

Estimating fMRI Timescale Maps

Gabriel Riegner

Halicioğlu Data Science Institute, University of California San Diego

Samuel Davenport

Division of Biostatistics, University of California San Diego

Bradley Voytek

Halicioğlu Data Science Institute, University of California San Diego
Department of Cognitive Science, University of California San Diego
Neurosciences Graduate Program, University of California San Diego

Armin Schwartzman

Halicioğlu Data Science Institute, University of California San Diego
Division of Biostatistics, University of California San Diego

Abstract

Brain activity unfolds over hierarchical timescales that reflect how brain regions integrate and process information, linking functional and structural organization. While timescale studies are prevalent, existing estimation methods rely on the restrictive assumption of exponentially decaying autocorrelation and only provide point estimates without standard errors, limiting statistical inference. In this paper, we formalize and evaluate two methods for mapping timescales: a time-domain fit of an autoregressive (AR1) model and an autocorrelation-domain fit of an exponential decay model. The assumptions are generalized to include all stationary and ergodic processes by incorporating robust standard errors that account for non-exponential autocorrelation decay. We introduce theoretical properties of timescale estimators and show parameter recovery in realistic simulations, as well as applications to resting-state fMRI from the Human Connectome Project. Comparatively, the time-domain method produces more accurate estimates under model misspecification, remains computationally efficient for high-dimensional fMRI data, and yields maps aligned with known functional brain organization. In this work we show valid statistical inference on fMRI timescale maps, and provide python implementations of all methods.

1 Introduction

1.1 Scientific Motivation

Neural processes spans multiple timescales, from millisecond synaptic events to slower activity coordinating distributed brain networks [2]. Multimodal evidence links these differences in timescales to intrinsic brain organization that reveal how regions integrate and process information over time. Timescale maps of the brain align with functional hierarchy – sensory areas that process rapidly changing stimuli show faster timescales than association areas involved in cognitive processes that unfold over longer durations [28, 6, 9, 24, 34]. This hierarchy is also associated with anatomical organization including myelination levels and gene expression patterns, as shown in studies using human electrophysiology, MEG, and gene expression profiling [6, 32].

Computational modeling by Li and Wang [15] suggests that hierarchical timescales emerge from: (1) brain-wide gradients in synaptic excitation strength, (2) electrophysiological differences between excitatory and inhibitory neurons, and (3) balance between distant excitatory and local inhibitory inputs. In addition to these intrinsic mechanisms, there is growing evidence that neuronal timescales are dynamic and modulated by experimental manipulations or behavioral demands. For example, pharmacological agents like propofol and serotonergic drugs alter intrinsic timescales, affecting the temporal integration of information in the brain [12, 33]. Timescale changes have also been observed during development, sleep deprivation, wakefulness, neuropsychiatric disorders (autism and schizophrenia), and naturalistic behaviors [21, 22, 38, 39, 20]. These findings demonstrate that timescales are broadly relevant to both structural and functional properties of the brain.

Seminal research on timescales has primarily used invasive electrophysiology in non-human animals [24, 3, 27, 19]. While these methods provide high temporal resolution for studying neural activity at the single-neuron level, they are limited by sparse spatial sampling. Investigating the large-scale spatial organization of timescale maps requires non-invasive methods like resting-state functional MRI (rfMRI), which measures spontaneous fluctuations in the blood oxygen level-dependent (BOLD) signal. rfMRI provides full-brain coverage of hemodynamic processes at frequencies below 0.1 Hz [28, 10], offering dense spatial sampling compared to techniques like EEG or MEG. Although the BOLD signal does not directly measure neural activity, it reflects hemodynamic changes associated with underlying electrophysiological signals [16], making it a valuable tool for investigating high-resolution cortical timescale maps. Studies have shown that rfMRI-derived timescale maps align spatially with those from other imaging modalities across human and animal models [28, 31, 17]. The present study will focus on rfMRI from the Human Connectome Project dataset [36].

1.2 Methodological Motivation

Timescales are generally estimated using three main methods in the (1) time-domain, (2) autocorrelation-domain, or (3) frequency-domain. The most common is the autocorrelation domain, where timescales are defined by fitting an exponential decay model to the sample autocorrelation function (ACF) [29, 3, 13, 30, 40, 27, 37, 25, 18, 15, 31]. Similar approaches compute timescales directly from the sample ACF as the sum of positive autocorrelations [39, 19, 38], or by identifying where the sample ACF crosses a specified threshold [39, 41]. Alternatively, the time-domain method uses a first-order autoregressive (AR1) model to estimate timescales directly from time-series data [14, 22, 12, 17, 33, 31], and has shown better test-retest reliability than autocorrelation-domain methods for rfMRI [12]. Finally, frequency-domain methods use the sample power-spectral density (PSD) to estimate and remove neural oscillations, as timescales are properties of the aperiodic signal [4, 6, 20, 40, 5]. Since previous research shows that rfMRI is predominantly aperiodic with scale-free spectral properties [11, 10], the present paper focuses only on the time- and autocorrelation-domain methods.

1.3 Problem Statement and Proposed Solution

A key challenge in applied timescale research is the lack of standardized model definitions, where diverse approaches have led to inconsistent findings across studies. Many parameterization methods rely on restrictive assumptions, such as exponential autocorrelation decay, which may bias timescale estimates and (more often) their standard errors. The distributional properties of estimators are often ignored, resulting in studies reporting only point estimates without quantifying uncertainty, which limits statistical inference and hypothesis testing.

To address these issues, this paper formalizes and evaluates two commonly applied timescale methods: the time-domain fit of an autoregressive (AR1) model and the autocorrelation-domain fit of an exponential decay model. The goal is to estimate accurate timescale maps that enable robust statistical testing and inference across brain regions. (1) The assumptions are generalized to include all stationary and ergodic processes, not only those with exponential autocorrelation decay. (2) Robust standard errors account for the inevitable mismatch between the data-generating process and fitted model, enabling valid inference despite model misspecification. (3) Theoretical properties demonstrate that both time- and autocorrelation-domain methods converge to different values due to their distinct definitions, and are consistent and asymptotically normal. (4) Simulations confirm that both methods yield unbiased estimates across autoregressive and realistic settings, with standard errors that appropriately capture uncertainty even with approximating models. (5) Empirical analysis of fMRI from the Human Connectome Project reveals that both approaches produce similar t-ratio maps (timescales relative to their standard errors) which display a hierarchical organization of timescales across the brain's cortical surface. (6) Comparative analyses show that the time-domain method performs as well as, and often better than, the autocorrelation-domain method, while maintaining greater computational efficiency for high-dimensional fMRI data analysis.

The proposed methods address important limitations in neural timescale research by providing rigorous statistical methods that move beyond point estimates to incorporate uncertainty quantification. Through formal definitions, theoretical validation, and extensive testing across simulations and empirical data, we demonstrate that both time- and autocorrelation-domain estimators yield consistent standard errors under broad conditions, enabling reliable inference and hypothesis testing. This work establishes a methodological foundation for future research investigating the functional and structural organization of timescales in the brain.

2 Methods

2.1 Assumptions

Let $\{X_t, t \in \mathbb{Z}\}$ be a discrete-time stochastic process that is *weakly stationary* and *ergodic*, and let $x_t = \{x_1, x_2, \dots, x_T\}$ be a finite sample of X_t . For simplicity, assume X_t and x_t are mean zero. Stationarity implies a constant mean and variance (independent of time index t), and an autocovariance function that only depends on time lag k :

$$\gamma_k = \text{cov}[X_t, X_{t-k}] = \mathbb{E}[X_t X_{t-k}]. \quad (1)$$

For analysis we use a normalized measure of the autocovariances, the *autocorrelation function (ACF)*:

$$\rho_k = \text{corr}(X_t, X_{t-k}) = (\gamma_0)^{-1}(\gamma_k) \quad (2)$$

where γ_k is the autocovariance at lag k and γ_0 is the variance. Cauchy-Schwarz bounds $|\rho_k| \leq 1$, but stationarity alone does not guarantee decay of ρ_k with increasing lag; constant or periodic processes can maintain nonzero correlations indefinitely. Ergodicity imposes a stronger dependence constraints

than stationarity, yet still allows for a wide set of time series processes. In particular, summability of the ACF, $\sum_{k=1}^{\infty} |\rho_k| < \infty$, guarantees that ρ_k decays to zero asymptotically at a rate that is absolutely summable [8, Chapter 14.7]. Therefore, as the time separation between X_t and X_{t-k} increases, the degree of correlation decreases. These conditions allow for defining a timescale by analyzing how its autocorrelations decay.

As introduced by Murray et al. [24], the timescale τ represent the lag where exponentially decaying autocorrelations reach $1/e \approx 0.37$ (e-folding time), analogous to the time constants of many physical systems. While it provides an intuitive description of the memory or persistence of that process, assuming an exponential function imposes stricter constraints than ergodicity, which alone does not prescribe any specific type of decay (exponential, linear, damped periodic, etc.). This highlights an important distinction between the data-generating process and the simplified parametric model used to describe the timescale over which such a process becomes decorrelated. In the present paper, we adopt broad assumptions, requiring only that the process is stationary and ergodic, to account for cases where the ACF decay may not be strictly exponential. Acknowledging that the data-generating process and the fitted model will likely be different in practice, we describe standard error estimation methods that account for this mismatch, enabling valid inference despite model misspecification.

2.2 Timescale Definitions

We approximate the dominant exponential decay in autocorrelations by a single timescale parameter τ , and formally evaluate two timescale methods that are commonly applied across neuroimaging modalities (fMRI, EEG, ECoG, MEG). The time-domain linear model estimated with linear least squares [14, 22, 12, 17, 33, 31], and the autocorrelation-domain nonlinear model estimated with nonlinear least squares [29, 3, 13, 30, 40, 27, 37, 25, 18, 15, 31].

2.2.1 Time-Domain Linear Model

A first order autoregressive model (AR1) provides a linear approximation of timescale. The AR1 model:

$$X_t = \phi X_{t-1} + e_t \quad (3)$$

models as linear regression between X_t and X_{t-1} in the time domain. In the autocorrelation domain, it implies that the theoretical ACF decays exponentially at a rate determined by ϕ , such that $\rho_k = \phi^k$ [8, Chapter 14.22]. For a stationary process with $|\phi| < 1$, the exponential decay rate can be directly extrapolated from ϕ , with a timescale τ equal to the lag at which the AR1-projected ACF reaches $1/e \approx 0.37$ [$\rho_\tau = \phi^\tau = 1/e$], resulting in a change of variable [$\tau = g(\phi) = -1/\log(|\phi|)$]. The timescale τ is expressed as a nonlinear function of ϕ , denoted by $g(\phi)$. This defines τ to be a real number even though the ACF only includes integer indices, and the absolute value allows for $\phi < 0$.

Importantly, we do not assume X_t follows AR1 with *iid* errors, allowing for projections errors that may exhibit unequal variance and residual autocorrelation. Relaxing the constraints on the errors allow for AR1 approximations where deviations from AR1 are captured by the error term. Thus, this model can be applied to any stationary and ergodic process, even if the true data-generating process is not AR1, making the resulting fit an AR1 projection. The parameter ϕ^* represents the best approximation of the process X_t by an AR1 model. It is the value that minimizes the expected squared error function $S(\phi)$:

$$S(\phi) = \mathbb{E}[(X_t - \phi X_{t-1})^2], \quad \phi^* = \operatorname{argmin}_{\phi} S(\phi). \quad (4)$$

$S(\phi)$ is minimized by taking its derivative with respect to ϕ , setting it to zero, and solving for ϕ^* :

$$\frac{d}{d\phi} S(\phi) = -2\mathbb{E}[X_{t-1}(X_t - \phi X_{t-1})] = 0 \quad (5)$$

Differentiating this quadratic function yields a linear equation in ϕ , and solving this results in a closed-form expression for the optimal ϕ^* . Therefore, ϕ^* is defined by *linear projection* and the timescale parameter τ^* by a change of variable:

$$\phi^* = (\mathbb{E}[X_{t-1}^2])^{-1}(\mathbb{E}[X_t X_{t-1}]) \quad (6)$$

$$\tau^* = g(\phi^*) = -\frac{1}{\log(|\phi^*|)} \quad (7)$$

Since X_t is stationary with finite variance, the parameters ϕ^* and τ^* defined by projection are unique; in fact, any approximating AR1 model is identifiable if $\mathbb{E}[X_{t-1}^2]$ is non-negative [8, Theorem 14.28].

2.2.2 Autocorrelation-Domain Nonlinear Model

Alternatively, timescales can be defined in the autocorrelation domain by an exponential decay function, as introduced by Murray et al. [24]. For consistent notation, we write the autocorrelation-domain nonlinear model as:

$$\rho_k = \phi^k + e_k, \text{ for } k \in \{0, 1, \dots, K\}, \quad (8)$$

where ρ_k denotes the autocorrelation at lag k and e_k is the error term. The relationship between ρ_k and k is nonlinear in ϕ which determines the exponential decay rate. Unlike the Time-Domain Linear Model above, this definition captures exponential decay across multiple (K) lags of the ACF rather than by a single lag, capturing longer-range temporal dependencies. Consequently, the parameter ϕ here is not the same as the AR1 projection parameter, since both its value and interpretation differ when influenced by multiple lags.

Here, the projection parameter ϕ^* is the value that minimizes the expected squared error function $S(\phi)$:

$$S(\phi) = \mathbb{E}[(\rho_k - \phi^k)^2], \quad \phi^* = \underset{\phi}{\operatorname{argmin}} S(\phi) \quad (9)$$

$S(\phi)$ is minimized by taking its derivative with respect to ϕ , setting it to zero, and solving for ϕ :

$$\frac{d}{d\phi} S(\phi) = -2\mathbb{E}[(k\phi^{k-1})(\rho_k - \phi^k)] = 0 \quad (10)$$

However, the derivative is nonlinear in ϕ , preventing a closed-form solution for least squares minimization. Therefore, optimization methods are needed to approximate ϕ^* by *nonlinear projection*. The corresponding timescale can be expressed by a change of variable:

$$\tau^* = g(\phi^*) = -\frac{1}{\log(|\phi^*|)} \quad (11)$$

2.3 Timescale Estimation

2.3.1 Time-Domain Linear Least Squares Estimator

The linear least squares (LLS) estimator of the Time-Domain Linear Model has the following closed-form expression:

$$\hat{\phi}_{\text{LLS}} = \left(\sum_{t=2}^T x_{t-1}^2 \right)^{-1} \left(\sum_{t=2}^T x_t x_{t-1} \right) \quad (12)$$

$$\hat{\tau}_{\text{LLS}} = g(\hat{\phi}_{\text{LLS}}) = -\frac{1}{\log(|\hat{\phi}_{\text{LLS}}|)}, \quad (13)$$

where $\hat{\phi}_{\text{LLS}}$ and $\hat{\tau}_{\text{LLS}}$ are the sample versions of the population parameters from equations (6) and (7), respectively [8, Chapter 14.3].

2.3.2 Autocorrelation-Domain Nonlinear Least Squares Estimator

The nonlinear least squares (NLS) estimator of the Autocorrelation-Domain Nonlinear Model is fit to the ACF, so the time series is first transformed into the autocorrelation domain. For a finite and centered time series, the population ACF from equation (2) is estimated by:

$$\hat{\rho}_k = (\hat{\gamma}_0)^{-1}(\hat{\gamma}_k) = \left(\sum_{t=1}^T x_t^2 \right)^{-1} \left(\sum_{t=k+1}^T x_t x_{t-k} \right), \quad (14)$$

where $\hat{\gamma}_k$ is the sample covariance at lag k and $\hat{\gamma}_0$ is the sample variance. The population ACF (2) by ergodicity approaches zero as lag k increases. However, sampling variability may yield non-zero autocorrelations even when true values are zero. To mitigate this, the sample ACF estimator (14) imposes a bias towards zero by scaling the autocovariances ($\hat{\gamma}_k$, calculated using $T - k$ terms) by the total sample variance ($\hat{\gamma}_0$, calculated using all T timepoints).

By the model definition (8), the exponential decay parameter ϕ^* that minimizes the cost function, $S(\phi)$ in equation (9), is estimated by minimizing the sample analog $\hat{S}(\phi)$:

$$\hat{S}(\phi) = \frac{1}{K} \sum_{k=0}^K (\hat{\rho}_k - \phi^k)^2 \quad (15)$$

$$\hat{\phi}_{\text{NLS}}^* = \underset{\phi}{\operatorname{argmin}} \hat{S}(\phi) \quad (16)$$

$$\hat{\tau}_{\text{NLS}}^* = g(\hat{\phi}_{\text{NLS}}^*) = -\frac{1}{\log(|\hat{\phi}_{\text{NLS}}^*|)}. \quad (17)$$

In this paper we use the Levenberg-Marquart algorithm to iteratively update the estimate of $\hat{\phi}_{\text{NLS}}^*$ until convergence (i.e., when the step size goes below a 10^{-6} tolerance).

2.4 Standard Error of the Estimators

2.4.1 Time-Domain Method

We provide a standard error expression for τ^* under model misspecification. When the data-generating process is not AR1 and consequently the errors are not independent, the usual (naive) standard errors

will have a downward bias. This renders invalid confidence intervals or hypothesis tests that rely on them. To correct for this, the Newey and West [26] (NW) expression takes a sandwich form and explicitly accounts for misspecification by summing the covariance structure of the errors, ensuring that the resulting standard errors are asymptotically valid [8, Theorem 14.32].

Given that X_t is stationary and ergodic, so too are the errors from equation (3), so their covariances vanish as the time lag ℓ increases (see Assumptions). Further, because the timescale τ^* is given by the nonlinear function $g(\phi^*)$ (see equation (7)) with derivative $\frac{d}{d\phi}g(\phi^*)$, its standard error can be approximated using the delta method:

$$\text{se}_{\text{NW}}(\phi^*) = \sqrt{q^{-1} \omega q^{-1}}, \quad \text{se}_{\text{NW}}(\tau^*) \approx \text{se}_{\text{NW}}(\phi^*) \cdot \frac{d}{d\phi}g(\phi^*), \quad (18)$$

where

$$q = \mathbb{E}[X_{t-1}^2] \quad \text{and} \quad \omega = \sum_{\ell=-\infty}^{\infty} \mathbb{E}[(X_{t-1} \cdot e_t)(X_{t-1-\ell} \cdot e_{t-\ell})]. \quad (19)$$

The covariance terms in ω capture deviations in the error structure from the standard *iid* case. For the special case of correct specification, when X_t is a true AR1 process, the standard error of the AR1 coefficient ϕ^* reduces to:

$$\text{se}_{\text{Naive}}(\phi^*) = \sqrt{\sigma^2 q^{-1}} \quad (20)$$

where σ^2 is the error variance.

2.4.2 Autocorrelation-Domain Method

To account for model misspecification, although the model defines a parametric regression function with an exponential decay form, it can accommodate deviations from this decay pattern by incorporating the error term (8) covariance structure, allowing for more general stationary and ergodic processes.

Following the description in Hansen [8, Chapter 22.8 and Chapter 23.5], if ϕ^* uniquely minimizes $S(\phi)$, such that $S(\phi) > S(\phi^*)$ for all $\phi \neq \phi^*$, the precision of ϕ^* can be computed using a Newey and West [26] (NW) form that reflects both the curvature of the squared loss function at its minimum and the covariance of the errors. Further, because the timescale τ^* is given by the nonlinear function $g(\phi^*)$ with derivative $\frac{d}{d\phi}g(\phi^*)$, its standard error can be approximated using the delta method:

$$\text{se}_{\text{NW}}(\phi^*) = \sqrt{q^{-1} \omega q^{-1}}, \quad \text{se}_{\text{NW}}(\tau^*) \approx \text{se}_{\text{NW}}(\phi^*) \cdot \frac{d}{d\phi}g(\phi^*). \quad (21)$$

The components q and ω are derived from the regression function $m(k, \phi) = \phi^k$ in (8) and its derivative $m_{\phi,k} = \frac{d}{d\phi}m(k, \phi) = k\phi^{k-1}$, defined as:

$$q = \mathbb{E}[m_{\phi^*,k}^2] = \mathbb{E}[(k\phi^{*k-1})^2] \quad \text{and} \quad \omega = \sum_{\ell=-\infty}^{\infty} \mathbb{E}[(m_{\phi^*,k} \cdot e_k)(m_{\phi^*,k-\ell} \cdot e_{k-\ell})]. \quad (22)$$

The derivative of the regression function evaluated at ϕ^* ($m_{\phi^*,k}$) locally approximates the nonlinear model by a linear one, ensuring that the standard errors are asymptotically valid even with model deviations (see (42)). This is a realistic scenario under the mild conditions of stationarity and ergodicity. In the special case where the errors are *iid*, the standard error of ϕ^* simplifies to the usual formula:

$$\text{se}_{\text{Naive}}(\phi^*) = \sqrt{\sigma^2 q^{-1}} \quad (23)$$

where σ^2 is the error variance.

2.4.3 Autocorrelation/Time-Domain Method

As discussed, ϕ defined in the autocorrelation domain by nonlinear projection (9) captures longer range autocorrelations than when it is defined in the time domain (6). However, it assumes a signal + noise form for the ACF [$\rho_k = \phi^k + e_k$ (8)] which is not realistic for many stochastic processes. For example, a correctly specified AR1 process has an ACF with no additive error ($\rho_k = \phi^k$). For approximating higher order autoregressive processes, the deviations e_k represent misspecification error – the part of the ACF that the approximating model fails to explain. In many cases this error might be systematic and not random, and therefore the definition of standard error from (21) would be incorrect. To address this problem, we propose a hybrid approach where the timescale is defined in the autocorrelation domain by (9) but its standard error is defined in the time domain by (18).

2.5 Standard Error Estimation

2.5.1 Time-Domain Method

The sample standard error estimator takes the form:

$$\widehat{se}_{NW}(\hat{\phi}_{LLS}^*) = \sqrt{\hat{q}^{-1} \hat{\omega} \hat{q}^{-1}}, \quad \widehat{se}_{NW}(\hat{\tau}_{LLS}^*) \approx \widehat{se}_{NW}(\hat{\phi}_{LLS}^*) \cdot \frac{d}{d\phi} g(\hat{\phi}_{LLS}^*) \quad (24)$$

where

$$\hat{q} = \frac{1}{T} \sum_{t=2}^T x_{t-1}^2 \quad \text{and} \quad \hat{\omega} = \sum_{\ell=-M}^M \left(1 - \frac{|\ell|}{M+1}\right) \quad \frac{1}{T} \sum_{1 \leq t-\ell \leq T} (x_{t-1} \cdot \hat{e}_t)(x_{t-1-\ell} \cdot \hat{e}_{t-\ell}). \quad (25)$$

This estimator calculates a weighted sum of the regression scores $x_{t-1} \cdot \hat{e}_t$, where $\hat{e}_t = x_t - \hat{\phi}_{LLS}^* \cdot x_{t-1}$. The true ω is approximated by $\hat{\omega}$ by taking a finite sum of the regression score covariances up to lag M , where M is the lag-truncation (or bandwidth). The weights used in the sum decrease linearly with lag ℓ , following a Bartlett kernel [26]. This kernel not only ensures the standard errors remain non-negative but also regularizes $\hat{\omega}$ to change smoothly with M [8, Chapter 14.35].

For comparison we also include the naive estimator which simplifies under correct specification:

$$\widehat{se}_{Naive}(\hat{\phi}_{LLS}) = \sqrt{\hat{\sigma}^2 \hat{q}^{-1}} \quad (26)$$

$$\widehat{se}_{Naive}(\hat{\tau}_{LLS}) \approx \widehat{se}_{Naive}(\hat{\phi}_{LLS}) \frac{d}{d\phi} g(\hat{\phi}_{LLS}) \quad (27)$$

where $\hat{\sigma}^2 = 1/T \sum_{t=2}^T \hat{e}_t^2$ is an estimate of the error variance.

2.5.2 Autocorrelation-Domain Method

The sample standard error estimator takes the form:

$$\widehat{se}_{NW}(\hat{\phi}_{NLS}^*) = \sqrt{\hat{q}^{-1} \hat{\omega} \hat{q}^{-1}}, \quad \widehat{se}_{NW}(\hat{\tau}_{NLS}^*) \approx \widehat{se}_{NW}(\hat{\phi}_{NLS}^*) \cdot \frac{d}{d\phi} g(\hat{\phi}_{NLS}^*) \quad (28)$$

where

$$\hat{q} = \frac{1}{K} \sum_{k=0}^K (\hat{m}_{\phi,k})^2 = \frac{1}{K} \sum_{k=0}^K (k \hat{\phi}_{NLS}^{*k-1})^2, \quad (29)$$

$$\hat{\omega} = \sum_{\ell=-M}^M \left(1 - \frac{|\ell|}{M+1}\right) \quad \frac{1}{K} \sum_{1 \leq k-\ell \leq K} (\hat{m}_{\phi,k} \cdot \hat{e}_k)(\hat{m}_{\phi,k-\ell} \cdot \hat{e}_{k-\ell}). \quad (30)$$

This estimator calculates a weighted sum of the linearized regression scores $\hat{m}_{\phi,k} \cdot \hat{e}_k$, where $\hat{e}_k = \hat{\rho}_k - (\hat{\phi}_{\text{NLS}}^*)^k$. The estimate of $\hat{\omega}$ takes a finite sum of these scores up to lag M , weighted by a Bartlett kernel, so that $\hat{\omega}$ changes smoothly with M .

In the case of correct specification the equation simplifies to:

$$\widehat{\text{se}}_{\text{Naive}}(\hat{\phi}_{\text{NLS}}^*) = \sqrt{\hat{\sigma}^2 \hat{q}^{-1}} \quad (31)$$

$$\widehat{\text{se}}_{\text{Naive}}(\hat{\tau}_{\text{NLS}}^*) \approx \widehat{\text{se}}_{\text{Naive}}(\hat{\phi}_{\text{NLS}}^*) \frac{d}{d\phi} g(\hat{\phi}_{\text{NLS}}^*) \quad (32)$$

where $\hat{\sigma}^2 = 1/K \sum_{k=0}^K \hat{e}_k^2$ is an estimate of the error variance.

2.5.3 Autocorrelation/Time-Domain Method

The sample standard error for the hybrid method is equivalent to (24), except that the LLS estimator $\hat{\phi}_{\text{LLS}}^*$ is replaced with the NLS estimator $\hat{\phi}_{\text{NLS}}^*$, which redefines the errors to be in the time domain $\hat{e}_t = x_t - \hat{\phi}_{\text{NLS}}^* \cdot x_{t-1}$.

2.6 Estimator Properties

In this section, we describe the large-sample properties of both the Time-Domain Linear Model and Autocorrelation-Domain Nonlinear Model, focusing on the consistency and limiting variance of their respective estimators. Under general conditions — when the linear method is applied to a process that is not AR(1), or the nonlinear method is applied to a decay process that is not exponential — we demonstrate that the asymptotic distribution is Gaussian, with a limiting variance that can be consistently estimated. Consequently, the resulting t-ratios (see equation (54)) are also asymptotically Gaussian. This allows for the construction of hypothesis tests and confidence intervals across timescale maps of the brain.

2.6.1 Time-Domain Method

Following the description in Hansen [8, Theorem 14.29], the ergodic theorem shows that ergodicity is a sufficient condition for *consistent estimation*. Since X_t is stationary and ergodic, so too are $X_t X_{t-1}$ and X_{t-1}^2 , and as $T \rightarrow \infty$:

$$\frac{1}{T} \sum_{t=2}^T x_t x_{t-1} \xrightarrow{p} \mathbb{E}[X_t X_{t-1}] \quad (33)$$

$$\frac{1}{T} \sum_{t=2}^T x_{t-1}^2 \xrightarrow{p} \mathbb{E}[X_{t-1}^2]. \quad (34)$$

Applying the continuous mapping theorem yields:

$$\hat{\phi}_{\text{LLS}}^* = \left(\frac{1}{T} \sum_{t=2}^T x_{t-1}^2 \right)^{-1} \left(\frac{1}{T} \sum_{t=2}^T x_t x_{t-1} \right) \xrightarrow{p} (\mathbb{E}[X_{t-1}^2])^{-1} (\mathbb{E}[X_t X_{t-1}]) = \phi^*. \quad (35)$$

This shows that the coefficients of the Time-Domain Linear Model can be consistently estimated by least squares, for any stationary and ergodic process with parameters defined by projection (equation (6)). Similarly for the regression score estimator in (25):

$$\hat{\omega} \xrightarrow{p} \omega \quad (36)$$

Following Hansen [8, Theorem 14.33], the asymptotic distribution under general dependence states that the *limiting variance* of $\hat{\phi}$ can be approximated using a central limit theorem for correlated observations. As $T \rightarrow \infty$:

$$\frac{\hat{\phi}_{\text{LLS}}^* - \phi^*}{\widehat{\text{se}}_{NW}(\hat{\phi}_{\text{LLS}}^*)} \xrightarrow{d} \mathcal{N}(0, 1). \quad (37)$$

And by the delta method we obtain the limiting variance for the timescale τ :

$$\frac{\hat{\tau}_{\text{LLS}}^* - \tau^*}{\widehat{\text{se}}_{NW}(\hat{\phi}_{\text{LLS}}^*) \cdot \frac{d}{d\phi} g(\hat{\phi}_{\text{LLS}}^*)} \xrightarrow{d} \mathcal{N}(0, 1). \quad (38)$$

2.6.2 Autocorrelation-Domain Method

To show *consistent estimation*, unlike the time-domain method above where we apply the ergodic theorem to the explicit closed-form expression of the estimator, this is not possible for nonlinear estimators because there is no algebraic expression. Instead, nonlinear least squares minimizes the sample objective function $\widehat{S}(\phi)$ from equation (15), which is itself a sample average. By Hansen [8, Theorem 22.1], for any ϕ , the weak law of large numbers shows that:

$$\widehat{S}(\phi) \xrightarrow{p} S(\phi). \quad (39)$$

Further, if the minimizer ϕ^* is unique, $S(\phi) > S(\phi^*)$ for all $\phi \neq \phi^*$, then the sample minimizer from (16) converges in probability to the true minimum as $K \rightarrow \infty$:

$$\hat{\phi}_{\text{NLS}}^* \xrightarrow{p} \phi^* \quad (40)$$

This shows that the parameters of the Autocorrelation-Domain Nonlinear Model can be consistently estimated by least squares. Similarly for the regression score estimator in (30):

$$\hat{\omega} \xrightarrow{p} \omega \quad (41)$$

With the additional assumption that the objective function (9) is Lipschitz-continuous for ϕ near ϕ^* , following Hansen [8, Theorem 23.2], we can approximate the *limiting variance* of ϕ^* and τ^* using a central limit theorem for correlated observations. Under general conditions, the nonlinear least squares estimator has an asymptotic distribution with a similar structure to that of the linear least squares estimator above; it converges to a Gaussian distribution with a sandwich-form variance:

$$\frac{\hat{\phi}_{\text{NLS}}^* - \phi^*}{\widehat{\text{se}}_{NW}(\hat{\phi}_{\text{NLS}}^*)} \xrightarrow{d} \mathcal{N}(0, 1). \quad (42)$$

And by the delta method we obtain the limiting variance for the timescale τ^* :

$$\frac{\hat{\tau}^* - \tau^*}{\widehat{\text{se}}_{NW}(\hat{\phi}_{\text{NLS}}^*) \cdot \frac{d}{d\phi} g(\hat{\phi}_{\text{NLS}}^*)} \xrightarrow{d} \mathcal{N}(0, 1) \quad (43)$$

3 Simulations

3.1 Simulation Settings

Monte Carlo simulations with $N = 10,000$ replications were used to evaluate time- and autocorrelation-domain methods. Time series realizations $x_t = \{x_1, x_2, \dots, x_T\}$ with $T = 4800$ were based on three

data-generating processes, each characterized by a different autocorrelation structure: AR1, AR2, and autocorrelations derived from rfMRI data. All autocorrelation structures shared the same AR1 projection (ϕ_{AR1}) for comparable timescales. That is, there is always a ϕ_{AR1} value that represents the AR1 projection, even if the time series was generated by a more complex process. To define a feasible parameter range for simulation, we referred to the Human Connectome Project dataset. Accordingly, the range of simulated parameters was restricted to positive values. Autocorrelation strength was varied using five ϕ_{AR1} values (0.1 to 0.8) with corresponding timescales τ_{AR1} from equation (7). This design resulted in a total of 15 simulation settings (three data-generating models \times five autocorrelation strengths). For each setting, estimator performance was assessed by relative root mean squared error:

$$\text{rRMSE}(\hat{\tau}) = \frac{\sqrt{\frac{1}{N} \sum_{n=1}^N (\hat{\tau}_n - \tau)^2}}{\tau}, \quad \text{rRMSE}(\widehat{\text{se}}(\hat{\tau})) = \frac{\sqrt{\frac{1}{N} \sum_{n=1}^N (\widehat{\text{se}}(\hat{\tau}_n) - \text{se}(\hat{\tau}))^2}}{\text{se}(\hat{\tau})}. \quad (44)$$

3.1.1 Time-Domain Simulations

In the *AR1 setting*, the data-generating process matches the fitted model, with time series simulated from an AR1 model:

$$x_t = \phi x_{t-1} + e_t, \quad e_t \stackrel{\text{iid}}{\sim} \mathcal{N}(0, 1) \quad (45)$$

The *AR2 setting* introduces a mismatch between AR2 data-generating process and AR1 fitting. While AR2 allows for more complex dynamics (e.g., periodic signals), this study focused on stationary, aperiodic processes typical of rfMRI signals [10]. As a result, the simulations were limited to stationary and aperiodic AR2 processes, with five pairs of AR2 coefficients selected so that the AR1 projections matched the above setting. The following model was used:

$$x_t = \phi_1 x_{t-1} + \phi_2 x_{t-2} + e_t, \quad e_t \stackrel{\text{iid}}{\sim} \mathcal{N}(0, 1) \quad (46)$$

The *HCP setting* did not follow an autoregressive process, using instead empirical ACFs from five brain regions of subject #100610 from the HCP dataset (see [Dataset Description](#)). These regions were selected to match the ϕ_{AR1} projections above. To simulate time series with the same autocorrelation structure as the empirical data, we sampled from a multivariate normal distribution $\mathcal{N}(0, \hat{\Sigma})$, where $\hat{\Sigma} \in \mathbb{R}^{K \times K}$ is the covariance matrix constructed from the sample ACFs. Under stationarity, the matrix $\hat{\Sigma}$ has a Toeplitz structure, meaning its k^{th} off-diagonal elements represent the sample ACF at lag k : $\hat{\Sigma}_{i-k,j-k} = \hat{\Sigma}_{i+k,j+k} = \hat{\rho}_k$. A Cholesky decomposition of the covariance matrix $\hat{\Sigma} = LL^\top$, where L is a lower triangular matrix, was multiplied with Gaussian white noise:

$$x_t = Let_t, \quad e_t \stackrel{\text{iid}}{\sim} \mathcal{N}(0, 1). \quad (47)$$

3.1.2 Autocorrelation-Domain Simulations

Lastly, generating data in the time domain as described above creates an inherent disadvantage for the autocorrelation-domain method when evaluating estimator performance. This method requires a two-step estimation process: (1) computing the sample autocorrelation function (14) and (2) fitting the exponential decay function (16), resulting in cumulative estimation errors. To isolate and evaluate parameter recovery specific to step (2), we directly generated ACFs $\rho_k = \{\rho_0, \rho_1, \dots, \rho_K\}$ with $K = 4800$ using the following signal + noise models:

$$AR1 : \rho_k = \phi^k + e_k \quad (48)$$

$$AR2 : \rho_k = \phi_1 \rho_{k-1} + \phi_2 \rho_{k-2} + e_k \quad (49)$$

$$HCP : \rho_k = \hat{\rho}_k + e_k. \quad (50)$$

Here, $e_k \stackrel{iid}{\sim} \mathcal{N}(0, 1)$ for all settings, and $\hat{\rho}_k$ represents the empirical autocorrelations obtained from subject #100610 from the HCP dataset. While this setting is unrealistic in practice when working with time series data, it enables us to decouple the estimation steps of the autocorrelation-domain method and make fair comparisons with the single-step time-domain method.

3.2 Simulation Results

3.2.1 Results for Autoregressive Simulations

AR1 simulations (Figure 1) are correctly specified because the data-generating process aligns with the fitted models, where each time series is generated from a AR1 process and each ACF by an exponential decay. The figure panels show how accurately the timescale estimators recover the true parameters and their standard errors as the timescale increases. **Panel B** shows larger timescales increase estimate variability, but maintain a relative RMSE below 10% (except that NLS shows an upward bias at small timescales in the ACF + noise setting in **panel B row 3**). **Panels C-D row 1** shows minimal difference in naive versus Newey-West standard errors under correct specification, and both give accurate estimates. **Panels C-D row 2** (dashed lines) illustrate the **Autocorrelation-Domain Method** fit to sample ACFs from time-series data, showing that both naive and Newey-West standard errors are underestimated with high rRMSE. This occurs because the method incorrectly assumes a signal + noise ACF for an AR1 process. While the timescale estimates in **panel B row 2** remain unbiased due to the correct signal specification, the standard errors are biased toward zero because the model introduces variability that is absent in the true process. **Panels C-D row 2** (solid lines) show that this variability is present in the time domain, allowing the **Autocorrelation/Time-Domain Method** to uncover true standard errors, particularly for the Newey-West estimator with less than 20% error. Finally, **panels C-D row 3** show that when the data are generated from AR1 ACFs with added noise, then the **Autocorrelation-Domain Method** accurately estimates standard errors.

AR2 simulation results (Figure 2) explore how timescale estimators perform when the data-generating process is AR2, introducing misspecification since the estimators fit AR1 projections. In this setting, the goal is to assess the impact of specification error on timescale estimation. **Panel A** illustrates the AR2 ACF and its AR1 projection, emphasizing the mismatch between the true and fitted models. **Panel B** shows that the true timescale differs between the two estimators due to their respective definitions. Despite misspecification, both timescale estimators remain mostly unbiased (except that NLS shows an upward bias at small timescales in the ACF + noise setting). **Panel C** illustrates underestimation of standard errors by the naive estimator across all time- and autocorrelation-domain methods, where the downward bias is corrected by the Newey-West standard errors of **Panel D**. As with the AR1 results, **panels C-D row 2** (dashed lines) show that the **Autocorrelation-Domain Method** fit to sample ACFs from time-series data have high rRMSE and the true standard errors are unrecoverable at large timescales even with the Newey-West approach – another example that ACFs of AR processes cannot be represented as signal + noise models. **Panels D row 2** (solid lines) show that the **Autocorrelation/Time-Domain Method** is unbiased, as with **panel D row 3** for data generated from AR2 ACFs with added noise.

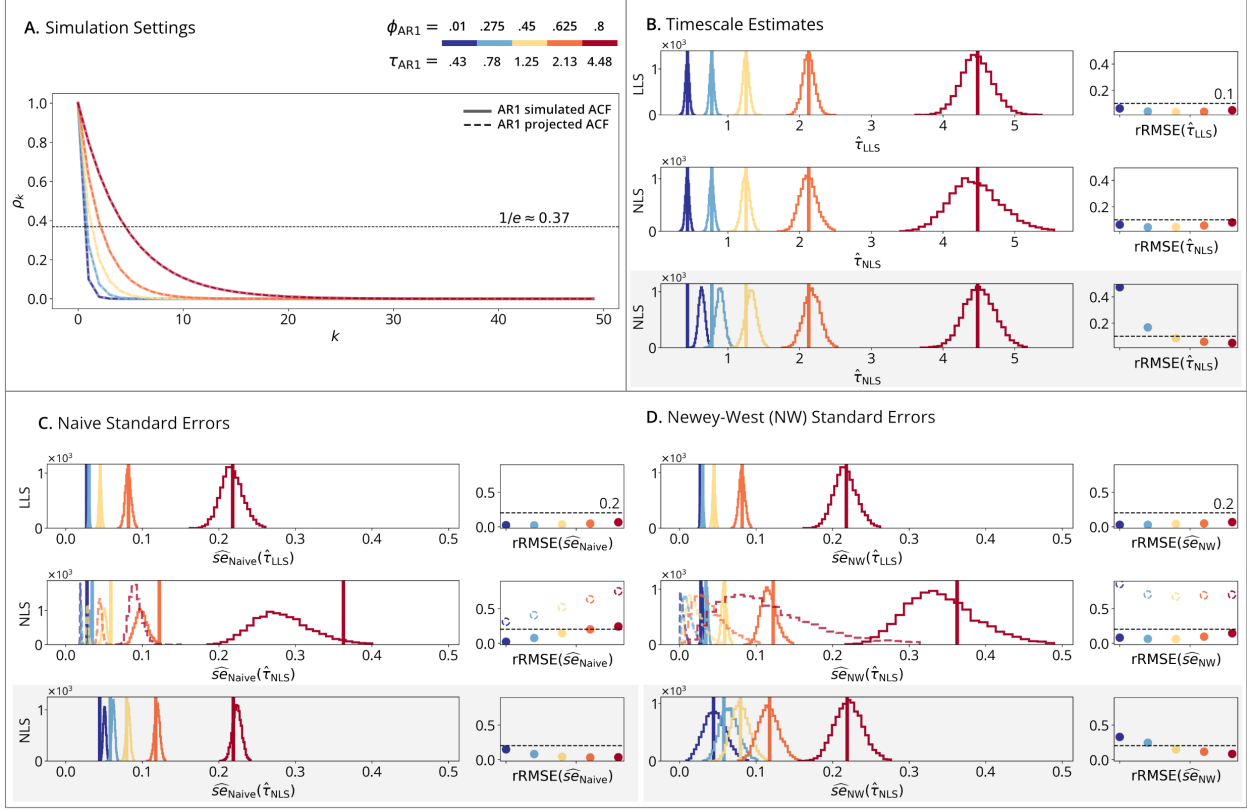


Figure 1: AR1 simulations.

(A) Simulation Setting: solid lines show the simulated ACFs; dashed lines show the AR1-projected ACFs, which are overlapping as both follow AR1. Horizontal line marks the timescale where the AR1-projected ACF reaches $1/e \approx 0.37$. **(B)** Timescale Estimates: vertical lines show true timescales; histograms show estimates across $N = 10,000$ replications; points show rRMSE versus a 10% error line. **(Row 1):** LLS estimator fit to time-series data. **(Row 2):** NLS estimator fit to sample ACFs from time-series data. **(Row 3):** NLS estimator fit to theoretical ACFs with added noise, which is grayed out to indicate it is not a realistic setting. **(C)** Naive and **(D)** Newey-West Standard Errors: vertical lines show standard deviations from panel B; histograms show standard error estimates; points show rRMSE versus a 20% error line. **(Row 1):** time-domain standard errors fit to time-series data. **(Row 2 dashed lines):** autocorrelation-domain standard errors fit to sample ACFs from time-series data. **(Row 2 solid lines):** autocorrelation/time-domain standard errors fit to time-series data. **(Row 3):** autocorrelation-domain standard errors fit to theoretical ACFs with added noise.

3.2.2 Results for Realistic rfMRI Simulations

Realistic rfMRI simulations (Figure 3) explore the performance of timescale estimators when simulating empirical processes derived from five distinct brain regions from a single subject. Like the AR2 results described above, this simulation method is designed to test estimator performance under model misspecification, as the data-generating process reflects realistic brain dynamics, while the fitted models project a simpler AR1 process. **Panel A** shows the mismatch between empirical ACFs and AR1 projections. **Panel B** presents the timescale estimates, where the LLS and NLS estimators yield different timescales due to their respective definitions. **Panels C-D** demonstrate that naive standard errors are underestimated; these errors are largely corrected by applying the Newey-West method. As with the AR simulations, the Autocorrelation-Domain Method approach is effective only when data are generated from ACFs with added noise. Otherwise, standard errors are only accurate when fit in the time

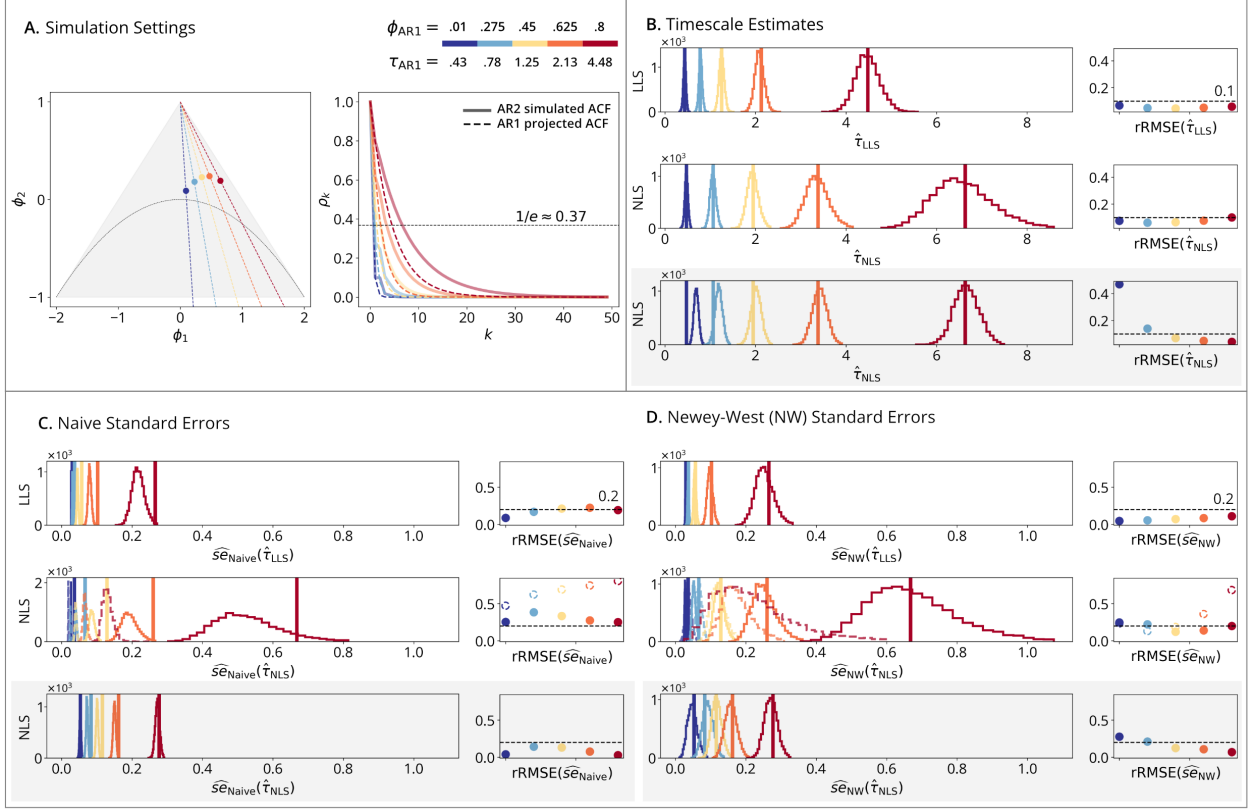


Figure 2: AR2 simulations.

(A) Simulation Setting: triangle shows AR2 stationary region in the (ϕ_1, ϕ_2) plane with a periodic/aperiodic boundary at $\phi_2 = -\phi_1^2/4$. Points show five AR2 (ϕ_1, ϕ_2) pairs with AR1 projections given by the colorbar. Solid lines show the simulated ACFs; dashed lines show the AR1-projected ACFs, which are not overlapping because the simulated AR2 is different from the fitted AR1. (B) Timescale Estimates: vertical lines show true timescales; histograms show estimates across $N = 10,000$ replications; points show rRMSE versus a 10% error line. (Row 1): LLS estimator fit to time-series data. (Row 2): NLS estimator fit to sample ACFs from time-series data. (Row 3): NLS estimator fit to theoretical ACFs with added noise. (C) Naive and (D) Newey-West Standard Errors: vertical lines show standard deviations from panel B; histograms show standard error estimates; points show rRMSE versus a 20% error line. (Row 1): time-domain standard errors fit to time-series data. (Row 2 dashed lines): autocorrelation-domain standard errors fit to sample ACFs from time-series data. (Row 2 solid lines): autocorrelation/time-domain standard errors fit to time-series data. (Row 3): autocorrelation-domain standard errors fit to theoretical ACFs with added noise.

domain, regardless of the domain used for timescale estimation. These results are consistent with the AR2 simulations using more realistic settings.

These figures highlight differences in finite sample bias and variance between the time- and autocorrelation-domain methods. (1) All timescale estimators yield largely unbiased results, while LLS performs as well or better than NLS in terms of rRMSE. (2) Standard error estimates show pronounced differences between naive and Newey-West methods for misspecified settings (AR2, realistic rfMRI). (3) Standard errors cannot be estimated in the autocorrelation domain from sample ACFs of time-series data, and require a hybrid Autocorrelation/Time-Domain Method. (4) Time-domain methods perform as well or better than any of the autocorrelation-domain methods across all simulation results.

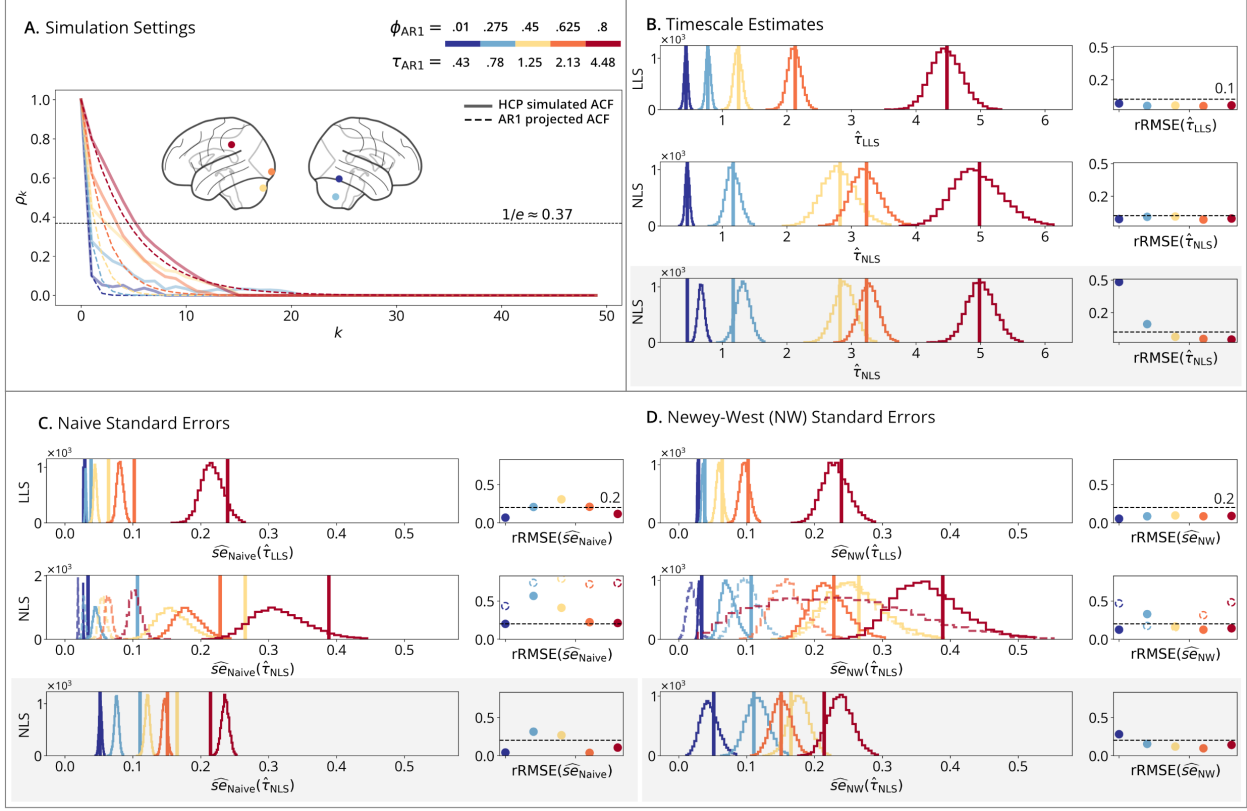


Figure 3: *Realistic rfMRI simulations.*

(A) Simulation Setting: solid lines show simulated ACFs from five brain regions of HCP subject #100610; dashed lines show the AR1-projected ACFs, which do not overlap. (B) Timescale Estimates: vertical lines show true timescales; histograms show estimates across $N = 10,000$ replications; points show rRMSE versus a 10% error line. (Row 1): LLS estimator fit to time-series data. (Row 2): NLS estimator fit to sample ACFs from time-series data. (Row 3): NLS estimator fit to theoretical ACFs with added noise. (C) Naive and (D) Newey-West Standard Errors: vertical lines show standard deviations from panel B; histograms show standard error estimates; points show rRMSE versus a 20% error line. (Row 1): time-domain standard errors fit to time-series data. (Row 2 dashed lines): autocorrelation-domain standard errors fit to sample ACFs from time-series data. (Row 2 solid lines): autocorrelation/time-domain standard errors fit to time-series data. (Row 3): autocorrelation-domain standard errors fit to theoretical ACFs with added noise.

4 Data Analysis

4.1 Dataset Description

Resting fMRI (rfMRI) scans were provided by the Human Connectome Project (HCP), WU-Minn Consortium (led by principal investigators David Van Essen and Kamil Ugurbil; 1U54MH091657) funded by the 16 NIH Institutes and Centers supporting the NIH Blueprint for Neuroscience Research, and by the McDonnell Center for Systems Neuroscience at Washington University [36]. Two subsets of the dataset were used: one for methods development and defining realistic simulation parameters (see Simulations), and the other for estimating high-resolution timescale maps of the cortex.

The *methods development subset* included 10 subjects (#100004 - #101410) scanned with a 3T gradient-

echo EPI sequence (TR=720ms, slice thickness=2mm). Each subject completed four 15-minute runs (4800 timepoints total), preprocessed with standard steps including motion regression and artifact removal (see Glasser et al. [7] for details). The resulting dataset dimensions were {10 subjects, 4800 timepoints, 300 regions}. The *timescale mapping subset* included 180 subjects scanned with a 7T gradient-echo EPI sequence (TR=1000ms, slice thickness=1.6mm) over four 16-minute runs (3600 timepoints total), using the same preprocessing steps. Functional data were analyzed on the cortical surface down-sampled to 2mm spatial resolution, yielding a dataset with the dimensions {180 subjects, 3600 timepoints, 64984 vertices}. The time- and autocorrelation-domain methods were fit to each vertex independently, a mass-univariate analysis approach that resulted in subject-level maps of timescale estimates and their standard errors.

4.1.1 Group-level Analysis

Group-level maps combined individual timescales and standard errors, accounting for within- and between-subject variability. While remaining within the mass-univariate framework, for simplicity, we express the group timescale for the $N = 180$ individual subjects at a single cortical vertex:

$$\hat{\tau}_n \text{ for } n \in \{1, 2, \dots, N\} \quad (51)$$

$$\hat{\tau}_N = \frac{1}{N} \sum_{n=1}^N \hat{\tau}_n. \quad (52)$$

The group-level standard error for the timescale is given by the law of total variance:

$$\widehat{\text{se}}(\hat{\tau}_N) = \sqrt{\frac{1}{N} \sum_{n=1}^N \widehat{\text{se}}(\hat{\tau}_n)^2 + \frac{1}{N} \sum_{n=1}^N (\hat{\tau}_n - \hat{\tau}_N)^2}. \quad (53)$$

Here, the first term under the square root is the within-individual variance and the second term is the between-individual variance.

For visualization, brain-wide t-statistic maps tested whether timescales exceeded 0.5 seconds ($H_0 : \tau \leq 0.5$) by the ratio:

$$t_N = \frac{\hat{\tau}_N - 0.5}{\widehat{\text{se}}(\hat{\tau}_N)}. \quad (54)$$

Additionally, relative standard error (RSE) maps are presented to visualize the spatial precision of timescale estimates by the ratio:

$$\text{rse}(\hat{\tau}_N) = \frac{\widehat{\text{se}}(\hat{\tau}_N)}{\hat{\tau}_N} \quad (55)$$

4.2 Data Analysis Results

4.2.1 Results for rfMRI Timescale Maps

Subject-level maps were generated by mass-univariate fitting of LLS and NLS estimators to cortical surface data. **Panel A** presents the spatial distribution of timescale estimates, which show that NLS estimates tend to yield larger timescales than LLS, consistent with simulation results. **Panel B** shows the corresponding maps of standard errors, which appear align with the timescale maps, consistent with the simulation finding that larger timescales are associated with greater sampling variability. **Panel**

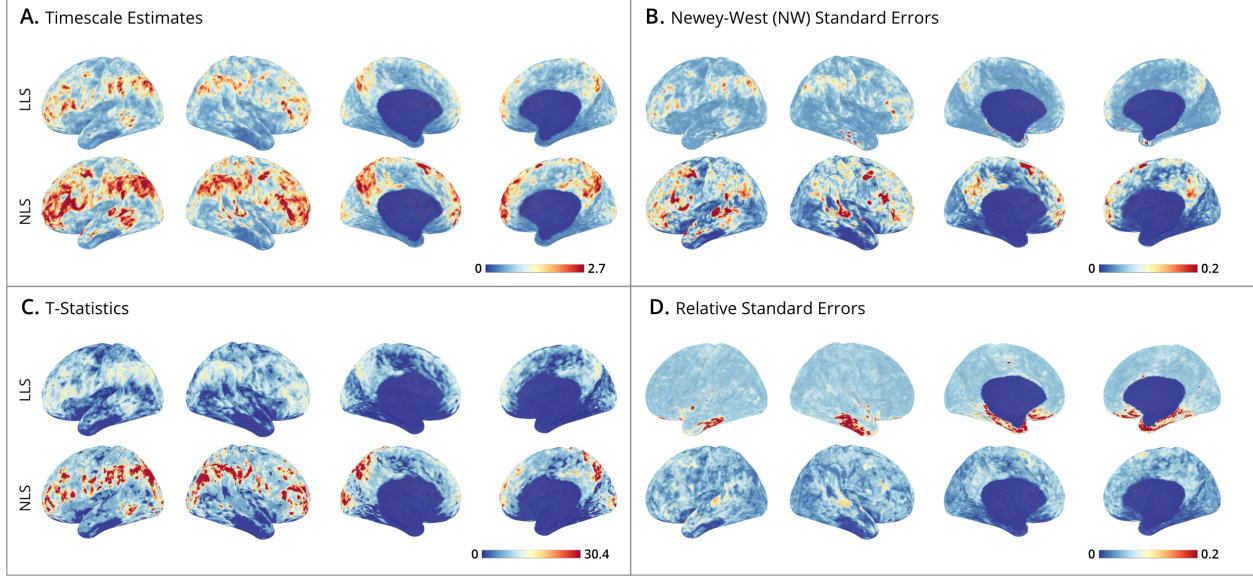


Figure 4: *Human Connectome Project subject-level timescale maps.*

(A-D) Cortical surface maps from HCP subject #100610. Displays show lateral-left, lateral-right, medial-left, and medial-right views (excluding the medial wall). The upper bounds on the colorbars are set for each panel at the 99th percentile of cortical map values. For each panel, the top row shows results of the time-domain method (LLS), and the bottom row the autocorrelation-domain method (NLS). **(A)** Timescale estimates: maps display the timescales (in seconds) estimated at each vertex. **(B)** Newey-West standard errors: shows the spatial distribution of standard errors, where smaller values indicate greater estimation precision. **(B row 2)** shows the hybrid autocorrelation/time method. **(C)** T-statistics: unthresholded and uncorrected t-ratios testing where timescales exceed 0.5 seconds. **(D)** Relative Standard Errors (RSEs): relative reliability of estimates, where low RSE (near zero) indicates high precision with small uncertainty.

C depicts the relative ratio of timescale estimates to their standard errors (i.e., t-statistics), testing whether the timescales significantly exceed a half second ($H_0 : \tau \leq 0.5$). Despite larger standard errors for higher timescales, these regions still exhibit higher t-statistics. **Panel D** shows low RSEs across much of the brain indicating high estimation reliability.

Group-level timescale maps were generated by combining individual estimates to account for both within-individual and between-individual variability, providing an aggregate view of timescale distributions across subjects. **Panel A** shows that the average timescale maps for both LLS and NLS estimators are smoother than the individual maps, displaying a well-organized spatial pattern across the cortex, with NLS estimates generally being larger than LLS. **Panel B** presents the standard error maps, which combine variances from within-subject Newey-West estimates and between-subject timescale estimates. As expected from the simulation results, the standard errors are larger for NLS than for LLS. **Panel C** depicts t-statistics testing whether timescales exceed a half second, showing that both methods yield comparable results. **Panel D** plots relative standard errors (RSEs), illustrating the general trend that regions with larger timescales are easier to estimate, while areas with smaller timescales exhibit greater uncertainty. This is particularly apparent in the limbic network comprising orbital frontal cortex and anterior temporal cortex. **Panel E** highlights the spatial organization of timescales into networks, by mapping the t-statistic at each vertex to one of seven networks from the Thomas Yeo et al. [35] atlas. The ordering {limbic, somatomotor, ventral attention, visual, dorsal attention, default, frontoparietal} aligns with the sensory-to-association axis of brain organization, and is consistent with a large literature on the hierarchical organization of timescales [24, 9, 34, 28, 6, 9]. Sensory networks

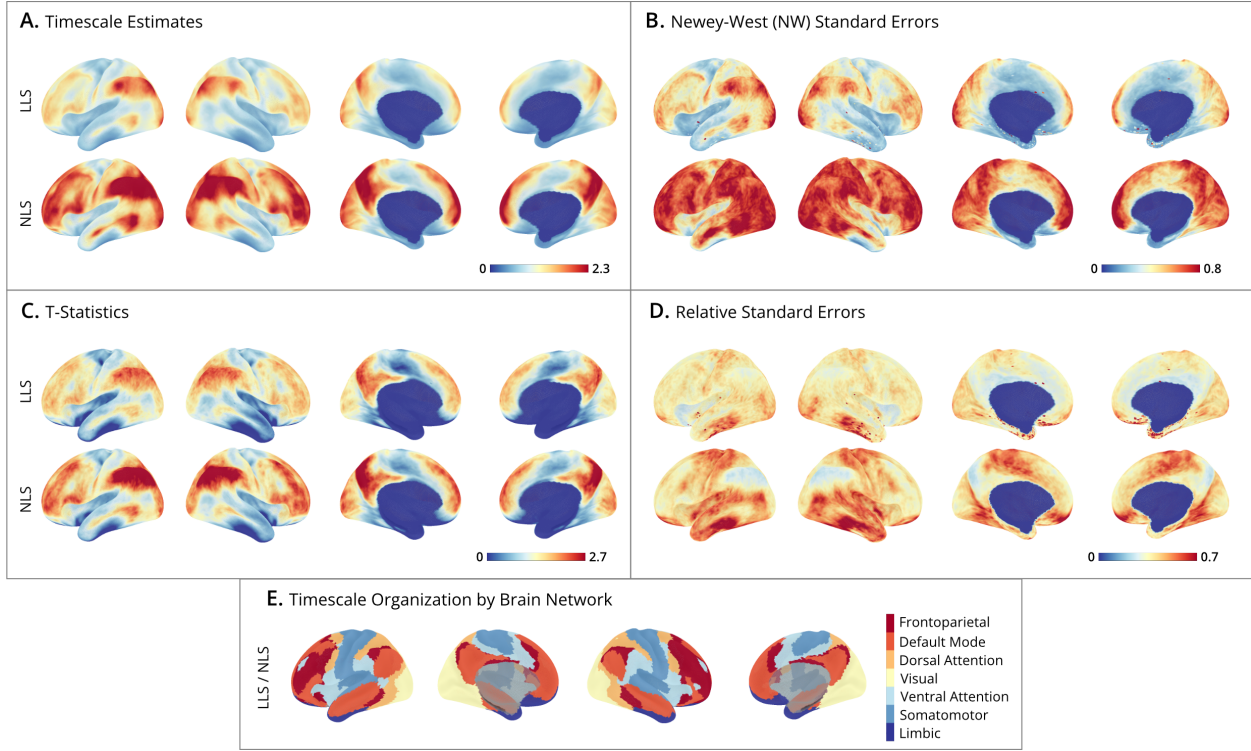


Figure 5: *Human Connectome Project group-level timescale maps.*

(A-D) Cortical surface maps from $N = 180$ HCP subjects. Displays show lateral-left, lateral-right, medial-left, and medial-right views (excluding the medial wall). The upper bounds on the colorbars are set for each panel at the 99th percentile of cortical map values. For each panel, the top row shows results of the time-domain method (LLS), and the bottom row the autocorrelation-domain method (NLS). (A) Timescale estimates: maps display the group-level timescales (in seconds) at each vertex, averaged over subjects. (B) Newey-West standard errors: group-level spatial distribution of estimates, accounting for within- and between-subject variability. Smaller values indicate greater precision. (C) T-statistics: unthresholded and uncorrected t-ratios testing where group-level timescales exceed 0.5 seconds. (D) Relative standard errors (RSEs): relative reliability of estimates, where low RSE (near zero) indicates high precision with small uncertainty across subjects. (E) Timescale organization by brain network: maps display brain networks from the Yeo 7 Network Atlas, ordered by the network-averaged t-statistics (from panel C). This ordering is the same for LLS and NLS methods, and highlights the hierarchical organization of timescales, progressing from sensory networks (e.g., somatomotor and limbic in blues) to association networks (e.g., frontoparietal and default mode in reds).

(limbic and somatomotor) have short timescales, followed by attentional networks (ventral and dorsal attention), and finally higher-order association networks (default mode and frontoparietal) contain the largest timescales.

This empirical analysis highlights methodological considerations for estimating fMRI timescale maps. (1) Both time-domain (LLS) and autocorrelation-domain (NLS) methods produce similar maps, but diverge at extremes – NLS yields larger timescales. The corresponding standard errors for NLS are also larger, so the resulting t-ratio maps remain similar between the two methods, highlighting why point estimates can be misleading without considering standard errors. Likewise, estimates are on average more precise for LLS, consistent with simulation results. (2) In this mass-univariate analysis, the computational cost of LLS is substantially lower than NLS because of its simple analytical solution. (3) The t-statistic maps organized by brain network exhibit a clear hierarchical timescale organization,

reflecting how networks integrate and process information over time. And this pattern is consistent regardless of the estimation method. Taken together, these findings suggest that time-domain method may be preferable for large-scale neuroimaging studies due to its computational efficiency and higher precision, while producing maps consistent with previously reported timescale hierarchies.

5 Conclusions

This study introduces statistical methods for mapping fMRI timescales. We detail the large-sample properties of time- and autocorrelation-domain methods, showing both give estimates that converge, but to different values. This difference arises from the distinct definitions of the timescale parameter inherent to each method. We also demonstrate that both estimators yield consistent standard errors under broad conditions, allowing reliable inference and hypothesis testing. This addresses a major limitation in neural timescale studies that typically report only point estimates without uncertainty measures.

Simulations confirm that both methods provide valid timescale estimates and standard errors, assessing estimation accuracy. Specifically, the time-domain linear method (3) is more computationally efficient and stable for small timescales, while the autocorrelation-domain nonlinear method (8) can capture longer-range autocorrelations at the expense of inaccuracy. Newey-West corrected standard errors reduce bias across methods, enhancing inference reliability [26]. This is evident in AR(2) and empirical rfMRI simulations when autocorrelation decay is not strictly exponential. Applied to HCP rfMRI data, both methods yield timescale and t-statistic maps consistent with known functional hierarchies [36], as other studies similarly show larger timescales in associative versus sensory cortices [28, 31, 17, 23, 14, 39, 33, 19, 13, 25]. More detailed interpretation of the mechanistic interpretation of timescales is reserved for future work.

Although rfMRI provides high spatial resolution compared to EEG, MEG, or ECoG, its sparse temporal sampling can introduce finite sample variability. Asymptotic properties (see Estimator Properties) assume large samples, but in practice, low-frequency sampling of strongly dependent hemodynamic processes can yield too small an effective sample size [1, 14]. Moreover, the mixed metabolic and neuronal origins of the hemodynamic signal complicate mechanistic interpretations [28, 10]. Methodologically, deviations from stationarity and ergodicity can still affect reliability, despite our methods handling common forms of misspecification [8, Chapter 14.7]. Extending model definitions to account for nonstationarity, where autocorrelations are time-dependent, might provide more accurate maps, especially in task-based or dynamic fMRI paradigms [10]. Additionally, adding standard errors to the frequency-domain approach to timescale estimation would allow for the direct modeling of oscillations, which is important when working with electrophysiological recordings of the brain [4, 6].

In conclusion, we introduce robust rfMRI estimators for timescales and standard errors, enabling rigorous statistical comparisons across regions, conditions, and subjects. This advances the accuracy and interpretability of neural timescale maps. Our methods move beyond point estimates by incorporating variability for inference and testing. The work lays the methodological foundation for future research on the role of timescales in brain structure and function.

Code and Data Availability

All simulation results and fMRI timescale maps, inclusive of the code by which they were derived, can be accessed on github.com/griegner/fmri-timescales. The code is under the open source MIT license, allowing access and reuse with attribution. The Human Connectome Project young adult dataset (ages

22-35; 2018 release) used in this study is publicly accessible under a data usage agreement, which describes specific terms for data use and sharing.

Disclosure Statement

The authors declare no conflicts of interest.

References

- [1] S. Afyouni, S. M. Smith, and T. E. Nichols. Effective degrees of freedom of the Pearson's correlation coefficient under autocorrelation. *NeuroImage*, 199:609–625, Oct. 2019. ISSN 10538119. doi: 10.1016/j.neuroimage.2019.05.011. URL <https://linkinghub.elsevier.com/retrieve/pii/S1053811919303945>.
- [2] G. Buzsáki. Large-scale recording of neuronal ensembles. *Nature Neuroscience*, 7(5):446–451, May 2004. ISSN 1097-6256, 1546-1726. doi: 10.1038/nn1233. URL <https://www.nature.com/articles/nn1233>.
- [3] R. Cirillo, V. Fascianelli, L. Ferrucci, and A. Genovesio. Neural Intrinsic Timescales in the Macaque Dorsal Premotor Cortex Predict the Strength of Spatial Response Coding. *iScience*, 10:203–210, Dec. 2018. ISSN 25890042. doi: 10.1016/j.isci.2018.11.033. URL <https://linkinghub.elsevier.com/retrieve/pii/S2589004218302232>.
- [4] T. Donoghue, M. Haller, E. J. Peterson, P. Varma, P. Sebastian, R. Gao, T. Noto, A. H. Lara, J. D. Wallis, R. T. Knight, A. Shestyuk, and B. Voytek. Parameterizing neural power spectra into periodic and aperiodic components. *Nature Neuroscience*, 23(12):1655–1665, Dec. 2020. ISSN 1097-6256, 1546-1726. doi: 10.1038/s41593-020-00744-x. URL <https://www.nature.com/articles/s41593-020-00744-x>.
- [5] J. Fallon, P. G. D. Ward, L. Parkes, S. Oldham, A. Arnatkevičiūtė, A. Fornito, and B. D. Fulcher. Timescales of spontaneous fMRI fluctuations relate to structural connectivity in the brain. *Network Neuroscience*, 4(3):788–806, Jan. 2020. ISSN 2472-1751. doi: 10.1162/netn_a_00151. URL <https://direct.mit.edu/netn/article/4/3/788-806/95846>.
- [6] R. Gao, R. L. van den Brink, T. Pfeffer, and B. Voytek. Neuronal timescales are functionally dynamic and shaped by cortical microarchitecture. *eLife*, 9:e61277, Nov. 2020. ISSN 2050-084X. doi: 10.7554/eLife.61277. URL <https://elifesciences.org/articles/61277>.
- [7] M. F. Glasser, S. N. Sotiropoulos, J. A. Wilson, T. S. Coalson, B. Fischl, J. L. Andersson, J. Xu, S. Jbabdi, M. Webster, J. R. Polimeni, D. C. Van Essen, and M. Jenkinson. The minimal preprocessing pipelines for the Human Connectome Project. *NeuroImage*, 80:105–124, Oct. 2013. ISSN 10538119. doi: 10.1016/j.neuroimage.2013.04.127. URL <https://linkinghub.elsevier.com/retrieve/pii/S1053811913005053>.
- [8] B. E. Hansen. *Econometrics*. Princeton University Press, Princeton, 2022. ISBN 978-0-691-23589-9.
- [9] U. Hasson, E. Yang, I. Vallines, D. J. Heeger, and N. Rubin. A Hierarchy of Temporal Receptive Windows in Human Cortex. *The Journal of Neuroscience*, 28(10):2539–2550, Mar. 2008. ISSN 0270-6474, 1529-2401. doi: 10.1523/JNEUROSCI.5487-07.2008. URL <https://www.jneurosci.org/lookup/doi/10.1523/JNEUROSCI.5487-07.2008>.

- [10] B. J. He. Scale-Free Properties of the Functional Magnetic Resonance Imaging Signal during Rest and Task. *The Journal of Neuroscience*, 31(39):13786–13795, Sept. 2011. ISSN 0270-6474, 1529-2401. doi: 10.1523/JNEUROSCI.2111-11.2011. URL <https://www.jneurosci.org/lookup/doi/10.1523/JNEUROSCI.2111-11.2011>.
- [11] B. J. He, J. M. Zempel, A. Z. Snyder, and M. E. Raichle. The Temporal Structures and Functional Significance of Scale-free Brain Activity. *Neuron*, 66(3):353–369, May 2010. ISSN 08966273. doi: 10.1016/j.neuron.2010.04.020. URL <https://linkinghub.elsevier.com/retrieve/pii/S0896627310002916>.
- [12] Z. Huang, X. Liu, G. A. Mashour, and A. G. Hudetz. Timescales of Intrinsic BOLD Signal Dynamics and Functional Connectivity in Pharmacologic and Neuropathologic States of Unconsciousness. *The Journal of Neuroscience*, 38(9):2304–2317, Feb. 2018. ISSN 0270-6474, 1529-2401. doi: 10.1523/JNEUROSCI.2545-17.2018. URL <https://www.jneurosci.org/lookup/doi/10.1523/JNEUROSCI.2545-17.2018>.
- [13] T. Ito, L. J. Hearne, and M. W. Cole. A cortical hierarchy of localized and distributed processes revealed via dissociation of task activations, connectivity changes, and intrinsic timescales. *NeuroImage*, 221:117141, Nov. 2020. ISSN 10538119. doi: 10.1016/j.neuroimage.2020.117141. URL <https://linkinghub.elsevier.com/retrieve/pii/S1053811920306273>.
- [14] Y. Kaneoke, T. Donishi, J. Iwatani, S. Ukai, K. Shinosaki, and M. Terada. Variance and Autocorrelation of the Spontaneous Slow Brain Activity. *PLoS ONE*, 7(5):e38131, May 2012. ISSN 1932-6203. doi: 10.1371/journal.pone.0038131. URL <https://dx.plos.org/10.1371/journal.pone.0038131>.
- [15] S. Li and X.-J. Wang. Hierarchical timescales in the neocortex: Mathematical mechanism and biological insights. *Proceedings of the National Academy of Sciences*, 119(6):e2110274119, Feb. 2022. ISSN 0027-8424, 1091-6490. doi: 10.1073/pnas.2110274119. URL <https://pnas.org/doi/full/10.1073/pnas.2110274119>.
- [16] N. K. Logothetis. What we can do and what we cannot do with fMRI. *Nature*, 453(7197):869–878, June 2008. ISSN 0028-0836, 1476-4687. doi: 10.1038/nature06976. URL <https://www.nature.com/articles/nature06976>.
- [17] D. J. Lurie, I. Pappas, and M. D’Esposito. Cortical timescales and the modular organization of structural and functional brain networks. *Human Brain Mapping*, 45(2):e26587, Feb. 2024. ISSN 1065-9471, 1097-0193. doi: 10.1002/hbm.26587. URL <https://onlinelibrary.wiley.com/doi/10.1002/hbm.26587>.
- [18] D. J.-N. Maisson, T. V. Cash-Padgett, M. Z. Wang, B. Y. Hayden, S. R. Heilbronner, and J. Zimmermann. Choice-relevant information transformation along a ventrodorsal axis in the medial prefrontal cortex. *Nature Communications*, 12(1):4830, Aug. 2021. ISSN 2041-1723. doi: 10.1038/s41467-021-25219-w. URL <https://www.nature.com/articles/s41467-021-25219-w>.
- [19] A. M. Manea, A. Zilverstand, K. Ugurbil, S. R. Heilbronner, and J. Zimmermann. Intrinsic timescales as an organizational principle of neural processing across the whole rhesus macaque brain. *eLife*, 11:e75540, Mar. 2022. ISSN 2050-084X. doi: 10.7554/eLife.75540. URL <https://elifesciences.org/articles/75540>.
- [20] A. M. G. Manea, D. J.-N. Maisson, B. Voloh, A. Zilverstand, B. Hayden, and J. Zimmermann. Neural timescales reflect behavioral demands in freely moving rhesus macaques. *Nature Communications*, 15(1):2151, Mar. 2024. ISSN 2041-1723. doi: 10.1038/s41467-024-46488-1. URL <https://www.nature.com/articles/s41467-024-46488-1>.

- [21] B. Martin-Burgos, T. S. McPherson, R. Hammonds, R. Gao, A. R. Muotri, and B. Voytek. Development of neuronal timescales in human cortical organoids and rat hippocampus dissociated cultures. *Journal of Neurophysiology*, 132(3):757–764, Sept. 2024. ISSN 0022-3077, 1522-1598. doi: 10.1152/jn.00135.2024. URL <https://journals.physiology.org/doi/10.1152/jn.00135.2024>.
- [22] C. Meisel, K. Bailey, P. Achermann, and D. Plenz. Decline of long-range temporal correlations in the human brain during sustained wakefulness. *Scientific Reports*, 7(1):11825, Sept. 2017. ISSN 2045-2322. doi: 10.1038/s41598-017-12140-w. URL <https://www.nature.com/articles/s41598-017-12140-w>.
- [23] A. Mitra, A. Z. Snyder, C. D. Hacker, and M. E. Raichle. Lag structure in resting-state fMRI. *Journal of Neurophysiology*, 111(11):2374–2391, June 2014. ISSN 0022-3077, 1522-1598. doi: 10.1152/jn.00804.2013. URL <https://www.physiology.org/doi/10.1152/jn.00804.2013>.
- [24] J. D. Murray, A. Bernacchia, D. J. Freedman, R. Romo, J. D. Wallis, X. Cai, C. Padoa-Schioppa, T. Pasternak, H. Seo, D. Lee, and X.-J. Wang. A hierarchy of intrinsic timescales across primate cortex. *Nature Neuroscience*, 17(12):1661–1663, Dec. 2014. ISSN 1097-6256, 1546-1726. doi: 10.1038/nn.3862. URL <http://www.nature.com/articles/nn.3862>.
- [25] E. J. Müller, B. Munn, L. J. Hearne, J. B. Smith, B. Fulcher, A. Arnatkevičiūtė, D. J. Lurie, L. Cocchi, and J. M. Shine. Core and matrix thalamic sub-populations relate to spatio-temporal cortical connectivity gradients. *NeuroImage*, 222:117224, Nov. 2020. ISSN 10538119. doi: 10.1016/j.neuroimage.2020.117224. URL <https://linkinghub.elsevier.com/retrieve/pii/S1053811920307102>.
- [26] W. Newey and K. West. A simple, positive semi-definite, heteroskedasticity and autocorrelation-consistent covariance matrix. *Econometrica*, 55(3):703–08, 1987.
- [27] S. Nougaret, V. Fascianelli, S. Ravel, and A. Genovesio. Intrinsic timescales across the basal ganglia. *Scientific Reports*, 11(1):21395, Nov. 2021. ISSN 2045-2322. doi: 10.1038/s41598-021-00512-2. URL <https://www.nature.com/articles/s41598-021-00512-2>.
- [28] R. V. Raut, A. Z. Snyder, and M. E. Raichle. Hierarchical dynamics as a macroscopic organizing principle of the human brain. *Proceedings of the National Academy of Sciences*, 117(34): 20890–20897, Aug. 2020. ISSN 0027-8424, 1091-6490. doi: 10.1073/pnas.2003383117. URL <https://pnas.org/doi/full/10.1073/pnas.2003383117>.
- [29] R. Rossi-Pool, A. Zainos, M. Alvarez, S. Parra, J. Zizumbo, and R. Romo. Invariant timescale hierarchy across the cortical somatosensory network. *Proceedings of the National Academy of Sciences*, 118(3):e2021843118, Jan. 2021. ISSN 0027-8424, 1091-6490. doi: 10.1073/pnas.2021843118. URL <https://pnas.org/doi/10.1073/pnas.2021843118>.
- [30] C. A. Runyan, E. Piasini, S. Panzeri, and C. D. Harvey. Distinct timescales of population coding across cortex. *Nature*, 548(7665):92–96, Aug. 2017. ISSN 0028-0836, 1476-4687. doi: 10.1038/nature23020. URL <https://www.nature.com/articles/nature23020>.
- [31] G. Shafiei, R. D. Markello, R. Vos De Wael, B. C. Bernhardt, B. D. Fulcher, and B. Misic. Topographic gradients of intrinsic dynamics across neocortex. *eLife*, 9:e62116, Dec. 2020. ISSN 2050-084X. doi: 10.7554/eLife.62116. URL <https://elifesciences.org/articles/62116>.
- [32] G. Shafiei, B. D. Fulcher, B. Voytek, T. D. Satterthwaite, S. Baillet, and B. Misic. Neurophysiological signatures of cortical micro-architecture. *Nature Communications*, 14(1):6000, Sept. 2023. ISSN 2041-1723. doi: 10.1038/s41467-023-41689-6. URL <https://www.nature.com/articles/s41467-023-41689-6>.

- [33] M. Shinn, A. Hu, L. Turner, S. Noble, K. H. Preller, J. L. Ji, F. Moujaes, S. Achard, D. Scheinost, R. T. Constable, J. H. Krystal, F. X. Vollenweider, D. Lee, A. Anticevic, E. T. Bullmore, and J. D. Murray. Functional brain networks reflect spatial and temporal autocorrelation. *Nature Neuroscience*, pages 1–12, Apr. 2023. ISSN 1546-1726. doi: 10.1038/s41593-023-01299-3. URL <https://www.nature.com/articles/s41593-023-01299-3>. Publisher: Nature Publishing Group.
- [34] G. J. Stephens, C. J. Honey, and U. Hasson. A place for time: the spatiotemporal structure of neural dynamics during natural audition. *Journal of Neurophysiology*, 110(9):2019–2026, Nov. 2013. ISSN 0022-3077, 1522-1598. doi: 10.1152/jn.00268.2013. URL <https://www.physiology.org/doi/10.1152/jn.00268.2013>.
- [35] B. T. Thomas Yeo, F. M. Krienen, J. Sepulcre, M. R. Sabuncu, D. Lashkari, M. Hollinshead, J. L. Roffman, J. W. Smoller, L. Zöllei, J. R. Polimeni, B. Fischl, H. Liu, and R. L. Buckner. The organization of the human cerebral cortex estimated by intrinsic functional connectivity. *Journal of Neurophysiology*, 106(3):1125–1165, Sept. 2011. ISSN 0022-3077, 1522-1598. doi: 10.1152/jn.00338.2011. URL <https://www.physiology.org/doi/10.1152/jn.00338.2011>.
- [36] D. C. Van Essen, S. M. Smith, D. M. Barch, T. E. Behrens, E. Yacoub, and K. Ugurbil. The WU-Minn Human Connectome Project: An overview. *NeuroImage*, 80:62–79, Oct. 2013. ISSN 10538119. doi: 10.1016/j.neuroimage.2013.05.041. URL <https://linkinghub.elsevier.com/retrieve/pii/S1053811913005351>.
- [37] D. F. Wasmuht, E. Spaak, T. J. Buschman, E. K. Miller, and M. G. Stokes. Intrinsic neuronal dynamics predict distinct functional roles during working memory. *Nature Communications*, 9(1):3499, Aug. 2018. ISSN 2041-1723. doi: 10.1038/s41467-018-05961-4. URL <https://www.nature.com/articles/s41467-018-05961-4>.
- [38] T. Watanabe, G. Rees, and N. Masuda. Atypical intrinsic neural timescale in autism. *eLife*, 8:e42256, Feb. 2019. ISSN 2050-084X. doi: 10.7554/eLife.42256. URL <https://elifesciences.org/articles/42256>.
- [39] K. Wengler, A. T. Goldberg, G. Chahine, and G. Horga. Distinct hierarchical alterations of intrinsic neural timescales account for different manifestations of psychosis. *eLife*, 9:e56151, Oct. 2020. ISSN 2050-084X. doi: 10.7554/eLife.56151. URL <https://elifesciences.org/articles/56151>.
- [40] R. Zeraati, T. A. Engel, and A. Levina. A flexible Bayesian framework for unbiased estimation of timescales. *Nature Computational Science*, 2(3):193–204, Mar. 2022. ISSN 2662-8457. doi: 10.1038/s43588-022-00214-3. URL <https://www.nature.com/articles/s43588-022-00214-3>.
- [41] F. Zilio, J. Gomez-Pilar, S. Cao, J. Zhang, D. Zang, Z. Qi, J. Tan, T. Hiromi, X. Wu, S. Fogel, Z. Huang, M. R. Hohmann, T. Fomina, M. Synofzik, M. Grosse-Wentrup, A. M. Owen, and G. Northoff. Are intrinsic neural timescales related to sensory processing? Evidence from abnormal behavioral states. *NeuroImage*, 226:117579, Feb. 2021. ISSN 10538119. doi: 10.1016/j.neuroimage.2020.117579. URL <https://linkinghub.elsevier.com/retrieve/pii/S1053811920310648>.