\mathbf{y} \sim \mathcal{N}(\hat{\mathbf{y}}, \Sigma)$, with $\hat{\mathbf{y}} = X\hat{\mathbf{w}}$ and $\Sigma = XCX^{\top}$. Changing the integration variable to $\mathbf{y} = G_k\mathbf{w}_k$ is then straightforward:

$$F = \int d\mathbf{w}_k \, \mathcal{N}(\mathbf{w}_k | \hat{\mathbf{w}}_k, B_*^{-1}) \cdot f(G_k \mathbf{w}_k)$$

$$= \int d\mathbf{y} \, \mathcal{N}(\mathbf{y} | \hat{\mathbf{y}}, \Sigma) \cdot f(\mathbf{y}), \qquad \Sigma = G_k B_*^{-1} G_k^{\top}. \quad (56)$$

Standardization. Finally, in order to deal with the numerical integration, it is convenient to have the normal distribution standardized. We can use the Cholesky decomposition for the covariance matrix,

$$LL^{\top} = \Sigma_{t+1}, \tag{57} \quad {}_{1497}$$

such that the new variable $\theta = L^{-1}(\mathbf{y} - \hat{\mathbf{y}}_{t+1})$ is standard normal distributed. From the above formulation, L can be written directly in terms of the Cholesky decomposition of B_* : 1500

$$L = G_k R^{-1}$$
 where $R^{\top} R = B_*$. (58) 150

Importantly, with this transformation, each dimension of θ is independently and identically distributed. The objective function to be evaluated is now 1504

$$F(\mathbf{x}) = \int d\mathbf{y} \cdot \mathcal{N}(\mathbf{y}|\hat{\mathbf{y}}_{t+1}, \Sigma_{t+1}) \cdot f(\mathbf{y}, \mathbf{x})$$
1505

$$= \int d\boldsymbol{\theta} \cdot \mathcal{N}(\boldsymbol{\theta}|\mathbf{0}, I) \cdot f(\phi(\boldsymbol{\theta}), \mathbf{x})$$
¹⁵⁰⁶
¹⁵⁰⁷
¹⁵⁰⁸

where $\phi(\theta) = \hat{\mathbf{y}}_{t+1} + L\theta$. Once the integration is standardized this ¹⁵⁰⁸ way, there are a number of efficient numerical methods that can be ¹⁵⁰⁹ applied. ¹⁵¹⁰