Nonequilibrium dynamics of adaptation in sensory systems

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Adaptation is used by biological sensory systems to respond to a wide range of environmental signals, by adapting their response properties to the statistics of the stimulus in order to maximize information transmission. We derive rules of optimal adaptation to changes in the mean and variance of a continuous stimulus in terms of Bayesian filters and map them onto stochastic equations that couple the state of the environment to an internal variable controlling the response function. We calculate numerical and exact results for the speed and accuracy of adaptation and its impact on information transmission. We find that, in the regime of efficient adaptation, the speed of adaptation scales sublinearly with the rate of change of the environment. Finally, we exploit the mathematical equivalence between adaptation and stochastic thermodynamics to quantitatively relate adaptation to the irreversibility of the adaptation time course, defined by the rate of entropy production. Our results suggest a means to empirically quantify adaptation in a model-free and nonparametric way.

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I. INTRODUCTION

To make informed decisions, biological organisms sense and internally represent their environment using sensory systems, often with accuracy approaching physical limits [1,2]. The response function of a sensory system maps input stimuli onto output signals, whose nature depends on the encoding system. For example, cells respond to environmental stimuli through biochemical signaling [3–5]. The fate of cells in development is controlled by patterns of gene expression [6]. Neural systems encode information using spikes, voltage differences, and ionic currents [7–9].

A common challenge posed to these sensory devices is the breadth of variation of external stimuli, which is often much larger than the response range of the sensory system. The activity of photoreceptors in the retina saturate over two orders of magnitude, yet they must cope with light intensities spanning ten decades [10]. E. coli can navigate gradients of chemoattractant concentrations over five orders of magnitude, using the binary response of its rotary motors [11]. This is possible because over short timescales, stimuli are typically restricted to a much narrower distribution which depends on the immediate surrounding environment, e.g., ambient light level in vision or location concentration in chemical sensing. To produce efficient responses, sensory systems must adapt their response properties, as changes in the environment modify the statistical properties of the stimulus [9,12].

Theories have been developed to understand general principles of sensory adaptation and predict response properties, in particular in the context of sensory neuroscience [13]. One central idea is Barlow’s efficient coding hypothesis [14,15], which posits that neural systems maximize information transmission under the constraints of metabolic costs, dynamic range, and internal noise [16–19]. Efficient coding predicts that the dynamic range of the response function should be matched to the distribution of inputs so as to make the output distribution as balanced as possible, thus maximizing information [20,21]. As the distribution of input signals changes with time, the response function should then adapt accordingly. This argument rationalizes adaptation and makes specific predictions about its properties, with successful applications in vision [13,22–24]. However, progress towards a general theory is hindered by the multiplicity of systems which differ in their constraints, costs, and relevant features to be encoded [25,26].

A possible overarching principle of sensory adaptation lies in its analogy to nonequilibrium statistical mechanics. When a thermal system is driven out of equilibrium by an external forcing, its energy landscape evolves and the system “adapts” by following the new equilibrium with a delay, while losing heat to the reservoir. The resulting dynamics is irreversible and the dissipated work can be estimated using the measure of entropy production, which quantifies irreversibility [27]. Similarly, sensory adaptation creates irreversible dynamics that carries an intrinsic energetic cost, as was studied in the case the E. coli chemotactic signaling network [28]. The main result of this paper is to formalize the link between adaptation and irreversibility by studying in detail analytically solvable systems of sensory adaptation, where the mean or the variance of the stimulus changes over time.

We first derive the dynamics of optimal adaptation from principles of Bayesian inference (Sec. II). While previous approaches have used Bayesian estimates to explain adaptation [24,26,29,30], their impact on the response function either was not made explicit or was obtained by minimizing a loss function reflecting particular coding constraints. By contrast, we derive the optimal response function from the maximization of Shannon’s mutual information [19], assuming a noiseless internal adaptation variable. A previous limitation of Bayesian update rules is that they usually
assume discrete time, while both the stimulus and response are continuous. We show that in the case of a varying mean, the adaptation dynamics has a well-defined continuous-time limit, which can be mapped onto a system of coupled Langevin equations with nonlinear forces and position-dependent diffusivities, allowing for an explicit analogy with nonequilibrium statistical mechanics (Sec. III). We derive this mapping for the classical case of a stimulus mean switching between two values, but also for the case of a stimulus whose mean follows a random walk, which admits an analytical solution. In that continuous limit, we obtain explicit expressions for two quantities that characterize adaptation, accuracy and speed, which are in a trade-off relationship (Sec. IV). To relax the assumption of optimality, we also explore performance in the case where the dynamical rules of the stimulus statistics are not precisely known (Sec. V). The case of variance switching, which was treated in [29], is revisited with predictions on how information transmission should drop and recover following switching events (Sec. VI). Finally, we generalize and formalize the analogy between sensory adaptation and nonequilibrium statistical mechanism by proposing entropy production (a nonparametric measure of temporal irreversibility) as a signature of adaptation (Sec. VI). We calculate it explicitly in the case of adaptation to a changing mean and discuss its general relevance to experimental recordings of sensory neurons.

II. ADAPTATION AS INFERENCE

The first requirement to do efficient coding is to build a good estimate of the statistics of the environment, e.g., the mean and variance of the stimulus. To do so, the sensory system only has access to stimuli experienced in the past. It must integrate stimulus information far enough into the past to collect enough statistics, but not too far, since stimulus statistics themselves may change over time. This trade-off reflects the need to adapt fast, in order to track the stimulus statistics reliably, but not too fast, in order to build enough evidence that changes actually occurred and not be prone to random fluctuations of the stimulus. This task can be naturally framed as an ongoing inference problem [24,26,31,32]. Its optimal solution is given by a Bayesian formulation, which estimates the distribution of possible stimulus parameters given the stimulus history [24,25,33]. Our approach postulates that the sensory system has access to the raw stimulus, rather than just the sensory response, to implement its adaptation mechanism. In this view, the sensory system “compresses” information by transforming the input into a noisy response, but can adapt this compression scheme based on the stimulus statistics, which it has access to through (unaltered) past stimuli. This is particularly relevant for the early processing of visual information, which mostly proceeds in a feedforward way. For instance, adaptation to the mean light level occurs directly at the level of photoreceptors, with no feedback from the retinal ganglion cells (the output of the retinal), and similarly for contrast adaptation at the level of signal transduction between photoreceptors and bipolar cells [34], although some biochemical feedback internal to each cell plays an important role [35]. This view stands in contrast with adaptation mechanisms through direct feedback, whereby the response function is adapted based on the knowledge of past sensory responses only, with the input and intermediates masked [26]. We also assume that the processing linked to the adaptation module is noiseless, which is a simplification with respect to realistic biological adaptation mechanisms. While inspired by aspects of visual processing, our approach does not aim to model the details of the circuitry or biochemical pathways, but rather takes a global view of the system as an input-output relationship.

The optimal Bayesian estimator [24,29,36], also called Bayesian filter, is defined as follows. At each time step $n$, the model estimates the state of the environment given the stimulus history $P(y_n|s_{j\leq n})$, where $y_n$ represents the state of the environment at time $n$ and $s_{j\leq n}$ is the vector of past stimuli up until time $n$. In general, $y_n$ parametrizes the distribution of stimuli at each time $P(s_n|y_n)$. In this paper this distribution will be assumed to be Gaussian and $y_n$ will then simply denote its mean or variance. Bayes’s rule states that

$$P(y_n|s_{j\leq n}) = \frac{1}{\Omega} P(s_n|y_n)P(y_n|s_{j\leq n}),$$

where $\Omega = P(s_n|s_{j\leq n})$ normalizes the distribution.

Assuming that the dynamics of the environment is Markovian, one can write a recursive form

$$P(y_n|s_{j\leq n}) = \frac{1}{\Omega} P(s_n|y_n) \sum_{y_{n-1}} P(y_{n-1}|y_{n-1})P(y_{n-1}|s_{j\leq n}).$$

This formula combines the new observation $s_n$ with the estimate of $y_{n-1}$ at the previous time step [29], taking into account the way $y_n$ may have evolved. Prior knowledge of $P(y_n|y_{n-1})$ is essential as it sets the timescale over which past samples are discarded. Ignoring it, as was done in [24], leads to assuming an infinite memory timescale and everlasting dependence on initial conditions.

Armed with an estimate of the environment statistics, the sensory system may use this knowledge to adapt its response $r_n$ to the stimulus. Following previous proposals [15,21,22,37], one may assume that the stochastic encoding $P(r_n|s_n)$ is chosen to maximize information transmission

$$I(s_n, r_n) = \int d\sigma d\gamma d\gamma d\sigma P(s_n)P(r_n|s_n) \ln \left( \frac{P(r_n|s_n)}{P(r_n)} \right),$$

where the form of the response function $P(r_n|s_n)$ is set by biophysical constraints. Alternative choices of objectives to optimize include decoding accuracy, metabolic costs [38], or the ability to infer the environmental variable $y_n$ itself [26]. The simplest assumption, which is equivalent to Laughlin’s original argument [20], is to assume a constant Gaussian output noise

$$r_n = g(s_n) + \epsilon_n,$$

where $\epsilon_n$ is a uncorrelated Gaussian noise of zero mean and variance $\sigma^2$ and $g(s)$ is a function constrained between 0 and $r_{max}$ so that $P(r) \propto \exp\left\{ -[r - g(s)]^2/2\sigma^2 \right\}$. In the small noise limit $\sigma \ll \sigma_{max}$, the response function $g(s)$ maximizing the mutual information is the one that maximizes the entropy of the response [21], which is realized by the uniform distribution $P(r) = 1/r_{max}$ for $0 \leq r \leq r_{max}$, which in turns gives $P(r)dr \approx \hat{P}(s)ds$, hence $dr/ds = \hat{g}(s) = r_{max}\hat{P}(s)$, or
which is optimized to maximize information transmission based on the internal variable the new adapted phase (intermediate colors). (e) Information theory dictates that the optimal gain function should be most sensitive where the raw stimulus B the internal representation of the expected stimulus statistics (distribution of inputs), encoded by then relaxes back to a basal value as the gain function adapts to the new state. (d) Changes in the gain function can be rationalized based on the internal state of the environment given by the ratio of the output dynamic range \( r_{\text{max}} \) to the resolution of the response \( \sigma_r \). This number, which can also be interpreted as a signal-to-noise ratio, encodes the resource constraints of the sensory system and is the main determinant of information transmission. However, an interesting feature of (8) is that the information loss due to nonoptimal adaptation encoded in the Kullback-Leibler divergence does not depend on that resource parameter, making the analysis robust to that choice.

The full adaptation scheme is summarized in Fig. 1 with the example of a Gaussian stimulus whose mean switches between two values. Shortly after the mean changes [Fig. 1(a)],

equivalently
\[
g(s) = r_{\text{max}} \int_{-\infty}^{s} ds' \hat{P}(s'),
\]
so that the response function’s sensitivity follows the assumed statistics \( \hat{P} \) of the stimulus at time \( n \), derived from the belief about the state of the environment \( y \),
\[
\hat{P}(s|s_{j<n}) = \int dy P(s|y)P(y|s_{j<n}),
\]
where for the rest of this section we drop the \( n \) index to simplify notation. The mutual information being actually transmitted can be rewritten as
\[
I(s, r) = S[r] - S[r|s],
\]
where in the second term we recognize a Kullback-Leibler divergence or cross entropy, which is always non-negative. The mutual information is maximized when the inferred statistics of the environment exactly matches the true one, \( \hat{P}(s) = P(s|y) \). In that case, the Kullback-Leibler divergence is zero and only the first term survives, the channel capacity, which has an intuitive interpretation: It is the logarithm of the number of distinct responses that can be resolved from each other, given by the ratio of the output dynamic range \( r_{\text{max}} \) to the resolution of the response \( \sigma_r \). This number, which can also be interpreted as a signal-to-noise ratio, encodes the resource constraints of the sensory system and is the main determinant of information transmission. However, an interesting feature of (8) is that the information loss due to nonoptimal adaptation encoded in the Kullback-Leibler divergence does not depend on that resource parameter, making the analysis robust to that choice.

The full adaptation scheme is summarized in Fig. 1 with the example of a Gaussian stimulus whose mean switches between two values. Shortly after the mean changes [Fig. 1(a)],
the system progressively updates its expected distribution of stimuli as it accumulates more samples [Fig. 1(d)] and adapts its response function [Fig. 1(b)] according to (5), as illustrated in Fig. 1(e). The resulting response dynamics [Fig. 1(c)] shows epochs of strong upward and downward changes right after each switching transition, followed by relaxation to intermediate values as the system adapts to its new response function, closely mimicking experimental observations.

In the following we will study and solve particular cases of adaptation to a Gaussian stimulus with varying mean or variance.

III. ADAPTATION TO A VARYING MEAN

Let us start with the most intuitive and simple case: adaptation to changes in the mean of the stimulus. We will take two very simple processes as case studies: a telegraph process, in which the mean randomly switches between two values, and an Ornstein-Uhlenbeck process, in which the mean varies continuously around a fixed value. Discontinuous switches such as those produced by the telegraph process are commonly studied in experiments on sensory adaptation, because of the strong adaptive transient they trigger. The Ornstein-Uhlenbeck process is interesting because of the analytical insight it affords.

A. Two models of fluctuating mean

1. Telegraph process

We consider a system where the environmental parameter \(y_n\) switches between two values \(y_{-}\) and \(y_{+}\) with probability \(k\) at each time step. The sensory system has access only to the \(s_n\), which we assume to be Gaussian distributed with mean \(y_n\) and fixed variance \(\sigma^2\),

\[s_n = y_n + \sigma \eta_n,\]

with \(\langle \eta_n \rangle = 0\) and \(\langle \eta_n \eta_n' \rangle = \delta_{nn'}\), so that

\[P(s_n|y_n) = \frac{1}{\sqrt{2\pi\sigma^2}} e^{-(y_n-s_n)^2/2\sigma^2}.\]

We substitute this expression into (2) and use the fact that the distribution of \(y_n\) given the stimulus history \(s_{j<n}\) is entirely determined by its mean \(x_n \equiv \langle y_n \rangle |_{s_{j<n}}\) because of its binary nature. Assuming \(y_{\pm} = \pm 1\) without any loss of generality, by averaging (2) we then obtain an update equation for \(x_n\):

\[x_n = \frac{\sinh (\frac{\eta_n}{2}) + (1 - 2k)x_{n-1} \cosh (\frac{\eta_n}{2})}{\cosh (\frac{\eta_n}{2}) + (1 - 2k)x_{n-1} \sinh (\frac{\eta_n}{2})}.\]

For this estimator to be optimal, the variance \(\sigma^2\) in (11) should be equal to the true variance of the stimulus distribution. This can be achieved, for instance, by a variance adaptation mechanism, as we will see in Sec. VI. In Sec. V we will also deal with the case where the system does not use optimal parameters.

2. Autoregressive (Ornstein-Uhlenbeck) process

Another simple example of stochastic dynamics for the mean is given by an autoregressive process, which we also call Ornstein-Uhlenbeck process by abuse of language, as it reduces to it in the continuous time limit (discussed later),

\[y_{n+1} = ay_n + \sigma^2 \eta_n',\]

where \(\eta'\) is a Gaussian white noise, \(\langle \eta_n' \rangle = 0, \langle \eta_n' \eta_n'' \rangle = \delta_{nn''}\), and \(a < 1\). Again we assume that signals are distributed according to a Gaussian of mean \(y_n\) and variance \(\sigma^2\), \(s_n = y_n + \sigma \eta_n\), with \(\langle \eta_n \rangle = 0, \langle \eta_n \eta_n' \rangle = \delta_{nn'}\), and \(\langle \eta_n^2 \rangle = 0\).

Since the elements of the recursion are Gaussian, we assume a Gaussian ansatz for the posterior:

\[P(y_n|s_{j<n}) = \frac{1}{\sqrt{2\pi u_n^2}} e^{-(y_n-x_n)^2/2u_n^2}.\]

Plugging it into (2) yields recursive equations for the mean and variance of the posterior of \(y_n\), which are equivalent to a Kalman filter [39]:

\[x_n = \frac{a \sigma^2 x_{n-1} + \left[ \sigma^2 + a^2 u_{n-1}^2 \right] \eta_n}{\sigma^2 + a^2 u_{n-1}^2 + \sigma^2 \eta_n^2},\]

\[u_n^2 = \frac{\sigma^2 \left[ a^2 + a^2 u_{n-1}^2 \right]}{\sigma^2 + a^2 u_{n-1}^2}.\]

The dynamics of the variance (15) does not depend on the mean or the stimulus, so it converges to a fixed point \(u^2\) at steady state.

B. Continuous-time limit

Our adaptation dynamics was derived at discrete times, using a Bayesian estimator and exploiting the Markovian property of the system. We now take the limit of continuous time to obtain stochastic differential equations. Denoting by \(\delta t\) the time step between two observations, the continuous time is defined as \(t = n\delta t\), with \(\delta t \to 0\). As we take this limit, we must scale the various parameters of the dynamics to ensure that all the relevant quantities remain finite.

For both types of environmental dynamics, the stimulus variance may scale with \(\delta t\) as

\[\sigma^2 = \frac{\theta}{\delta t}.\]

This scaling allows us to write the stochastic term in (9) as the discretization of a Gaussian white noise \(\xi(t)\): \(\eta_n = (\sqrt{\theta}/\delta t) \int_{t-\delta t}^{t} dt \xi(t),\) with \(\langle \xi(t) \rangle = 0\) and \(\langle \xi(t) \xi(t') \rangle = \delta(t-t')\). In the continuous-time limit this gives

\[s(t) = y(t) + \sqrt{\theta} \xi(t).\]

In the telegraph process, the switching probability between two infinitesimal time steps should scale with \(\delta t\). We define \(k = \delta t/2\tau\). The switching rate is \((2\tau)^{-1}\), so \(\tau\) is the correlation timescale of the process.

Similarly, a continuous Ornstein-Uhlenbeck (OU) process may be obtained from the autoregressive process (12) in the \(\delta t \to 0\) limit with the scalings \(a = 1 - \delta t/\tau\) and \(\sigma^2 = 2D\delta t\),

\[\frac{dy}{dt} = -\frac{y}{\tau} + \sqrt{2D} \xi(t),\]

where \(\xi(t)\) is a Gaussian white noise, \(\tau\) is a relaxation time, and \(D\) may be interpreted as a diffusion coefficient in stimulus space. We set it to \(D = 1/\tau\), so the variance of \(y\) is 1. With
these choices, the first two moments of $y(t)$ are the same for both the telegraph and OU processes,

$$\langle y(t) \rangle = 0, \quad \langle y(t)y(t') \rangle = e^{-|t-t'|/\tau}. \quad (19)$$

With these scalings, the adaptation dynamics are described in the $\delta t \to 0$ limit by stochastic differential equations (with Itô convention). For the telegraph process, Eq. (11) becomes

$$\frac{dx}{dt} = -\frac{x}{\tau} + \frac{1-x^2}{\theta} s = -\frac{x}{\tau} + \frac{1-x^2}{\theta} y + \frac{1-x^2}{\sqrt{\theta}} \xi. \quad (20)$$

For the OU process, Eq. (14) becomes

$$\frac{dx}{dt} = -\frac{\sqrt{1+2\pi/\theta}}{\tau} x + \frac{u^2}{\theta} s = -\frac{\sqrt{1+2\pi/\theta}}{\tau} x + \frac{u^2}{\theta} y + \frac{u^2}{\sqrt{\theta}} \xi, \quad (21)$$

where the posterior variance

$$u^2 = \text{Var}[y(t)|s(t' \leq t)] = \frac{2}{1+\sqrt{1+2\pi/\theta}} \quad (22)$$

is obtained from the fixed-point condition (15). The adaptation dynamics for the OU process thus follows an exactly solvable Gaussian process described by the coupled equations (18) and (21).

Two timescales $\tau$ and $\theta$ govern the adaptation dynamics of both processes (20) and (21): $\tau$ is the timescale over which the environment varies, while $\theta$ gives the typical timescale over which the state variable $y$ and the fluctuations of the stimulus $\sqrt{\theta} \xi$ have equal contributions: $\langle (1/T) \int_0^T \sqrt{\theta} \xi(t) \, dt \rangle^2 = \theta/T \sim \langle y^2 \rangle = 1$ for $\tau \sim \theta$. The adaptation dynamics is expected to depend crucially on the ratio between these two timescales

$$\alpha = \frac{\theta}{\tau}, \quad (23)$$

which can be viewed as an inverse signal-to-noise ratio and is the main control parameter of adaptation. A low $\alpha$ means low stimulus fluctuations and thus precise adaptation, while high $\alpha$ means high fluctuations and thus poor adaptation.

The information transmitted under the optimal scheme described in the preceding section is given by (8) with

$$\tilde{P}(s) = \frac{1}{\sqrt{2\pi} (\sigma^2 + u^2)} \exp\left[-\frac{(s-x)^2}{2(\sigma^2 + u^2)}\right] \quad (24)$$

for the OU process and

$$\tilde{P}(s) = \frac{1}{\sqrt{2\pi} \sigma^2} \left[ \frac{1}{\alpha} e^{-(s-x)^2/2\alpha^2} + 1 - \frac{x}{\alpha} e^{-(s-x)^2/2\alpha^2} \right] \quad (25)$$

for the telegraph process. In the OU case the expression for the information simplifies to

$$I(s, r) = \frac{1}{2} \ln \frac{2\pi e \max}{\sigma^2} - \frac{1}{2} \ln \left[ 1 + \frac{u^2}{\sigma^2} \right] - \frac{1}{2} \frac{(x-y)^2 - u^2}{\sigma^2 + u^2}. \quad (26)$$

Figure 2 shows numerical simulations of the optimal adaptation dynamics (20) and (21) for the two processes, as well as the expected sensory response and information rate under the assumption of optimal information transmission (3)–(6). For the telegraph process, the response shows typical adaptive behavior, with fast changes in the response following a switch, followed by a slower relaxation to the baseline. The information rate drops right after each switch and climbs back up to its maximum as the system adapts its response function to the new statistics.

IV. ACCURACY AND SPEED OF OPTIMAL ADAPTATION

To study the adaptation dynamics of (20) and (21), we focus on two fundamental properties: the speed of adaptation, measured by the typical time it takes for the system to adapt to the changing environment, and the accuracy of adaptation, measured by the discrepancy between $x$ and $y$.

A. Adaptation time

We formally define the adaptation time as the time $t = t_A$ at which the cross-correlation function

$$C(t) = \langle y(t_0) x(t_0 + t) \rangle \quad (27)$$

reaches its maximum

$$t_A = \arg\max_t C(t). \quad (28)$$

This defines the time delay after which the adaptation variable $x$ is maximally aligned with the state variable $y$, i.e., the time it takes for $x$ to “catch up” with $y$.

We first compute this adaptation time numerically (Fig. 3) for both processes and for different values of the control parameter $\alpha$, using the Euler-Maruyama method [40]. In general, the adaptation timescale $t_A$ grows with the timescale of switching $\tau$. This suggests that sensory systems should modify their dynamics of adaptation as a function of what their expected rate of change is. Such meta-adaptation of the relaxation timescale has indeed been observed in the context of mean adaptation to the mean [24].

Two broad regimes can be distinguished. In the low signal-to-noise regime $\alpha \gtrsim 1$, $t_A$ scales linearly with $\tau$: The optimal adaptation timescale is proportional to the timescale of the environment. However, in the well-adapted regime $\alpha \ll 1$, this scaling breaks down and adaptation happens much faster than the rate of change of the environment would suggest. In that regime, it is the reliability of the observed signals that drives how fast the system adapts, so small values of $\theta$ lead to fast adaptation.

To gain analytical insight into these different scalings, we first consider the OU process, for which the cross correlation can be calculated analytically

$$C(t) = e^{-t/\tau} \left[ 1 - \frac{2}{1+\sqrt{1+2/\alpha}} e^{-(\sqrt{1+2/\alpha} - 1)t/\tau} \right], \quad (29)$$

yielding

$$t_A = \frac{\alpha(1+\sqrt{1+2/\alpha})}{2} \ln \left( \frac{2\sqrt{1+2/\alpha}}{1+\sqrt{1+2/\alpha}} \right). \quad (30)$$

For large $\alpha$, this expression becomes at leading order $t_A \sim \tau/2$, confirming the linear scaling observed in that regime. For small $\alpha$, we get $t_A \sim \sqrt{\tau/2} \ln(2)$, meaning that the

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FIG. 2. Adaptation dynamics for two mean-varying processes: the telegraph process and the Ornstein-Uhlenbeck process. (a) and (b) The state variable $y(t)$, the mean of the stimulus, varies over time with unit variance and correlation timescale $\tau$. (c) and (d) Perceived stimulus $s(t) = y(t) + \sqrt{\theta}\xi(t)$. (e) and (f) Mean value of the state variable $x = \int dy y P(y|s(t' \leq t))$ inferred by the system from past observations. (g) and (h) Sensory response obtained with the optimal gain function $r = g(s)$, derived by optimizing information transmission. (i) and (j) Information (3) actually transmitted per time bin ($\delta t = 10^{-3}$ and $\sigma_e = 0.01$). Angular brackets denote averages over 500 repetitions of the state variable evolution. The parameter is $\alpha = 0.1$.

adaptation scale is the geometric mean of the environmental timescale and the noise timescale. Such a scaling, which was previous derived in the context of Bayesian filtering for concentration sensing [41], results from a trade-off between the requirements to integrate information about the stimulus for as long as possible ($t_A \gg \theta$), but not too long to avoid including out-of-date information ($t_A \ll \tau$).

While the adaptation dynamics for the telegraph process does not have an analytical solution, we may approximate $t_A$ by the typical time $x$ takes to cross 0 following a switch. This allows us to cast the problem as a first-passage calculation.

We start by writing the Fokker-Planck equation describing the density evolution of (20) as a function of time

$$
\frac{\partial \rho}{\partial t} = -\frac{\partial}{\partial x} \left[ \left(-\frac{x}{\tau} + \frac{1-x^2}{\theta} y \right) \rho \right] + \frac{\partial^2}{\partial x^2} \left( \frac{1-x^2}{2\theta} \rho \right),
$$

(31)

whose steady-state solution at constant $y$ reads

$$
\rho_{eq}(x|y) = \frac{N}{(1-x^2)^2} \left( \frac{1+x}{1+x^2} \right)^y \exp \left( -\frac{\alpha}{1-x^2} \right),
$$

(32)

where $N$ is a normalization constant.
In the $\alpha \ll 1$ limit, $x$ is typically close to $y$ and its difference with it is of order $\alpha$. $1 - xy = O(\alpha)$, as can be checked by solving $dP_{eq}/dx = 0$. Suppose that at the time $t = 0$ a switch from $y = +1$ to $y = -1$, $x = x_0$ is drawn from the steady-state distribution $P_{eq}(x|y = +1)$ [Eq. (32)]. We define $T(x_0)$ as the mean first-passage time of $x$ from $x_0$ to 0, with $x = +1$ acting as a reflecting boundary. This is given by

$$T(x_0) = 2\theta \int_{x_0}^{+\infty} \frac{dx}{\psi(x)} \int_{-\infty}^{x} \frac{\psi(x')}{(1 - x'^2)^2},$$

(33)

where

$$\psi(x) = \left(\frac{1 - x}{1 + x}\right) \exp\left(-\frac{\alpha}{1 - x^2}\right).$$

(34)

The adaptation time is then $t_A \approx \langle T(x_0) \rangle_{x_0 \sim P_{eq}(x|y = +1)}$.

Examining Eq. (20) in the small-$\alpha$ regime, we see that when $x$ is close to $+1$, the $1 - x^2$ terms scale with $\alpha$, meaning that $x$ has sluggish dynamics dominated by the timescale $\tau$. When $x$ finally gets away from $+1$, the dynamics accelerates according to the fast timescale $\theta$, quickly reaching zero. This motivates us to use the change of variable $x = 1 - \alpha/2v$ and expand in $\alpha$ to obtain (see the Appendix)

$$\frac{t_A}{\tau} \simeq \frac{\alpha}{4} \left[\ln^2(\alpha) + [2\gamma_\text{e} - 2 \ln(2)] \ln(\alpha) + C\right] + o(\alpha),$$

(35)

where $\gamma_\text{e}$ is Euler’s constant and $C \approx 7.59$ is a numerical constant. We checked the validity of this expression against simulation results [Fig. 3(a)].

The leading order in $\alpha$ scales as

$$t_A \sim \frac{\theta}{4} \ln^2\left(\frac{\tau}{\theta}\right).$$

(36)

This sublinear scaling with $\tau$ stands in contrast with the theoretical results of Wark et al. [24], which reported a linear one. This difference is explained by the fact that in [24], the inference system assumed a vanishing switching rate. This leads to the absence of a well-defined steady state, requiring one to evaluate the adaptation scale over simulations of finite durations, with a dependence on initial conditions.

This scaling is also much more faster with $\theta$ than the square-root scaling obtained for the OU process. This is because the OU process assumes that the mean changes all the time, instead of by discontinuous jumps, making adaptation faster.

B. Accuracy

We introduce two definitions of accuracy adapted to each case. For the telegraph process, which admits only two values for the state variable $y = \pm 1$, we define the discrepancy as the probability for the posterior to be wrong $P_{\text{wrong}} = (1 - \langle xy \rangle)/2$, long after the adaptation transient. For the OU process, whose state variable is continuous, it is more natural to define the discrepancy of adaptation as the standard error between $x$ and $y$: $\Delta^2 = \langle (y - x)^2 \rangle$ [although note that $P_{\text{wrong}} = \langle |\hat{y} - y|^2 \rangle/4$, where $\hat{y}$ is drawn from the posterior $P(\hat{y}|s_{\text{in}}) = 1/(\hat{y} - 1) + 1/\hat{y}(\hat{y} + 1)$, can also be viewed as a mean-square difference].

I. Ornstein-Uhlenbeck process

Since the OU process is Gaussian, all moments can be calculated exactly. We can rewrite $\Delta^2 = \langle (y - x)^2 \rangle = \langle y^2 \rangle + \langle x^2 \rangle - 2\langle xy \rangle$. We already know $\langle y^2 \rangle = 1$ and calculate $\langle x^2 \rangle = \langle x0 \rangle = 1 - \alpha(\sqrt{T/2} + \sqrt{1/\alpha} - 1)$, from which we get

$$\Delta^2 = \alpha(\sqrt{1 + 2/\alpha} - 1) = u^2.$$

(37)

(Note that this error is equal to the uncertainty computed by the Bayesian inference system $u^2$ consistent with its optimality.) In Fig. 4(b) we plot this result and check its validity by comparing it to numerical integration of (18) and (21). In the
FIG. 4. Accuracy of adaptation to the mean stimulus as a function of rescaled stimulus variance $\alpha = \theta/\tau$. (a) Accuracy in the telegraph process, defined as the value of the probability of being in the wrong state $P_{\text{wrong}} = (1 - \langle xy \rangle)/2$. Circles are obtained by numerical simulation and the line is the analytical approximation given by (41) for $\alpha \ll 1$. (b) Accuracy in the Ornstein-Uhlenbeck process, defined as the posterior mean-square error $\Delta^2 = \langle (y - x)^2 \rangle$. Circles are from numerical simulations and the line is the analytical prediction given by (37). In both panels the dashed line shows chance prediction ($P_{\text{wrong}} = 1/2$ and $\Delta^2 = 1$).

The comparison between numerics and this expression is shown in Fig. 5(b). The limit of perfect adaptation $\alpha \ll 1$ we get, at leading order, $\Delta^2 \sim (2\alpha)^{1/2}$.  

The expression for the accuracy allows us to compute the average information between stimulus and response (26),

$$I(s, r) \approx \langle I(s, r) \rangle = \frac{1}{2} \ln \frac{2 \pi e r_{\text{max}}^2}{\sigma^2} - \frac{1}{2} \ln \left[ 1 + \frac{u^2}{\sigma^2} \right].$$ (39)

The comparison between numerics and this expression is shown in Fig. 5(b). In the $\alpha \ll 1$ regime, information approaches channel capacity, i.e., the maximal value allowed by the output noise, $I(s, r) \approx I_{\text{max}} = \frac{1}{2} \ln[2 \pi e r_{\text{max}}^2/\sigma^2]$.

2. Telegraph process

Because $P_{\text{wrong}}$ is a steady-state property, it only depends on the control parameter $\alpha$. We computed it by simulating a telegraph process $y(t)$ with switching rate $1/2\tau$ and by integrating (20) numerically [Fig. 4(a)]. For small $\alpha$, adaptation is very accurate, while for $\alpha \gtrsim 1$ it is very poor, with $P_{\text{wrong}}$ quickly reaching chance level approximately equal to $1/2$.

In the well-adapted phase $\alpha \ll 1$ ($\tau \gg \theta$), adaptation is fast compared to the switching of the state variable. After each transition, following an adaptation transient, the system quickly reaches the steady state of (20) with fixed $y$, given by (32). While the moments of $\rho_{\text{eq}}(x|y)$ cannot be calculated analytically, its expression simplifies in the small-$\alpha$ limit. With the change of variable $x = y(1 - \alpha u)$, we obtain the distribution of $u$ at leading order in $\alpha$, $\rho_{\text{eq}}(u) \approx \frac{1}{4 \tau^2} e^{-1/2u}$, (40)

from which we deduce $\langle u \rangle \approx \frac{1}{2}$ and thus

$$P_{\text{wrong}} = \frac{1 - \langle xy \rangle}{2} = \frac{\alpha \langle u \rangle}{2} \approx \frac{\alpha}{4}.$$ (41)

FIG. 5. Average information transmission as a function of the adaptation parameter $\alpha = \theta/\tau$: difference $\Delta I$ between the mutual information (8) averaged over time $\langle I(s, r) \rangle$ and the channel capacity $I_{\text{max}} = \frac{1}{2} \ln[2 \pi e r_{\text{max}}^2/\sigma^2]$ for (a) the telegraph process and (b) the Ornstein-Uhlenbeck process. Circles are from numerical simulations and the line is the analytical prediction given by (39).
This analytic prediction agrees very well with the simulation [see Fig. 4(a)]. This error is estimated assuming that the system has adapted, so long after the adaptation transient $t \gg t_A$. However, while $P_{\text{wrong}}$ is the error long after the adaptation transient, it is not equal to the average error including during the adaptation transients. To get that average error, we can approximate the error to 1 for $0 < t \leq t_A$ and to $P_{\text{wrong}}$ for $t > t_A$. Since the average time between switches is $2\tau$, we obtain

$$P_{\text{wrong, av}} \approx \left( 1 - \frac{t_A}{2\tau} \right) P_{\text{wrong}} + \frac{t_A}{2\tau} \approx 2(\ln^2(\alpha) + (22) + 21) \log(\alpha) + C + 21.
$$

Note that this scaling of the error with $\alpha$ is smaller than for the OU process (37). In the switching process, in the $\alpha \ll 1$ limit the system adapts almost perfectly to the value of $y$ after a transient lag. By contrast, in the OU process the state variable changes constantly, causing a permanent lag in the adaptation and thus a larger error. Figure 5(b) shows how the transmitted information depends on $\alpha$, in particular how it quickly converges to the maximal transmissible information as $\alpha$ goes to 0.

V. NONOPTIMAL ADAPTATION

So far we have assumed that the sensory system knows perfectly the statistics of the environmental dynamics, including its timescale $\tau$, mean, and variance, and type of dynamics. In realistic situations, however, the system may not know the statistics of the environment precisely or may have evolved to best respond to stimuli with particular statistics (e.g., natural ones), while the experimental stimulus was designed using artificial statistics (e.g., mean switching between two values). In this section we explore the impact of maladapted (nonoptimal) adaptation dynamics, where the inference system makes wrong assumptions about the environmental dynamics.

A. Misrepresented dynamics

We first consider adaptation when the sensory systems misrepresents the nature of the dynamics. We assume that the true dynamics follows a switching mean (telegraph process), while the Bayesian system assumes an Ornstein-Uhlenbeck dynamics. This leads to integrating the OU dynamics (21), but with $y$ switching between $-1$ and $+1$ with rate $(2\tau)^{-1}$.

Such an adaptation scheme is illustrated Figs. 6(a) and 6(b). The error [in the sense of the OU process, $\Delta = \sqrt{\langle (y - x)^2 \rangle}$] may be calculated analytically and is identical to the case where $y$ would actually follow an OU process with the same time constant [Eq. (37)]. This is due to the fact that in both cases the solution to (21) can be formally written as $y(t) = \int dt' e^{-(t-t')/\tau'} (\tau'/\theta)^{1/2} s(t')$, with $\tau' = \tau / \sqrt{1 + 2/\alpha}$, and that both the telegraph and the OU process have the same first two moments (19). However, it does much worse that the optimal adaptation scheme (20), as can be seen from Fig. 6(c) and confirmed by the scaling at small $\alpha$ of the adaptation time $\alpha \ln^2 \alpha \ll \alpha^{1/2}$; see (38). The impact of this nonoptimal adaptation can also be seen, to a lesser extent, on mutual information [Fig. 6(d)].

For the same reason cited above, the adaptation timescale $t_A$ is identical to that of the OU process (30). We observe
a broad range of parameters where adaptation with the OU assumption is faster than the optimal adaptation scheme [Fig. 6(e)]. We can understand this intuitively by noting that the OU assumption assumes constant change and thus will be quicker to react to sudden changes. The flip side is that even when the mean is stable between switches, it overestimates the uncertainty and tends to conservatively adapt less well [Figs. 6(a) and 6(b)].

B. Wrong parameters

We next consider the case where the type of dynamics is correctly assumed by the Bayesian system, but its parameters are wrongly estimated. The assumed values of the two parameters are off by a factor $r$ and $l$, respectively: $\tau_{\text{assumed}} = r\tau$ and $\theta_{\text{assumed}} = l\theta$. The resulting adaptation dynamics for both processes (20) and (21) are modified by substituting these assumed values

$$\frac{dx}{dt} = -\frac{x}{r\tau} + \frac{1-x^2}{l\theta}y + \frac{1-x^2}{1/\sqrt{\beta}}\xi,$$

$$\frac{dx}{dt} = -\frac{\sqrt{1+2r/4\alpha}}{r\tau} x + \frac{u^2}{l\theta}y + \frac{u^2}{1/\sqrt{\beta}}\xi$$

for the telegraph and OU processes, respectively, with $u^2 = 2/(1+\sqrt{1+2r/4\alpha})$. On the other hand, the dynamics of $y$ [Eq. (17)] [and switching with rate (22)] for the telegraph process and Eq. (18) for the OU process] are unchanged, with the true parameters $\tau$ and $\theta$.

For the OU process the problem is still solvable and the accuracy can be calculated as

$$\Delta^2 = u^2 \frac{1+lr + \beta (l+r+l-1) + \beta^2(l-r)}{2(1+b\beta r)},$$

with $\beta = (1+2r/\alpha)^{-1/2}$, which reduces to

$$\Delta^2 \sim \frac{1+lr}{2\sqrt{lr}}(2\alpha)^{1/2} \geq (2\alpha)^{1/2}$$

in the well-adapted regime $\alpha \ll 1$. The adaptation time can similarly be computed as

$$\frac{t_{\text{A}}}{\tau} \sim \frac{\beta r}{1-\beta r} \ln \frac{2}{1+\beta r},$$

which scales as $t_{\text{A}} \sim (r\tau l/\theta r)^{1/2}$ in the $\alpha \ll 1$ regime, i.e., the geometric mean of the two assumed timescales, as in the optimal case. In the $\alpha \gg 1$ regime, the adaptation timescales like $\tau$ (also like the optimal case): $t_{\text{A}} \sim \tau[r/(r-1)]\ln[(1+r)/2]$. These results indicate that while underestimating the noise and switching period leads to faster adaptation, doing so hurts the accuracy of the inference.

For the telegraph process, the equilibrium accuracy may be computed using the same method as described earlier:

$$P_{\text{wrong}} = \frac{\alpha l}{4r}.$$  

For the adaptation time, we obtain

$$\frac{t_{\text{A}}}{\tau} \sim \frac{\alpha l^2}{2l(1-1)} \ln(1/\alpha) \quad \text{for } l > 1$$

$$\sim \alpha l^2 \Gamma(l+1) \left( \frac{l^2}{2r} \right)^{l-1} \quad \text{for } l < 1$$

$$\approx \frac{\alpha}{4} \left[ \ln^2(\alpha) + [1 + 2\gamma_e - 2\ln(2) + \ln(r)] \ln(\alpha) + O(1) \right] \quad \text{for } l = 1,$$

where $\Gamma(\alpha)$ is the Gamma function.

Underestimating the switching rate ($r < 1$) allows a slightly faster adaptation, but at the cost of a lower equilibrium accuracy. Likewise, overestimating the noise ($l > 1$) also speeds up the response, but again at the expense of accuracy.

VI. ADAPTATION TO VARYING VARIANCE

A. Optimal adaptation dynamics

We now turn to the case of variance switching, where the sensory system tries to evaluate the variance of a random stimulus of fixed mean. The variance follows a telegraph process, alternating a between high-value $\sigma_h$ and a low-value $\sigma_l$. There are many examples of this type of adaptation [7,34,43,44] and it has been much studied experimentally [18,22–24,29,45]. As in the case of adaptation to the mean, we start with discrete time. At each time step, the variance switches with probability $k$. This defines a correlation time $\tau = 1/\ln(1-2k)$ of the underlying telegraph process. The stimulus is $s_n = s_0 + y_n \eta_n$, where $y_n$ is the varying stimulus (equal to $\sigma_h$ or $\sigma_l$) and $\eta_n$ is normally distributed ($\langle \eta_n \rangle = 0$ and $\langle \eta_n \eta_n \rangle = \delta_{nn}$). We set $s_0 = 0$, $\sigma_l = 1$, and $\sigma_h = r$ without loss of generality. The posterior probability of being in the low-variance phase given previously observed stimuli, $P_n = P(s_n | s_{n-1} \neq a)$, is then given by the recursive relation obtained from (2):

$$P_n = \left[ 1 + \frac{kP_{n-1}}{r(1-k)}(1-P_{n-1}) \right]^{-1}.$$  

A major difference with the case of adaptation to the mean is that this equation does not admit a well-defined continuous-time limit. Intuitively, this is because learning the amplitude of white noise is much faster than learning its mean. During any observation time $\Delta t$, the observer has access to an infinite number of independent samples. However, each sample is infinitely noisy. For the mean, these two infinities compensate exactly, but for the variance the high value of the noise is not an issue since the goal is to estimate its magnitude. As a result, in the continuous-time limit, the system can adapt instantly to the variance, as it receives an infinite amount of signal. A solution to this problem could be to replace the white noise in the stimulus by a colored noise with a finite correlation timescale $\tau_s$ so that the number of independent samples during
time $\Delta t$ scales as $\Delta t/\tau_s$. However, keeping with discrete time has a similar effect with $\tau_s = \delta t$.

### B. Speed and accuracy of variance adaptation

The dynamics of optimal adaptation can be studied by simulating (52) numerically. Figure 7 shows an example of input signal, inferred variance, optimal output response, and information transmission as a function of time.

The dynamics is characterized by an asymmetry in the adaptation to the two states [Fig. 7(b)]. This is due to the fact that the high-variance state is able to produce signals of low amplitude with high probability, while the low-variance state is very unlikely to produce large amplitudes. When in the low-variance state, the system receives few misleading signals, while in the high-variance state, frequent low-amplitude signals confound the posterior. The flip side is that the system adapts faster to switches towards the high-variance state, as already noted [29], because that state produces unlikely signals under the low-variance hypothesis. This asymmetric behavior is reflected in the dynamics of information transmission, which experiences a stronger drop after a low-to-high transition than after a high-to-low transition [Fig. 7(d)]. Notably, these predictions agree with experiments in the fly visual system, where this asymmetry in the loss of information transmission following variance switches was observed [23].

As in the case of adaptation to the mean, we can define the accuracy of adaptation as the probability that the posterior is wrong and the adaptation time $t_A$ as the delay $n$ maximizing $\langle y_n P_{y_n+y_n} \rangle$. In Fig. 8 we plot these two quantities, as well as the transmitted information, given by (8), as a function of the two parameters of the model: the switching rate $\tau$ and the variance ratio $r = \sigma_h/\sigma_l$. The error decreases with the switching period $\tau$, as the system has more time to adapt, as well as with the variance ratio $r$. The trend here depends also on the value of $r$. The adaptation time varies sublinearly with $\tau$. It is approximately linear at small $\tau$ and levels off without saturating at larger $\tau$. The larger the variance ratio $r$, the narrower the linear region. The loss in information transmission due to imperfect adaptation $\Delta I$ follows the same dependence on $\tau$ as the error. However, its dependence with respect to the variance ratio $r$ is reversed. The higher that ratio, the easier it is to distinguish between the two stimulus statistics, but errors are more costly in terms of information transmission because of the larger discrepancy between the assumed and actual dynamic ranges. Conversely, smaller differences between the two environments lead to less inefficiency in their encoding, despite a higher error in distinguishing them.
Experiments in the fly [23] and vertebrate [24] visual system suggest a linear relationship between the adaptation time and the switching timescale, although this linear assumption was not compared against alternative scalings. Our theory predicts that such a linear regime is only expected when the switching period is relatively short compared to the typical time scale of the adaptation variable $y(t)$. In this type of dynamics, the irreversible production of entropy was defined in terms of heat loss and dissipation and shown to be equal to (53) in the case of Markovian thermodynamics. For a Markov system, one has $\delta S^{\text{tot}}(\Omega) = \sum_{z,\zeta} w_{z\zeta} p_{z} \ln \frac{w_{z\zeta}}{w_{\zeta\zeta}}$, where $w_{z\zeta}$ is the transition probability from $z$ to $\zeta$, so at steady state the average entropy production reads

$$\frac{1}{N} \sum_{z,\zeta}^{N \to \infty} w_{z\zeta} p_{z} \ln \frac{w_{z\zeta}}{w_{\zeta\zeta}}.$$

where $p_{z}$ is the steady-state distribution of $z$ defined by the implicit equation $\sum_{\zeta} w_{z\zeta} p_{\zeta} = 1$. Taking the continuous limit, we can define an entropy production rate $\dot{S} = \frac{\delta S}{\delta t}$ as $\delta t \to 0$.

When the statistics of stimuli do not vary and are reversible in time, no adaptation occurs in an optimal estimator and we expect the temporal statistics of the response to be temporally reversible. When the statistics of stimuli change (abruptly or continuously), the response statistics are transiently poorly adapted and require time to relax to their efficient encoding state, similar to the relaxation of the equilibrium of a thermodynamic system after a change in an external control parameter (temperature, displacement, force, etc.), thus leading to the production of entropy.

The measure of adaptation through entropy production provides a means to detect and quantify adaptation in any sensory system, in a parameter-free manner and without having to know the fine details of the encoding strategy. In the following

![Graphs](attachment:entropy_production.png)

FIG. 8. Accuracy and speed of adaptation to variance switching. (a) Accuracy of the optimal adaptation dynamics for three different values of the variance ratio $r = \sigma_{l}/\sigma_{r}$, as a function of the inverse of the switching timescale $\tau$, which plays a similar role to $\alpha$ in mean switching. The lower the $r$, the larger the estimation error. (b) Adaptation time as a function of the switching timescale $\tau$. The dependence is linear for very short times and then displays strong sublinear behavior. (c) Information transmission as a function of $1/\tau$. Note that while a large ratio between the two variances induces a lower error in (a) (because they are easier to distinguish), this error has a larger impact on information transmission, so that information loss is overall larger.
we quantify the production of entropy in the models of mean and variance adaptation studied above.

B. Adaptation to mean

We start with the OU process, for which entropy production may be calculated analytically (see Fig. 9). We start with the Gaussian transition probabilities, with \( z' = (x(t), y(t)) \) and \( z = (x(t + \delta t), y(t + \delta t)) \),

\[
\begin{align*}
    P(y_{t+\delta t} | y_t) & = P(x_{t+\delta t} | x_t, y_t) \\
    & = \frac{1}{\sqrt{4\pi \Delta}} \exp \left( - \frac{(y_{t+\delta t} - y_t)^2}{4\Delta} \right), \\
    P(x_{t+\delta t} | x_t, y_t) & = \frac{1}{\sqrt{4\pi \Delta}} \exp \left( - \frac{(x_{t+\delta t} - x_t)^2}{4\Delta} \right),
\end{align*}
\]

with \( \Delta_t = u^2 / 2\gamma \). Plugging these expressions into (54) gives

\[
\delta S = \frac{\sqrt{1 + 2/\alpha} - 1}{C(\delta t)} - \frac{1}{C(\delta t)},
\]

where we recall \( C(t) = (y(t_0)x(t_0 + t)) \). Expanding at small \( \delta t \) yields the entropy production rate

\[
\dot{S} = \frac{1}{2\tau} [(U_+(x)|_{y = \rho(x/y+1)} - U_-(x)|_{y = \rho(x/y-1)}],
\]

which again diverges for \( \alpha \to 0 \), but this time logarithmically. In the \( \alpha \to \infty \) limit, the system never adapts and the dynamics is in effective equilibrium \( \dot{S} \to 0 \).

C. Variance switching

Since we have no analytical solution or continuous-time limit for the case of a switching variance, we take recourse to numerical estimates of (54). The results, shown in Fig. 10, express a picture similar to that for the mean switching case. When the switching period is large, the system is well adapted, spending most of its time in the equilibrium state, but dissipating heat during each adaptation transient. The average amount of entropy production per switching cycle does not generally saturate at large \( \tau \) and also grows with the variance ratio \( r \), as expected.

Taken together, these results suggest that entropy production provides a good signature of adaptation. It is larger when adaptation is more precise and when the environmental changes causing adaptation are more important.

During the adaptation transient, the system relaxes to its new equilibrium, lowering its energy by dissipation. The heat thus dissipated is exactly equal to the total entropy produced during this transient. Assuming that the system has completely adapted to the previous epoch prior to each switch (which is valid for \( \alpha \ll 1 \)), we may write the entropy production rate as the mean entropy production per switch, divided by the average time between switches:

\[
\dot{S} = \frac{1}{2\tau} [(U_+(x)|_{y = \rho(x/y+1)} - U_-(x)|_{y = \rho(x/y-1)}],
\]

\[
\dot{S} = \frac{1}{2\tau} \ln \frac{1+x}{1-x} \rho(x/y+1).
\]
that they both depend on a single parameter computed the speed and accuracy of adaptation and showed differential equation, which we can solve analytically. We reduces in the continuous-time limit to coupled stochastic dynamics.

response of the adaptation mechanism, leading to irreversible lay between changes in the statistics of the stimulus and the mathematical sense. This correspondence stems from the de-
plicit analogy to nonequilibrium systems allowed us to relate their study in terms of stochastic thermodynamics. This ex-
the state of the environment in a noisy manner, allowing for
ous changes to the stimulus statistics. The resulting Bayesian simple solvable models of adaptation to sudden or continu-
terms of Bayesian filters. In this work we have focused on
When

adaptation have reported linear scalings in adaptation to both

methods. For instance, Hopfield networks [53] encode information collectively through their interactions or two-photon calcium imaging techniques [51,52]. Adaptation has mostly been studied at the level of single neurons, because of its evident manifestation in terms

adaptation always scales sublinearly with the environmental timescale. However, the precise scaling depends of the precise model of the environment: logarithmic for abrupt changes in the stimulus mean (telegraph process) and square root for continuous ones (Ornstein-Uhlenbeck process). Since these are two extremes of one-dimensional stochastic process, we expect other types of environmental dynamics to have scalings that fall within that range. Most experiments studying visual adaptation have reported linear scalings in adaptation to both the mean and variance [23,24], although alternative scaling laws were not tested. In our model, a strict linear scaling is approached only in the limit of low fidelity, where adaptation is very poor. This suggests that these sensory systems are actually not optimal for these simplified stimulus statistics, but have instead evolved under other constraints, such as metabolic or biophysical constraints, and for efficiency under a broad range of complex stimulus contexts.

Our results on mean switching mostly hold true in the case of variance switching, including the sublinear scaling of adaptation time with environmental timescale. In that case, the continuous-time limit is not straightforward. We thus focused on the dynamics in discrete time using numerical simulations. We found an asymmetry in the adaptation dynamics, with a larger drop in information transmission following an increase of variance than a decrease, consistent with experimental ob-
servations in the fly visual system [23].

Optimality is only a guiding principle and not an assumption for what the system does. We also studied the effect of optimizing response properties when stimulus statistics are misrepresented. We found that while assuming continuous change is more conservative, the corresponding adaptation strategy fares just as well when the actual dynamics is discontinuous. Mischaracterizing the parameters of the environmental statistics affects the precision and speed of adaptation, tipping the balance between the two, but not their scaling with \( \alpha \).

To explore the impact of adaptation on the coding strategy, we assumed that the system changes its response function to optimize the expected information transfer. This relies on the implicit assumption that the sensory system also outputs the adaptation variable \( x \) in addition to the response \( r \). However, realistic coding strategies may combine the two together into a single response, as, for instance, in the case of variance coding, where the response encodes both stimulus fluctuations and its variance through the mean response [23]. How to efficiently compress these different signals into a single noisy response remains to be studied.

A main contribution of this paper is a mathematically pre-
cise link between adaptation and dissipation. This analogy emerges from the form of the dynamical Bayesian equa-
tions (20) and (21), where adaptation resembles relaxation to equilibrium. We showed that a high degree of adaptation, as determined by its speed and accuracy, can be quantified by the degree of deviation from equilibrium and reversibility. This measure of adaptation through entropy production could give a systematic way to detect and quantify adaptation in any sensory system, in a parameter-free manner and without having to know the fine details of the encoding strategy. In particular, it could be applied to recordings of populations of neurons in a variety of sensory systems in response to changes in the stimulus statistics, using increasingly available multielectrode recordings [50] or two-photon calcium imaging techniques [51,52]. Adaptation has mostly been studied at the level of single neurons, because of its evident manifestation in terms of spiking rates of individual neurons. However, more subtle adaptive changes could occur in the way multiple neurons encode information collectively through their interactions or shared variability, which would be missed by traditional meth-
ods. For instance, Hopfield networks [53] encode information in the collective state of many neurons, and it has been argued

![FIG. 10. Entropy production in adaptation to variance switching. The average entropy production rate \( \delta S/\tau \) is plotted as a function of the inverse correlation time for three different values of the variance ratio \( r = \sigma_n/\sigma_r \). Note that the entropy production is rescaled by \( 1/\tau \).](image-url)
that similar principles may govern the encoding of visual information in the retina [54]. In that case both the output $x$ and the adaptation variable $y$ could be implicitly encoded in the collective activity. The signatures of entropy production proposed here could be useful for detecting adaptation in such combinatorial codes.

Other sources of irreversibility, including in the stimulus statistics themselves or through the inherent irreversibility of the biophysical processes that implement the response, may confound this analysis and would have to be corrected for. Still, the relation between adaptation, time, and accuracy provides a testable hypothesis that could be explored in future experiments. It should also be emphasized that this dissipation may not directly correspond to actual energy consumption in the system, although it always provides a lower bound. In neural systems, energy consumption from electrical activity far outweighs dissipation estimated by the irreversibility of measurable quantities. On the other hand, the two may be surprisingly close for molecular systems. Previous work by Lan et al. [28] used entropy production to estimate energy consumption in the context of adaptation in E. coli chemotaxis, showing that the system realizes a near-optimal trade-off between energy, speed, and accuracy. By contrast, our proposal is to use entropy production as an intrinsic definition of adaptation in sensory systems, independently of energetic considerations.

Our framework could be expanded to account for meta-adaptation [55], which describes the possibility that the system dynamically learns the hyperparameters of the stimulus statistics (e.g., $\theta$, $\tau$, and $\sigma$) from the past stimulus. A similar form of meta-adaptation has been proposed and explored in the context of evolutionary adaptation [56], where microbial populations are assumed to adapt their composition not only to environmental changes, but also to the statistics with which that environment changes, using a strategy of Bayesian filtering similar to that employed here. Combining all hyperparameters into a collective variable $\Theta$, we could write a recursive equation similar to (2) assuming a Markovian dynamics for $\Theta$:

$$P(y_n, \Theta_n|s_{j\leq n}) = \frac{1}{\Omega} P(s_n|y_n) \left[ \sum_{y_{n-1}, \Theta_{n-1}} P(y_n|y_{n-1}, \Theta_{n-1}) \times P(k_n|\Theta_{n-1})P(y_{n-1}, \Theta_{n-1}|s_{j<n}) \right].$$

(A3)

In this scheme the sensory system learns both the parameters of the stimulus statistics and the hyperparameters that govern their own dynamics. Such meta-adaptation is needed to explain how sensory systems dynamically adapt their adaptation speed to the environmental timescale [23,24], on a longer timescale than adaptation itself. This hypothesis is consistent with the fact that changes in the world occur on many timescales [57] and is also suggested by the wide range of timescales in visual adaptation even at the level of the retina, from seconds to hours. Future experimental and theoretical work should determine the relevance of this theory in biological sensory systems.

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APPENDIX: MEAN FIRST-PASSAGE TIME AFTER A SWITCH

We start with (33) to operate the change of variables $x = 1-\alpha/2w$ and $x' = 1-\alpha/2v$. In the limit $\alpha \ll 1$, we assume that $w$ and $v$ are small for the dominating contribution of the integral. Because of the $1-x^2$ prefactor in the attraction term towards $y$, the dynamics of $x$ is initially “stuck” in the previous belief, $x \sim 1$. Once it reaches a value $1-O(1)$, it quickly (in time approximately equal to $\theta$) reaches 0, before continuing on to its next state approximately equal to $-1$.

The change of variable gives

$$T(x_0) \sim \frac{\theta}{2} \int_{\alpha/2}^{w_0} dw \ e^{e^{w_2}/(w-\alpha/4)} \frac{w - \frac{\alpha}{4}}{w^3} \times \int_{w}^{+\infty} dv \ e^{e^{v_2}/(v-\alpha/4)} \frac{v^2}{(v - \frac{\alpha}{4})^3},$$

(A1)

where $w_0 = (\alpha/2)/(1-x_0)$ is of $O(1)$ in $\alpha$. The internal integral in (A1) can be rewritten as the sum of three parts

$$\int_{w}^{+\infty} dv \ e^{-v} \frac{v^2}{(v - \frac{\alpha}{4})^3} \left[ \exp \left( -\frac{\alpha}{4} \frac{1}{v - \frac{\alpha}{4}} \right) - 1 \right].$$

(A2)

with $R(v) = v^2/(v - \frac{\alpha}{4})^3 - 1/v$.

The first term on the right-hand side in (A2) is by definition equal to the special function $E_1(w) \approx w - \gamma_e - \ln(w) + O(w)$ (related to exponential integral). Its contribution to (A1) is dominated by the behavior at small $w \to \alpha/2$ where the integral diverges when $\alpha \to 0$. As we will see, it also dominates the second and third terms, yielding

$$T(x_0) \approx \frac{\theta}{2} \int_{\alpha/2}^{w_0} dw \ \ln(1/w) - \frac{\alpha}{4} \frac{w - \frac{\alpha}{2}}{w^2}$$

$$\approx \frac{\theta}{4} \ln^2(\alpha) + [2\gamma_e - 2\ln(2) + 1] \ln(\alpha) + O(1).$$

(A3)

What remains to show is that the second and third terms in (A2) contribute at most $O(1)$ to the result. The second term’s contribution to (A1) is bounded by

$$\frac{\theta}{2} \int_{\alpha/2}^{w_0} dw \left( w - \frac{\alpha}{4}\right) \left[ \frac{8w}{3} - 3 + \ln \frac{2w}{\alpha} \frac{1}{2(2w/\alpha - 1)^2} \right]$$

$$= \theta O(1).$$

(A4)

As for the third term, it is approximately $-\frac{\alpha}{4}$

$$\int_{w}^{+\infty} \frac{w^3}{(w - \frac{\alpha}{4})^3} e^{-w} \propto \alpha \ln(1/w)$$

and will thus contribute $O(\theta\alpha \ln(1/\alpha)) = O(\theta)$ to the result.
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