
SpectraLDS: Provable Distillation for Linear Dynamical Systems

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Abstract

We present the first provable method for identifying symmetric linear dynamical systems (LDS) with accuracy guarantees that are independent of the system’s state dimension or effective memory. Our approach builds upon recent work that represents symmetric LDSs as convolutions learnable via fixed spectral transformations. We show how to invert this representation—recovering an LDS model from its spectral transform—yielding an end-to-end convex optimization procedure. This distillation preserves predictive accuracy while enabling constant-time and constant-space inference per token, independent of sequence length. We evaluate our method, SpectraLDS, as a component in sequence prediction architectures and demonstrate that accuracy is preserved while inference efficiency is improved on tasks such as language modeling.

1 Introduction

The emergence of attention-based transformer architectures has revolutionized sequence modeling tasks, particularly in natural language processing [34] and large-scale sequence-to-sequence learning [16][27]. These transformer models rely on the self-attention mechanism, which allows each token in a sequence to attend to every other token, enabling strong contextual understanding. However, this approach suffers from quadratic complexity in sequence length, making it computationally expensive for longer sequences. Recent research has thus explored alternative, more efficient architectures that preserve expressiveness while reducing computational costs for long sequences. Among these are attention-free approaches such as convolution-based or state-space models (SSMs) [19][2][22][26], which can offer sub-quadratic or even near-linear time generation.

The most basic SSM, which is the starting point for all aforementioned models, are linear dynamical systems (LDS), a foundational framework for modeling sequential dependencies in control theory, signal processing, and learning theory [20]:

$$x_t = Ax_{t-1} + Bu_t, \quad \hat{y}_t = Cx_t + Du_t. \quad (1)$$

Here u_t represents the input sequence, x_t encodes the past state information, and \hat{y}_t approximates the target sequence. By maintaining a fixed latent state, LDSs enable efficient inference and allow for efficient memory utilization. However, gradient-based approaches for learning LDSs suffer from exploding or vanishing gradients, particularly when modeling systems with long-term memory or large hidden state spaces [14].

To address these limitations, Agarwal et al. [2] leveraged the spectral filtering method [14] and introduced the Spectral Transform Unit (STU), a convex relaxation that shifts from learning the hidden

transition matrix A directly to a reparameterization of the LDS impulse response that learns how input signals convolve with fixed spectral filters. This approach provably preserves the expressiveness of an LDS while making the training problem far more tractable. In practice, it has proven robust for systems requiring long-range memory and empirical results show that hybrid STU models—with alternating attention and STU layers—can match or even surpass purely attention-based architectures, as well as other popular hybrid state-space models, on long-context tasks such as language modeling and time-series prediction [22]. Furthermore, unlike self-attention, which requires $O(L^2)$ operations per L tokens during training and identically for inference with KV-caching, the STU operates in $O(L \log L)$ operations per L tokens during training and $O(L \log^3 L)$ operations per L tokens during inference through algorithms based on the Fast Fourier Transform [1] [22].

However, although STUs capture LDS-like dynamics using spectral filtering, there has been no straightforward way to convert (or “distill”) a trained STU layer back into explicit LDS form. Moreover, it has remained unclear whether every STU layer can be represented as a LDS with provably small error. Such a distillation from an STU layer to an LDS layer would permit recurrent inference in place of convolution operations, reducing the cost of generating L tokens during inference to $O(L)$, while maintaining the training robustness guarantees of the STU.

Such a distillation is especially desirable in the light of the resurgence of SSMs as scalable alternatives to Transformers in long-context tasks [9] [10] [32]. Thus, our work allows to retain the STU’s performance on long sequences while enabling **constant-time per-token generation**—providing an appealing alternative to both long convolutions and Transformer-based self-attention caching.

1.1 Our Contribution

We present a novel technique for distilling STU filters into LDS form, achieving a substantial reduction in operations—from $O(\log^3 L)$ to $O(\log L)$ per token¹ during generation—while preserving the STU’s expressivity and performance. Moreover, due to the training stability guarantees of the STU architecture, even when learning a marginally stable symmetric LDS or an LDS with high hidden dimension, this distillation procedure provides the **first provable method** to directly learn the parameters of a **symmetric LDS of arbitrarily high effective memory and with bounded noise**. Specifically:

- We show how to convert a learned STU into explicit system matrix parameters whose recurrence can be computed in constant time. This is spelled out in Algorithm 2
- We provide theoretical analysis of such a distillation, in Theorem 1, and empirically demonstrate that the new LDS form incurs negligible degradation in modeling quality while improving autoregressive generation speed.
- We demonstrate in section 6 that by applying SpectraLDS to trained STUs, we maintain the same accuracy and attain constant per-token computational generation cost.
- As a consequence of our theoretical analysis we show in section 5.3 how we can efficiently convert a symmetric LDS of **arbitrary high dimension**, via an intermediate STU learning step, to one of constant dimension and maintain the same accuracy.

2 Related Work

State-space models have long been a cornerstone of control theory and signal processing. The simplest variants, linear dynamical systems, provide a succinct way to capture temporal dependencies via a hidden state propagated in constant time. Classical methods include the Kalman filter [20], which remain widely used due to its robust theoretical properties and computational efficiency.

In recent years, a new wave of SSMs have emerged as efficient alternatives to attention-based methods for long-sequence tasks, promising sub-quadratic or even near-linear complexity without compromising expressive power. Models like S4 [10] and its diagonal variants [11] exploit structured state matrices to learn long-range dependencies, while works such as Hyena, Mega, and Mamba [32] [24] [9] incorporate gating or convolution-based parameterizations to compete against (and

¹We expect the actual reduction to be much larger: from \sqrt{L} , which is a practical method for generation, as opposed to $\log^3 L$, which is a complex algorithm.

sometimes outperform) transformers in language modeling, time-series analysis, and other long-context applications. Their growing popularity and the challenges of training SSMs underscore the need for methods with greater training robustness, efficiency, and performance guarantees. The Spectral Transform Unit (STU) [2] lies squarely in this tradition, offering a powerful convex relaxation for training LDS-like systems that achieves impressive empirical results on long-context tasks. Our work builds directly on this line of research, introducing the first method to distill a learned STU layer into an explicit LDS with provable guarantees, thereby unifying the convex training advantages of spectral filtering with the real-time inference benefits of a recurrent LDS.

Our contributions also align with the long tradition of system identification for LDSs, where the aim is to learn the hidden transition and output matrices (A, B, C, D) from observed sequences. Early influential approaches—such as the Ho-Kalman algorithm [15], Kung’s method [21], and the Eigensystem Realization Algorithm (ERA) [18]—rely on linear-algebraic decompositions (e.g., SVD) of Hankel matrices consistent with observations. Modern variants allow for single-trajectory identification [29], and subsequent refinements like MOESP and N4SID [17] added stochastic noise modeling, while prediction-error and maximum-likelihood methods improved estimation accuracy and statistical efficiency. More recent lines of work incorporate regularization and spectral methods (e.g., stable-spline kernels, sparse identification) to yield more robust or interpretable LDS representations. In the context of distillation, these methods still require matrix decompositions that scale super-linearly in the problem size.

In contrast to many previous approaches, since the STU’s parameterization avoids direct reconstruction of the system matrix A , distillation from a STU layer into an LDS remains agnostic to hidden dimension. Moreover, our method uses a fixed and abstract Hankel matrix—rather than having to construct it anew from observed data—we can perform a significant part of the distillation computation offline.

Finally, we highlight the recent, closely related work of [26], which distills state-space-like recurrences from convolution-based sequence models. Their *Laughing Hyena* method accelerates long convolutional filters by approximating them with a diagonal LDS, thus allowing constant-time generation at inference. While this approach generalizes to any convolution-based model (e.g., Hyena [32]), it does not provide formal guarantees on the distillation quality. In contrast, we focus on the STU’s spectral filters, which have expressive power comparable to a symmetric LDS with real eigenvalues, and present the first theoretical framework to convert the filters into such an LDS with provable bounds on distillation quality. By leveraging the STU’s fixed bank of spectral filters, our method preserves long-sequence expressiveness while achieving a symmetric LDS realization with guaranteed approximation error (see Section 5).

3 Token Generation and Complexity for Language Modeling

In this section, we summarize the autoregressive generation costs for three model classes, Transformers, Convolutional Models, and RNNs, considering a prompt length of T and the generation of K tokens by each model, with L the length of the convolutional filters (i.e., the maximum sequence length). We show the runtimes and memory requirements for each of the listed models in Table 1.

Attention. Processing a prompt requires $O(T^2)$ time. However, token-by-token generation can be accelerated to $O(T + K)$ for each generated token via key-value caching, with a total of $O(T^2 + K(T + K)) = O(T^2 + TK + K^2)$ operations and requiring $O(T + K)$ space [34][26].

Convolutional Model. A Convolutional Model with k convolutional filters will require $O(kN)$ operations to autoregressively generate a new output given N inputs, and thus a naive autoregressive convolutional implementation will require $O(k \cdot K \cdot (T + K))$ operations to generate K tokens. A more refined “Epoched Future Fill” algorithm with prompt prefilling can reduce this to $O(k \cdot (T \log T + K^{3/2} \sqrt{\log K}))$ to generate K tokens. The “Continuous Future Fill” algorithm has theoretical guarantees of $O(k \cdot (T \log T + K \log^2 K))$ operations, although it suffers from numerical instability and has not been implemented or used in practice [1]. For the guarantees of the STU architecture, we require $k = O(\log L)$ and in practice we choose $k = 24$ [22].

RNN (LDS). For an RNN with state dimension h , autoregressive generation requires $O(h)$ operations per token generated and $O(h)$ memory, allowing generation of K tokens with $O(h \cdot (T + K))$

operations. As we will prove, for an LDS with representation capacity comparable to an STU with k filters, we require $h = O(k) = O(\log L)$.

Method	Prefill + Generation Runtime	Cache Size	Runtime with $K, T = O(L)$
Standard Conv	$(TK + T \log T + K^2)k$	$T + K$	$L^2 \log L$
Standard Attn.	$T^2 + TK + K^2$	$T + K$	L^2
EpochedFF	$(T \log T + K^{3/2} \sqrt{\log K})k$	K	$L^{3/2} (\log L)^{3/2}$
ContinuousFF	$(T \log T + K \log^2 K)k$	K	$L \log^3 L$
SpectraLDS (ours)	$(T + K)h$	h	$L \log L$

Table 1: Comparison of architecture runtime and memory requirements for generating K tokens from a length T prompt. $O(\cdot)$ is omitted for brevity. $k, h = O(\log L)$.

4 Problem Background

In this section, we survey the fundamentals relevant to our approach. First, we discuss linear dynamical systems and the inherent challenges of training them directly on tasks requiring long memory. We then present the main theoretical results of spectral filtering and outline how the Spectral Transform Unit (STU) leverages fixed spectral filters to model linear recurrences without explicitly learning the transition matrix A . Finally, we set the stage for our method of distilling STUs back into linear dynamical systems.

4.1 Linear Dynamical Systems

Linear dynamical systems (LDS) have been widely used in control theory to represent time-dependent processes, forming the basis of classical state-space formulations and optimal control methods [20] [23] [3] [19] [6]. Concretely, we consider an input sequence $u_1, u_2, \dots, u_t \in \mathbb{R}^n$, and the corresponding output sequence $y_1, y_2, \dots, y_t \in \mathbb{R}^m$. The hidden state $x_t \in \mathbb{R}^d$ summarizes the system’s memory of past inputs, with the evolution of the system being represented as

$$\begin{aligned} x_t &= A x_{t-1} + B u_t, \\ y_t &= C x_t + D u_t, \end{aligned}$$

where $A \in \mathbb{R}^{d \times d}$, $B \in \mathbb{R}^{d \times n}$, $C \in \mathbb{R}^{m \times d}$, and $D \in \mathbb{R}^{m \times n}$. We omit dynamics and observation noise terms for simplicity with this derivation, although we test our methods on signals with noise.

Expanding the LDS to the Convolutional Form. In a noiseless environment, we can expand the LDS equations as follows:

$$y_t = C x_t + D u_t = C(Ax_{t-1} + Bu_t) + Du_t = \dots = \sum_{i=0}^{t-1} CA^i B u_{t-i} + Du_t.$$

If any eigenvalue $|\lambda_i(A)| > 1$, the system becomes unstable and y_t may tend to infinity in magnitude. Even for $|\lambda_i(A)| < 1$, systems with $\|A\| \approx 1$ are prone to failure due to large A^i powers in backpropagation, as a noisy algorithm may approximate A with spectral radius greater than 1 during training. If $|\lambda_i(A)| < 1 - \delta$ for some spectral gap $\delta > 0$, then

$$y_t = \sum_{i=0}^{\tau} CA^i B u_{t-i} + \varepsilon_k, \quad \|\varepsilon_k\| \leq \varepsilon,$$

where $\tau = O\left(\frac{1}{\delta} \log \frac{1}{\varepsilon}\right)$ and the effective memory is thus on the order $\frac{1}{\delta}$ [2]. As $\delta \rightarrow 0$, learning A directly becomes unstable for large contexts [30] [28], highlighting the need for methods such as spectral filtering. Since the D matrix serves as a skip connection that can be easily computed, we fix it to a 0-matrix and omit its consideration for the remainder of this paper. We will sometimes use the shorthand LDS(C, A, B) to refer to a linear dynamical system with those parameters and $D = 0$.

Additionally, for the remainder of this paper, we restrict our attention to systems where A is a symmetric real matrix. An LDS with symmetric A can be diagonalized, making it equivalent to an LDS with a diagonal A . Without loss of generality, we therefore assume A is diagonal with eigenvalues $\alpha_1, \dots, \alpha_d$.

Spectral Filtering. With initial state $x_0 = \vec{0}$ and defining $\mu(\alpha) = (1, \alpha, \alpha^2, \dots)$, we can extend the convolutional representation:

$$y_t = \sum_{i=0}^{t-1} C A^i B u_{t-i} = \sum_{i=0}^{t-1} C \left(\sum_{\ell=1}^d \alpha_\ell^i (e_\ell \otimes e_\ell) \right) B u_{t-i} = \sum_{\ell=1}^d (c_\ell \otimes b_\ell) \sum_{i=1}^t \mu(\alpha_\ell)(i) \cdot u_{t-i+1}$$

To circumvent the non-convex optimization problem of finding α that best fit an LDS, [14] propose the spectral filtering algorithm, which learns an approximation of $\mu(\alpha)$ in a convex manner. They prove that given eigenvalue-eigenvector pairs $\{\sigma_j, \phi_j\}_j$ of the Hankel matrix Z ,

$$Z := \int_0^1 \mu(\alpha) \mu(\alpha)^\top d\alpha, \quad Z_{i,j} := \frac{2}{(i+j)^3 - (i+j)}$$

any $\mu(\alpha)$ with $0 \leq \alpha \leq 1$ can be approximated by the top k eigenvectors $\{\phi_1, \dots, \phi_k\}$ of Z with an exponentially decreasing error in k . Thus, if y_t is generated by a PSD linear dynamical system, we have the following result:

$$y_t \approx \sum_{\ell=1}^d (c_\ell \otimes b_\ell) \sum_{i=1}^t \tilde{\mu}(\alpha_\ell)(i) \cdot u_{t-i+1} = \sum_{j=1}^k M_j \left(\sum_{i=1}^t \phi_j(i) \cdot u_{t-i+1} \right)$$

where we define $\tilde{\mu}(\alpha) := \sum_{i=1}^k \langle \mu(\alpha), \phi_i \rangle \phi_i$ and learn suitable parameters M_j . Rather than depending on powers of A , learning an LDS with this parameterization remains convex in $\{M_j\}$, since eigenvectors $\{\phi_j\}$ from the matrix Z are computed offline.

The Spectral Transform Unit. To account for learning negative eigenvalues of A , the spectral filtering construction can be adapted by introducing positive and negative sets of feature maps. If $\{\sigma_j, \phi_j\}$ are the eigenvalue-eigenvector pairs of Z , then for each time t and each respective filter ϕ_j , we define the projections of the inputs onto the spectral basis:

$$U_{t,j}^+ = \sum_{i=1}^t u_{t-i+1} \cdot \phi_j(i), \quad U_{t,j}^- = \sum_{i=1}^t u_{t-i+1} \cdot (-1)^{i-1} \cdot \phi_j(i).$$

One then forms the output by learning linear combinations of both $U_{t,k}^+$ and $U_{t,k}^-$ and an optional autoregressive term [12][2]:

$$y_t^{\text{SF}} = \underbrace{\sum_{j=1}^k M_j^{\phi+} U_{t-2,j}^+ + \sum_{j=1}^k M_j^{\phi-} U_{t-2,j}^-}_{\text{Spectral Filtering component}} + \hat{y}_{t-2} + \underbrace{\sum_{i=1}^3 M_i^u u_{t+1-i}}_{\text{AR component}} \quad (2)$$

Without the autoregressive component, we compute $y_t^{\text{SF}} = \sum_{j=1}^k M_j^{\phi+} U_{t,j}^+ + \sum_{j=1}^k M_j^{\phi-} U_{t,j}^-$. The above expression is considered the *Spectral Transform Unit*, where $\{M_j\}$ is the set of parameters to be learned using a differentiable algorithm. Empirical evidence [22] shows that hybrid STU models can compete with or even outperform purely attention-based architectures.

Error Bounds for Spectral Approximation. We repeat the result from [25] and [2] stating that given any LDS parameterized by A, B, C where A is a symmetric matrix with $\|A\| \leq 1$, there exist matrices $M_1^{\phi+}, \dots, M_k^{\phi+}, M_1^{\phi-}, \dots, M_k^{\phi-}$ such that for all L and for all input sequences u_1, \dots, u_L , $\|u_t\| \leq 1$, the following holds for all $t \in [L]$:

$$\|y_t^{\text{LDS}} - y_t^{\text{SF}}\| \sim e^{-\frac{k}{\log L}}.$$

where k is the number of spectral filters, y_t^{LDS} is the sequence generated by the LDS, and y_t^{SF} is the sequence generated by spectral filtering.

Therefore, one can approximate any LDS that meets these specifications up to error ε by selecting $k = O(\log L \log(\frac{1}{\varepsilon}))$ spectral filters. Thus, the STU can capture LDS dynamics with only a logarithmic number of filters k , providing a compact and stable representation even for systems with high effective memory ($\|A\| \approx 1$).

4.2 Distilling STU into an LDS

While the STU avoids direct learning of A , it still implements LDS-like dynamics via spectral filtering. A natural question is whether we can recover explicit parameters $(\tilde{A}, \tilde{B}, \tilde{C})$ from the learned STU, enabling an equivalent recurrence evolving in $O(1)$ time per token, rather than $O(L)$ when convolving spectral filters with input sequences. This can be achieved by approximating the convolution kernel of the STU by the implicit convolution kernel of an LDS. Such a distillation would bridge the gap between the stable convex training of the STU and the fast inference of a recurrent LDS.

5 Algorithm and Main Result

We now present our main theoretical result and accompanying algorithm, which shows how to recover an accurate LDS representation from the learned spectral filters. Concretely, we demonstrate that each STU filter can be approximated by a linear combination of geometrically decaying LDS filters.

5.1 A General Transformation from Spectral Filters to LDS

Our main result is given in Algorithm 2. For the dynamics LDS($1 - \alpha, \alpha, 1$), we denote the impulse response filter by

$$\mu_L(\alpha) = (1 - \alpha)[1 \quad \alpha \quad \alpha^2 \quad \dots \quad \alpha^{L-1}].$$

Note that $\mu_L(\alpha) * u_{1:L} = \sum_{i=1}^L (1 - \alpha)\alpha^{i-1} \cdot u_{L-i+1} = \text{LDS}(1 - \alpha, \alpha, 1)(u_{1:L})$, and thus inference with the LDS is the same as convolution with its impulse filter. Let $\phi_1, \dots, \phi_k \in \mathbb{R}^L$ be the spectral filters of length L . We write the first k filters in matrix form as $\Phi_{1:k} \in \mathbb{R}^{k \times L}$, such that the i -th row is ϕ_i .

A first observation is that we can write any impulse filter approximately in the spectral basis. This is a direct consequence of the spectral filtering methodology for learning an LDS [14].

Algorithm 1 FindSpectralRepresentation

- 1: **Input:** Scalar LDS parameter $\alpha \in \mathbb{R}$, representation size k .
- 2: **Output:** Spectral parameters $m \in \mathbb{R}^k$.
- 3: Construct the impulse response vector $\mu_L(\alpha) \in \mathbb{R}^L$ as $\mu_L(\alpha) = (1 - \alpha)[1, \alpha, \alpha^2, \dots, \alpha^{L-1}]$.
- 4: **return** best spectral fit of the system over random inputs $u \in \mathbb{R}^L$ using gradient descent

$$m = \arg \min_{m \in \mathbb{R}^k} \mathbb{E}_{u \in \mathbb{R}^L} \left[|m^\top \Phi_{1:k} u - \mu_L(\alpha)^\top u|^2 \right].$$

As a consequence of results from spectral filtering for learning LDSs, we show in Appendix A.3 that the procedure `FindSpectralRepresentation` returns a vector m for which, for some $c > 0$,

$$\|m^\top \Phi_{1:k} - \mu_L(\alpha)\| \leq c e^{-\frac{k}{\log L}}.$$

We proceed to use this subroutine to find a distillation to the spectral filters.

Algorithm 2 Spectral Filters to LDS Filters

- 1: **Input:** The first k spectral filters matrix $\Phi_{1:k} \in \mathbb{R}^{k \times L}$; and parameter $h > k$.
- 2: **Output:** The transformation matrix \tilde{M} .
- 3: Sample h randomly chosen independent scalars $\alpha_1, \dots, \alpha_h$ with each $\alpha \in [0, 1]$.
- 4: Construct the matrix $\mu_L(\alpha_{1:h}) \in \mathbb{R}^{h \times L}$ with i -th row $\mu_L(\alpha_i) = (1 - \alpha_i)[1, \alpha_i, \alpha_i^2, \dots, \alpha_i^{L-1}]$.
- 5: For each scalar impulse response, find the spectral representation by

$$m_i = \text{FindSpectralRepresentation}(\alpha_i).$$

- 6: Let M be the $h \times k$ matrix whose rows are m_i .
 - 7: **return** $\tilde{M} := M^{-1} \in \mathbb{R}^{k \times h}$
-

Our main performance guarantee is given in the following theorem.

Theorem 1. As long as $h \geq k$, Algorithm 2 returns w.h.p. a matrix \widetilde{M} such that

$$\left\| \Phi_{1:k} - \widetilde{M} \mu_L(\alpha_1, \dots, \alpha_h) \right\| \leq c \lambda_{\max} h e^{-\frac{k}{\log L}},$$

where λ_{\max} is the largest eigenvalue of the Penrose-Moore pseudo inverse of the matrix M .

The significance of Theorem 1 is that it allows us to translate between the representation of spectral filters and linear dynamical systems.

We note that it is not immediate to upper bound λ_{\max} . Indeed, for $h \sim k$, this can be exponentially large in k , as it corresponds to the condition number of a Vandermonde matrix [4]. However, we note experimentally that as h grows, λ_{\max} quickly becomes smaller. This is an overparametrization effect, which we show experimentally in Appendix A.2. We provide analysis of Theorem 1 in Appendix A.3 and a separate Algorithm 3 with improved practical performance in Appendix A.4.

5.2 Converting \widetilde{M} into an LDS.

As a result of Theorem 1, with $A := \text{Diag}(\alpha_1, \dots, \alpha_h)$, $\Gamma := \text{Diag}(1 - \alpha_1, \dots, 1 - \alpha_h)$ and $\mu := \mu_L(\alpha_1, \dots, \alpha_h)$, we can replace the costly STU convolutions:

$$U_{t,j}^+ = \sum_{i=1}^t u_{t-i+1} \cdot \phi_j(i) \approx \sum_{i=1}^t (\widetilde{M}_j \mu)(i) \cdot u_{t-i+1} = \widetilde{M}_j \sum_{i=0}^{t-1} \Gamma A^i \mathbf{1}_h u_{t-i}^\top = \text{LDS}(\widetilde{M}_j \Gamma, A, \mathbf{1}_h)(u_{1:L}^\top)$$

$$U_{t,j}^- = \sum_{i=1}^t u_{t-i+1} \cdot \phi_j(i) \approx \widetilde{M}_j \sum_{i=0}^{t-1} \Gamma (-A)^i \mathbf{1}_h u_{t-i}^\top = \text{LDS}(\widetilde{M}_j \Gamma, -A, \mathbf{1}_h)(u_{1:L}^\top)$$

This provides the basis of our autoregressive inference advantage, as rather than computing $U_{t,j}^+$ and $U_{t,j}^-$ as pure convolutions, we can maintain the hidden state for $\text{LDS}(\widetilde{M}_j \Gamma, A, \vec{1})$ and $\text{LDS}(\widetilde{M}_j \Gamma, -A, \vec{1})$ for $O(h)$ computation during inference. For practical efficiency, we can compute all $U_{t,1}^+, \dots, U_{t,k}^+, U_{t,1}^-, \dots, U_{t,k}^-$ simultaneously with a single LDS by leveraging the similarities in the state updates (see Appendix A.9).

5.3 LDS to LDS Distillation

A direct consequence of our approach is that we can distill any high-dimensional symmetric LDS into a low-dimensional LDS with bounded error. For an LDS with input dimension d_{in} and output dimension d_{out} , spectral filtering provides an ε -approximation with only $O(d_{in} \cdot d_{out} \cdot \log L \cdot \log(\frac{1}{\varepsilon}))$ parameters regardless of the hidden dimension.

Accepting that $h \lambda_{\max}$ is $O(1)$ for $h \gg k$, as justified in Appendix A.2, we can then convert this spectral representation by application of our distillation procedure into an LDS with state dimension $d_{in} \cdot h$ with $O(d_{in} \cdot d_{out} \cdot \log L \cdot \log(\frac{1}{\varepsilon}))$ learned parameters and $O(d_{in} \cdot d_{out} \cdot \log L \cdot \log(\frac{1}{\varepsilon}) + d_{in} \cdot h)$ total parameters. In other words, the combination of spectral filtering for LDS learning and the following distillation step yields a practical method to reduce state dimension while preserving the system's dynamics within tight error bounds. Empirically, for tasks as difficult as modeling language, we only require $k \leq 24$ filters for strong performance, and state dimension $h \geq 80$ of the distilled LDS suffices to retain performance.

6 Experiments

To demonstrate the effectiveness of our distillation algorithm, we begin by illustrating that a low-dimensional LDS can effectively approximate the spectral filters in Figure 2 and further examine the eigenvalues of the resulting system in Appendix A.9. Building on this, to further quantify the effectiveness of Algorithm 3 in fitting the spectral filters with practically efficient LDSs, we plot the reconstruction error of the spectral filters across different initial subset sizes of LDS impulses

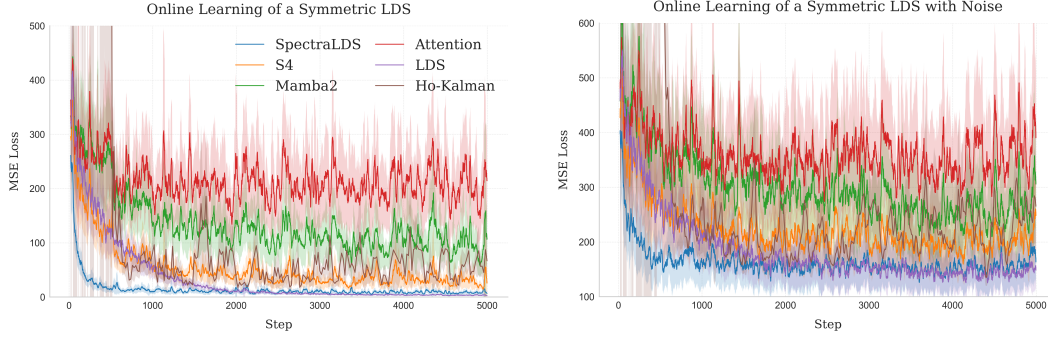


Figure 1: Comparison of SpectraLDS and other methods learning an arbitrary symmetric LDS with and without noise. The shaded region shows the 95% confidence interval over 8 runs. Each model leveraged default configurations except the LDS, which required a lower learning rate to converge. More details are available in Appendix A.7.

in Figure 9 in Appendix A.10. These results show that our algorithm, guided by Theorem 1, not only identifies an initial small state dimension that achieves a low error, but also, by greedily adding 1D-LDS increments, progressively reduces the approximation error to a sufficiently small level without an excessive increase in state dimension.

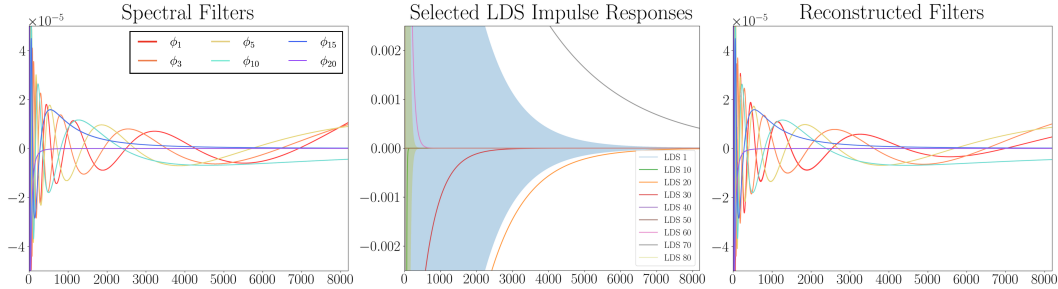


Figure 2: Fit of Spectral Filters by an LDS of state dimension 80 where x-axis represents the time domain. Filters are normalized to be comparable. The blue shading on the middle figure represents a filter quickly alternating (negative eigenvalue). A complete comparison for $k = 24$ without normalization is provided in Appendix A.8.

To validate our method quantitatively, we perform synthetic experiments comparing SpectraLDS against other benchmarks on learning high-memory LDSs (see Appendix A.11). We find that our method significantly outperforms strong baselines in both sample efficiency and reconstruction accuracy, confirming that our approach is well suited for learning systems with long-range dependencies.

Turning to the large-scale evaluation, we distill a 340M-parameter FlashSTU model [22] into an LDS-based architecture and compare their performance across a suite of language benchmarks. From the results in Table 2 we point out that despite the change from convolution-based spectral filters to an explicit LDS representation for the STU layers, the performance remains identical across all tasks. This observation supports our claim that the STU can be closely approximated by a low-dimensional LDS without compromising predictive accuracy. We provide details of the experimental setup and hyperparameters for the models used in Appendix A.14.

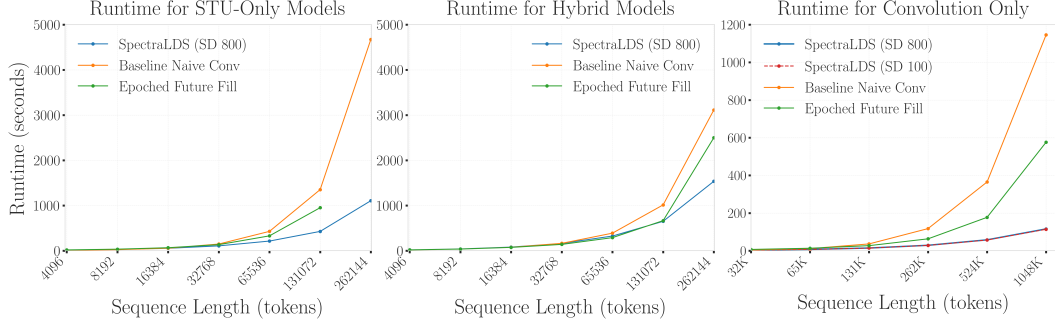


Figure 3: Runtime for generating sequences of increasing length across STU implementations. The naive convolution approach exhibits quadratic growth, the FutureFill variants show logarithmic growth, and the distilled STU-to-LDS layers achieve linear growth. The STU-Only Epoch Future Fill OOMs for the largest sequence length. As shown in the rightmost figure, the SpectralLDS models have nearly identical runtime despite varied state dimension. More results are available in [A.12](#)

Model	MMLU	Hella.	PIQA	BoolQ	Wino.	CSQA	OBQA	ARC-e	ARC-c	Average
Flash STU 340M	26.58	30.46	65.34	60.12	51.85	20.48	20.60	54.08	23.29	39.20
SpectraLDS 340M	26.58	30.45	65.29	60.12	50.99	20.15	20.20	54.17	23.29	39.03
Flash STU Std. Err.	0.37	0.47	1.11	0.86	1.40	1.16	2.06	1.02	1.24	—
SpectraLDS Std. Err.	0.37	0.46	1.11	0.86	1.40	1.15	2.07	1.02	1.24	—
Transformer 340M	26.81	30.41	64.64	61.10	51.62	19.98	18.80	55.47	21.84	38.96

Table 2: Evaluation of a 340M-parameter FlashSTU model and its corresponding distilled LDS representation of state dim. 160 across multiple language benchmarks. Despite converting convolution-based spectral filters into an explicit LDS formulation, performance remains statistically equivalent.

Finally, we present various measures of inference speed to illustrate the constant-time per token generation provided by the distilled STU. In Figure 3 we compare the inference speed of a distilled STU model against a naive convolutional approach and a numerically stable FutureFill variant. In the 12 STU layer model, the naive convolution exhibits quadratic runtime growth with sequence length; the FutureFill variants achieve logarithmic growth; and the distilled STU-to-LDS model demonstrates the best performance with linear growth.

In the hybrid model with 6 attention and 6 STU layers, we find the distilled LDS implementation still provides a significant performance increase. Additionally, we note increases in the LDS state dimension have little impact on the overall runtime, indicating that LDS operations are not a compute bottleneck (see Appendix A.12). For both the hybrid and STU-only models up to 16,384 tokens, the distilled LDS, naive convolution, and Epoch Future Fill all have similar runtimes. All performance benchmarks were conducted on a single H100 GPU, with each generation process evaluated separately to ensure consistent measurements. We provide full experiment details in Appendix A.13.

Looking ahead, further investigation is warranted to better understand how convolution, LDS, and attention layers interact at the hardware level, and to optimize their coordination for even greater speedup. Additionally, further work is required to determine if the LDS layer is stable below float64.

To conclude, as optimized transformer implementations suffer from KV-Cache memory bottleneck rather than compute bottlenecks [9], and the LDS layers have drastically lower memory requirements, we anticipate that, **with appropriate optimization, the inference speed of hybrid attention-STU architectures will be independent of the amount of LDS layers.**

7 Conclusion and Discussion

We have provided the first provable technique for learning the parameters of a symmetric LDS with arbitrarily high effective memory. By leveraging their convex methodology, we show how spectral filters can be distilled into an explicit LDS representation, enabling the construction of a state-space model with constant-time inference and theoretical guarantees on the loss bounds.

References

- [1] Naman Agarwal, Xinyi Chen, Evan Dogariu, Vlad Feinberg, Daniel Suo, Peter Bartlett, and Elad Hazan. Futurefill: Fast generation from convolutional sequence models, 2024.
- [2] Naman Agarwal, Daniel Suo, Xinyi Chen, and Elad Hazan. Spectral state space models, 2024.
- [3] Brian D.O. Anderson and John B. Moore. *Linear Optimal Control*. Prentice-Hall, 1971.
- [4] Bernhard Beckermann. The condition number of real vandermonde, krylov and positive definite hankel matrices. *Numerische Mathematik*, 85(4):553–577, 2000.
- [5] Yoshua Bengio, Patrice Simard, and Paolo Frasconi. Learning long-term dependencies with gradient descent is difficult. *IEEE Transactions on Neural Networks*, 5(2):157–166, 1994.
- [6] Dimitri P Bertsekas. *Dynamic Programming and Optimal Control*, volume 1. Athena Scientific, 2005.
- [7] Tri Dao. FlashAttention-2: Faster attention with better parallelism and work partitioning. In *International Conference on Learning Representations (ICLR)*, 2024.
- [8] Tri Dao, Daniel Y. Fu, Stefano Ermon, Atri Rudra, and Christopher Ré. FlashAttention: Fast and memory-efficient exact attention with IO-awareness. In *Advances in Neural Information Processing Systems (NeurIPS)*, 2022.
- [9] Albert Gu and Tri Dao. Mamba: Linear-time sequence modeling with selective state spaces, 2024.
- [10] Albert Gu, Karan Goel, and Christopher Ré. Efficiently modeling long sequences with structured state spaces, 2022.
- [11] Albert Gu, Ankit Gupta, Karan Goel, and Christopher Ré. On the parameterization and initialization of diagonal state space models, 2022.
- [12] E. Hazan, H. Lee, K. Singh, C. Zhang, and Y. Zhang. Spectral filtering for general linear dynamical systems, 2018.
- [13] Elad Hazan and Karan Singh. Introduction to online nonstochastic control. *arXiv preprint arXiv:2211.09619*, 2022.
- [14] Elad Hazan, Karan Singh, and Cyril Zhang. Learning linear dynamical systems via spectral filtering, 2017.
- [15] B. L. Ho and R. E. Kalman. Effective construction of linear state-variable models from input/output functions. *Automatica*, 3(1):49–59, 1966.
- [16] A. Huang, S. Subramanian, J. Sum, K. Almubarak, S. Biderman, and S. Rush. The annotated transformer, 2022.
- [17] I. W. Jamaludin, N. A. Wahab, N. S. Khalid, S. Sahlan, Z. Ibrahim, and M F. Rahmat. N4sid and moesp subspace identification methods. In *2013 IEEE 9th International Colloquium on Signal Processing and its Applications*, pages 140–145, 2013.
- [18] Jer-Nan Juang and Richard S. Pappa. An eigensystem realization algorithm for modal parameter identification and model reduction. *Journal of Guidance, Control, and Dynamics*, 8(5):620–627, 1985.
- [19] Thomas Kailath. *Linear Systems*. Prentice-Hall, 1980.
- [20] Rudolph Emil Kalman. A new approach to linear filtering and prediction problems. *Journal of Basic Engineering*, 82.1:35–45, 1960.
- [21] S. Y. Kung. A new identification and model reduction algorithm via singular value decompositions. In *Proc. 12th Asilomar Conf. Circuits, Systems and Computers*, pages 705–714, 1978.

- [22] Y. I. Liu, W. Nguyen, Y. Devre, E. Dogariu, A. Majumdar, and E. Hazan. Flash stu: Fast spectral transform units, 2024.
- [23] David Luenberger. An introduction to observers. *IEEE Transactions on Automatic Control*, 16(6):596–602, 1971.
- [24] Xuezhe Ma, Chunting Zhou, Xiang Kong, Junxian He, Liangke Gui, Graham Neubig, Jonathan May, and Luke Zettlemoyer. Mega: Moving average equipped gated attention, 2023.
- [25] Annie Marsden, Evan Dogariu, Naman Agarwal, Xinyi Chen, Daniel Suo, and Elad Hazan. Provable length generalization in sequence prediction via spectral filtering, 2024.
- [26] Stefano Massaroli, Michael Poli, Daniel Y. Fu, Hermann Kumbong, Rom N. Parnichkun, Aman Timalsina, David W. Romero, Quinn McIntyre, Beidi Chen, Atri Rudra, Ce Zhang, Christopher Ré, Stefano Ermon, and Yoshua Bengio. Laughing hyena distillery: Extracting compact recurrences from convolutions, 2023.
- [27] OpenAI. Gpt-4 technical report, 2024.
- [28] A. Orvieto, S. L. Smith, A. Gu, A. Fernando, C. Gulcehre, R. Pascanu, and S. De. Resurrecting recurrent neural networks for long sequences, 2023.
- [29] Samet Oymak and Necmiye Ozay. Non-asymptotic identification of Iti systems from a single trajectory. In *2019 American control conference (ACC)*, pages 5655–5661. IEEE, 2019.
- [30] R. Pascanu, T. Mikolov, and Y. Bengio. On the difficulty of training recurrent neural networks, 2013.
- [31] Adam Paszke, Sam Gross, Soumith Chintala, Gregory Chanan, Edward Yang, Zachary DeVito, Zeming Lin, Alban Desmaison, Luca Antiga, and Adam Lerer. Automatic differentiation in pytorch. 2017.
- [32] Michael Poli, Stefano Massaroli, Eric Nguyen, Daniel Y. Fu, Tri Dao, Stephen Baccus, Yoshua Bengio, Stefano Ermon, and Christopher Ré. Hyena hierarchy: Towards larger convolutional language models, 2023.
- [33] Ofir Press, Noah A. Smith, and Mike Lewis. Train short, test long: Attention with linear biases enables input length extrapolation, 2022.
- [34] Ashish Vaswani, Noam Shazeer, Niki Parmar, Jakob Uszkoreit, Llion Jones, Aidan N. Gomez, Łukasz Kaiser, and Illia Polosukhin. Attention is all you need, 2017.

A Appendix

A.1 Experimental details and notation

All experiments were performed on Nvidia H100-80GB GPUs in PyTorch [31]. All layers except the LDS leverage bfloat16 precision, whereas the LDS layers require float64 precision. All tests, unless otherwise stated, use $k = 24$ spectral filters and replace the k filters with an LDS with state-dimension $h = 80$. To fit the additional 24 negative filters of the negative component (i.e., ϕ_j^- , where $\phi_j^-[i] = (-1)^{i-1}\phi_j[i]$, as is needed to compute $U_{t,j}^-$ purely convolutionally), we expand to an LDS with state-dimension $h = 160$.

For the layer-level inference speed benchmarks, we leverage the STU with tensor-dot approximation (STU-T) [2], the same variant used in the FlashSTU language model [22]. It is worth noting that the STU-to-LDS distillation leads to LDS layers with comparable speed regardless of whether the tensor-dot approximation is employed, and thus this approximation only accelerates the STU layers in benchmarks. A formal definition of this approximation is as follows:

STU Tensor-Dot Approximation: [2] introduced an important optimization, the tensor-dot approximation, wherein each tensor $M \in \mathbb{R}^{d_{\text{in}} \times 2k \times d_{\text{out}}}$, representing a concatenation of the tensors $M_1^{\phi+}, \dots, M_k^{\phi+}, M_1^{\phi-}, \dots, M_k^{\phi-}$, is learned as $M^{(1)} \times M^{(2)}$ for $M^{(1)} \in \mathbb{R}^{d_{\text{in}} \times 2k}$ and $M^{(2)} \in \mathbb{R}^{d_{\text{in}} \times d_{\text{out}}}$. This approximation allows for a reduction in convolutions as, with input x_1, x_2, \dots, x_ℓ , we can compute $y_{\text{SF}}^{\text{SF}} \approx \sum_{i=1}^{\ell} (x_{\ell-i+1}^\top M^{(2)}) \odot M_{\text{filters}}[i]$, where $M_{\text{filters}} = [\phi_1, \dots, \phi_k, \phi_1^-, \dots, \phi_k^-]^\top M^{(1)} \in \mathbb{R}^{L \times d_{\text{out}}}$ and \odot refers to the Schur product (i.e. $(x \odot y)_j = x_j \cdot y_j$). This allows for only d_{out} convolutions with the tensor-dot approximation, as opposed to $k \cdot d_{\text{in}}$ convolutions without it. Although this method can reduce expressivity and does not inherit the same marginal-stability guarantees, it maintains competitive empirical performance while yielding significant improvements in efficiency.

Additionally, we frequently refer to the impulse response of the LDS and STU models. The impulse response of a linear sequence model $f : \mathbb{R}^L \rightarrow \mathbb{R}^1$ is the vector or convolutional kernel $\psi \in \mathbb{R}^L$ that is equivalent to f (i.e. $f(x_{[1, \dots, L]}) = \psi * x_{[1, \dots, L]}$), and thus:

$$\psi[t] = f(\underbrace{[0, \dots, 1, \dots, 0]}_{1 \text{ at position } L-t+1})$$

We only consider the impulse response of $f : \mathbb{R}^L \rightarrow \mathbb{R}^1$, but the impulse response is closely related to the derivative with respect to the inputs and is generalized identically. For an STU model with $d_{\text{in}} = d_{\text{out}} = 1$ and parameters $M_1^{\phi+}, \dots, M_k^{\phi+}, M_1^{\phi-}, \dots, M_k^{\phi-} \in \mathbb{R}$, the impulse response ψ_{SF} is thus described by $\psi_{\text{SF}}[t] = \sum_{j=1}^k M_j^{\phi+} \phi_j[t] + (-1)^{t-1} \sum_{j=1}^k M_j^{\phi-} \phi_j[t]$. Similarly, for an LDS with $d_{\text{in}} = d_{\text{out}} = 1$ and parameters $a, b, c \in \mathbb{R}$, the impulse response ψ_{LDS} is described by $\psi_{\text{LDS}}[t] = ca^{t-1}b$

A.2 Experimental Results on the Condition Number of M , the Spectral Coefficients Matrix

In Figure 4, we present experimental results on the condition number of M as defined in Section 5. Recall that $M \in \mathbb{R}^{h \times k}$ is the spectral coefficients matrix produced by Algorithm 2; its i -th row m_i^* stores the coefficients that express the LDS impulse response $\mu_L(\alpha_i)$ (with geometric decay factor α_i) in the spectral basis $\Phi_{1:k}$. Starting at $h = k$ with $k = 48$, we repeatedly add additional independent vectors α_i and measure the maximum singular value of the constructed M^{-1} . The blue central line shows the mean maximum singular value across 10 experiments for a given value of h , with the shaded region showing the maximum and minimum of the largest singular values across experiments. α were drawn independently from the distribution in Figure 6, which is near uniform but oversamples values in the range $[-1, -0.9] \cup [0.9, 1]$ and greatly oversamples in the range $[-1, -0.99] \cup [0.99, 1]$. To justify our statement in section 5.2 that $\lambda_{\max} \cdot h$ can be considered $O(1)$ in a practical domain, we additionally plot $\lambda_{\max} \cdot h$ under the same experimental setup in Figure 5.

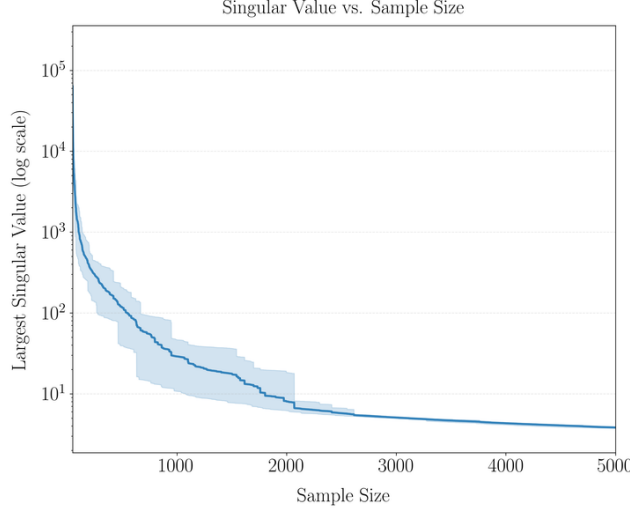


Figure 4: Largest Singular Value as we increase h .

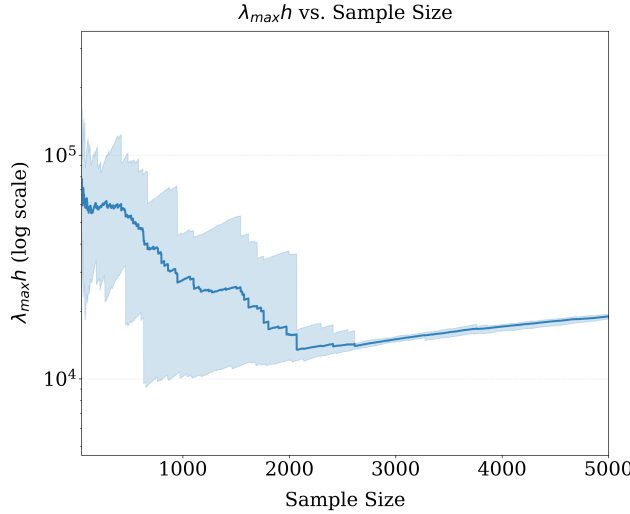


Figure 5: $\lambda_{\max} \cdot h$ as we increase h .

A.3 Analysis of Theorem 1

Proof of Theorem 1 Let

$$\mu_L(\alpha) = (1 - \alpha) [1 \quad \alpha \quad \alpha^2 \quad \dots \quad \alpha^{L-1}] \in \mathbb{R}^L,$$

be the L -impulse response of the one dimensional linear dynamical system with parameters $\alpha, b = 1, c = 1 - \alpha$. Lemma 11.3 from [13] asserts that for any $\alpha \in [0, 1]$, there exist real coefficients $m_1, \dots, m_k \in \mathbb{R}^k$ and constant $c > 0$, such that any sequence of inputs $u_{L:1} \in \mathbb{R}^L$ with $\|u_{L:1}\|_\infty \leq 1$,

$$\left| \sum_{i=1}^k m_i \langle \phi_i, u_{L:1} \rangle - \langle \mu_L(\alpha), u_{L:1} \rangle \right| \leq c e^{-\frac{k}{\log L}}.$$

The RHS term represents the evolution of the LDS, with system matrices b, c that are assumed to be identity, and $\mu_L(\alpha)$ is the system evolution.

As the above is true for all $u_{L:1}$, for each α_j , there exist m_j such that,

$$\forall j \quad , \quad \|m_j^\top \Phi_{1:k} - \mu_L(\alpha_j)\| \leq c e^{-\frac{k}{\log L}}. \quad (3)$$

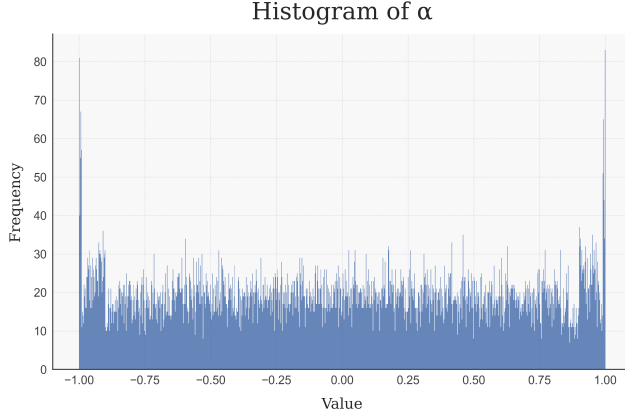


Figure 6: Distribution of α

Moreover, as the optimization problem is convex in m_j and Lipschitz continuous, FindSpectralRepresentation will return such m_j .

Let $M \in \mathbb{R}^{h \times k}$ be the matrix whose rows are $m^j \in \mathbb{R}^k$:

$$M = \begin{pmatrix} -m^1 - \\ -m^2 - \\ \vdots \\ -m^h - \end{pmatrix}.$$

Let $\mathcal{E} = M \Phi_{1:k} - \mu_L(\alpha_1, \dots, \alpha_h)$. By the triangle inequality and (3),

$$\|\mathcal{E}\|_1 \leq \sum_{j=1}^h \|m_j^\top \Phi_{1:k} - \mu_L(\alpha_j)\| \leq c h e^{-\frac{k}{\log L}}. \quad (4)$$

Thus, assuming that M is full rank, multiplying \mathcal{E} by the Penrose-Moore pseudo-inverse of M and using Holder's inequality, we get

$$\|\Phi_{1:k} - M^{-1} \mu_L(\alpha_1, \dots, \alpha_h)\| = \|M^{-1} \mathcal{E}\| \leq \|M^{-1}\|_\star \|\mathcal{E}\|_1 \leq \lambda_{\max} \cdot c h e^{-\frac{k}{\log L}}.$$

It remains to argue that M is full rank. This follows since $\Phi_{1:k}$ is an orthogonal basis, and the matrix $\mu_L(\alpha_{1:h})$ is a Vandermonde matrix. Thus, both matrices are full rank.

□

A.4 The Practical Algorithm

Algorithm 3 LDS Impulse Responses to Matrix Representation

```

1: Input: Learned STU filters  $\Phi_{1:k} \in \mathbb{R}^{k \times L}$ 
2: Output: System matrices  $\{A, B, C\}$  of a discrete-time LDS.
3: Initialize  $\Psi \in \mathbb{R}^{N \times L}$  and  $\Theta \in \mathbb{R}^{N \times k}$  as empty matrices.
4: for  $i = 1, \dots, N$  do
5:   Sample  $(a_i, b_i, c_i)$  with  $|a_i| \leq 1$ 
6:   Train an STU of filter length  $L$  on the 1D-LDS defined by  $(a_i, b_i, c_i)$ 
7:   Let  $\psi_i = (cb, cab, \dots, ca^{L-1}b) \in \mathbb{R}^{1 \times L}$  be the LDS impulse response; update the  $i^{\text{th}}$  row of
 $\Psi$  by  $\Psi[i] \leftarrow \psi_i$ .
8:   Let  $\theta_i \in \mathbb{R}^{1 \times k}$  be the corresponding learned STU weights; update the  $i^{\text{th}}$  row of  $\Theta$  by
 $\Theta[i] \leftarrow \theta_i$ .
9: end for
10: Repeatedly sample a row-subset  $I_{\text{sub}} \subset [1, \dots, N]$  of size  $h_{\text{start}}$  and keep the sample that
minimizes  $\|\Theta_{\text{sub}}^\dagger \Psi_{\text{sub}} - \Phi_{1:k}\|_F^2$ , where  $\Psi_{\text{sub}} := \Psi[I_{\text{sub}}]$  and  $\Theta_{\text{sub}} := \Theta[I_{\text{sub}}]$ .
11: for  $t = h_{\text{start}} + 1, \dots, h$  do
12:    $i^* \leftarrow \operatorname{argmin}_{i \in \{1, \dots, N\} \setminus I_{\text{sub}}} \|\Theta[I_{\text{sub}} \cup \{i\}]^\dagger \Psi[I_{\text{sub}} \cup \{i\}] - \Phi_{1:k}\|_F^2$ 
13:    $I_{\text{sub}} \leftarrow I_{\text{sub}} \cup \{i^*\}$ 
14: end for
15: Initialize  $\widetilde{M} \leftarrow \Theta_{\text{sub}}^\dagger$ 
16: repeat
17:   Update  $\widetilde{M} \leftarrow \widetilde{M} - \eta \nabla_{\widetilde{M}} (\|\Phi_{1:k} - \widetilde{M} \Psi_{\text{sub}}\|_F^2)$ 
18: until convergence
19: Set  $A \leftarrow \operatorname{diag}(\{a_i\}_{i \in I_{\text{sub}}})$ ,  $B \leftarrow (b_i)_{i \in I_{\text{sub}}}$ ,  $C \leftarrow \widetilde{M} \operatorname{diag}(\{c_i\}_{i \in I_{\text{sub}}})$ 
20: return  $\{A, B, C\}$ 

```

A.5 Recovering the Best-Fit LDS for the Spectral Filters

We now elaborate on Algorithm 3 which converts the STU filters into an explicit LDS representation.

1. Generating and Fitting Random LDSs. We first sample many random (a, b, c) system parameters to produce a large collection of one-dimensional LDSs:

$$(1\text{-D-LDS}) \quad x_t = a x_{t-1} + b u_t, \quad y_t = c x_t,$$

where $a \in \mathbb{R}$ satisfies $|a| \leq 1$. In particular, we skew toward generating LDSs with a matrices that have high eigenvalues, and therefore high effective memory, in order to accurately capture the full shape of the spectral filters later in the algorithm (i.e., see Figure 6).

Following the findings of [25], which show that an STU trained on sufficiently long LDS outputs can achieve minimal approximation error, we train an STU to approximate each LDS by feeding in a series of Gaussian inputs u_t into both the LDS and an STU of comparable filter length L . We optimize the STU weights to align its output with that of the LDS, and measure the approximation error using the mean squared error (MSE) between their outputs, discarding any instance whose approximation exceeds a threshold. This procedure corresponds to training the STU to best fit the impulse response for each 1D-LDS due to [14]'s learning guarantees (Appendix A.6). Thus, we yield a dataset of paired (LDS, STU) systems where the STU's learned spectral filters successfully replicate the LDS impulse response.

2. Collecting LDS Impulses and STU Weights. For each of the retained 1D-LDSs, we explicitly record its impulse response, $\psi_{(i)} \in \mathbb{R}^L$, as the vector $\psi_{(i)}[t] = c a^{t-1} b$ for $t = 1, \dots, L$. Stacking these impulse responses row by row yields a matrix

$$\Psi = \begin{pmatrix} -\psi_{(1)}^- \\ -\psi_{(2)}^- \\ \vdots \\ -\psi_{(N)}^- \end{pmatrix} \in \mathbb{R}^{N \times L},$$

where N is the number of LDS examples that we kept.

Meanwhile, we also obtain the STU weights learned to replicate each LDS. In our implementation, each STU layer is parameterized by spectral coefficients $M_1^{\phi+}, \dots, M_k^{\phi+}$ and, if negative filters are used, $M_{-1}^{\phi-}, \dots, M_k^{\phi-}$, which correspond to the positive and negative components of the spectral projection. We can combine these coefficients to form a vector $\theta_i \in \mathbb{R}^k$ (or \mathbb{R}^{2k} if we also incorporate the negative components). Stacking these vectors θ_i row-by-row produces a matrix $\Theta \in \mathbb{R}^{N \times k}$.

At a high level, each row of Ψ is an LDS impulse response, while each row of Θ describes how the STU forms that impulse vector from its spectral basis. Thus, Ψ and Θ capture a variety of 1D-LDSs whose shapes are well-represented by STU filters.

3. Subset Selection and Pseudoinverse Next, we use matrices Ψ and Θ to approximate the target STU filters $\Phi_{1:k}$. We wish to project the STU filters on the basis Ψ of LDS impulses in \mathbb{R}^L .

From the analysis in Section A.3, we look for a matrix \widetilde{M} such that

$$\Phi_{1:k} \approx \widetilde{M} \Psi.$$

We have shown that each row of the LDS impulse vectors Ψ can be approximated by the STU filters using the weights matrix Θ (i.e. $\Theta \Phi_{1:k} \approx \Psi$). Thus, one can invert the filter mapping and form \widetilde{M} :

$$\widetilde{M} = \Theta_{\text{sub}}^\dagger \quad \text{on a suitably chosen subset.}$$

Concretely, our implementation samples fixed-size subsets among the N LDSs and retains the subset I_{sub} that best reconstructs the spectral filters. We then greedily expand I_{sub} to reduce the approximation error of the reconstructed filters $\Phi_{1:k}$. Since each new example adds one dimension (i.e. one more row) to the potential LDS basis, our approach incrementally expands the underlying LDS state dimension only when necessary.

4. Gradient-Based Fine-Tuning. Once we have a suitable subset, we refine the matrix $\widetilde{M} = \Theta_{\text{sub}}^\dagger$ by gradient descent:

$$\widetilde{M} = \operatorname{argmin}_{\widetilde{M}} \|\Phi_{1:k} - \widetilde{M} \Psi_{\text{sub}}\|_F^2.$$

Since Ψ_{sub} and $\Phi_{1:k}$ are both fixed, this is a simple least-squares objective. Empirically, we often observe an approximately 1.4x decrease in the MSE for the fits of the spectral filters once this stage completes.

5. Combining Into a Single LDS. Finally, we merge the chosen 1D-LDSs into a single higher-dimensional discrete-time system. In particular, suppose our final subset Ψ_{sub} includes h different 1D systems, with parameters $\{(a_i, b_i, c_i)\}_{i=1}^h$. The gradient refined matrix \widetilde{M} helps determine how to combine these 1D systems to produce the best approximation of the spectral filters. We then form a system:

$$\mathbf{A} = \operatorname{diag}(a_1, \dots, a_h) \in \mathbb{R}^{h \times h}, \quad \mathbf{B} = \begin{pmatrix} b_1 \\ \vdots \\ b_h \end{pmatrix} \in \mathbb{R}^{h \times 1}, \quad \mathbf{C} = \widetilde{M} \operatorname{diag}(c_1, \dots, c_h) \in \mathbb{R}^{k \times h}$$

where each (a_i, b_i, c_i) corresponds to one of the original 1D LDS components. The net dimension (i.e., the number of included components.) of the hidden state is h .

Thus, for any vector $v \in \mathbb{R}^{1 \times k}$, $v \cdot \Phi_{1:k} \approx v \cdot \langle CB, CAB, CA^2B, \dots \rangle$, and the coefficients learned on the spectral filters can be directly ported to the LDS. In this manner, the STU's spectral filters are translated into a classical LDS. Empirically, $h \approx 160$ has high accuracy in our experiments. Once we have parameters $\{\mathbf{A}, \mathbf{B}, \mathbf{C}\}$ in standard form, we can generate new sequences in $\mathcal{O}(h)$ time per token, exactly like a conventional recurrent model, while achieving the same performance as the STU's filters.

For a practical performance boost, although we fit each 1D-LDS with an STU with positive and negative filters, we then fit I_{sub} and \widetilde{M} by only considering the reconstruction error on the positive

filters and return an LDS that solely reconstructs the positive filters. Then replacing A with $-A$ leads to a new LDS that fits the negative spectral filters, and we can thus fit both positive and negative filters with state dimension $2h$. This approach typically leads to better reconstruction accuracy.

A.6 Why Matching Outputs Corresponds to Matching Impulse Responses

We start by recalling that Theorem 2.1 of [2] ensures that the outputs of an LDS and the spectral filtering (SF) approximation are close for any bounded input sequence $\{u_t\}$ with $\|u_t\| \leq a$ when A is a PSD matrix with $\|A\| \leq 1$. In particular, if we set $\{u_t\}$ to the sequence of all ones (so $a = 1$), the theorem guarantees that for every time t ,

$$\|y_t^{\text{LDS}} - y_t^{\text{SF}}\|^2 \leq \varepsilon,$$

for some ε on the order of $c \|B\|_{\text{col}} \|C\|_{\text{col}} L^3 e^{-\left(\frac{\pi^2}{4}\right) \cdot \frac{k}{\log(L)}}$.

Let $\psi_{\text{LDS}}[i]$ denote the i -th impulse response of the LDS. Likewise, let $\psi_{\text{SF}}[i]$ be the corresponding impulse response of the spectral filtering approximation. We observe that if the input sequence is all ones up to time $r \geq t$, then the resulting output at time t is precisely the sum of the first t impulse responses:

$$y_t^{\text{LDS}} = \sum_{i=1}^t \psi_{\text{LDS}}[i], \quad \text{and} \quad y_t^{\text{SF}} = \sum_{i=1}^t \psi_{\text{SF}}[i].$$

Hence, the condition $\|y_t^{\text{LDS}} - y_t^{\text{SF}}\|^2 \leq \varepsilon$ for all t can be rewritten as

$$\left\| \sum_{i=1}^t \psi_{\text{LDS}}[i] - \sum_{i=1}^t \psi_{\text{SF}}[i] \right\|^2 \leq \varepsilon \quad \text{for each } t.$$

To see that each individual impulse $\psi_{\text{LDS}}[t]$ is close to $\psi_{\text{SF}}[t]$, note that

$$\psi_{\text{LDS}}[t] - \psi_{\text{SF}}[t] = \left(\sum_{i=1}^t \psi_{\text{LDS}}[i] - \sum_{i=1}^t \psi_{\text{SF}}[i] \right) - \left(\sum_{i=1}^{t-1} \psi_{\text{LDS}}[i] - \sum_{i=1}^{t-1} \psi_{\text{SF}}[i] \right).$$

As both $\|\sum_{i=1}^t \psi_{\text{LDS}}[i] - \sum_{i=1}^t \psi_{\text{SF}}[i]\|^2 \leq \varepsilon$ and $\|\sum_{i=1}^{t-1} \psi_{\text{LDS}}[i] - \sum_{i=1}^{t-1} \psi_{\text{SF}}[i]\|^2 \leq \varepsilon$, then

$$\|\psi_{\text{LDS}}[t] - \psi_{\text{SF}}[t]\|^2 \leq \left\| \sum_{i=1}^t \psi_{\text{LDS}}[i] - \sum_{i=1}^t \psi_{\text{SF}}[i] \right\|^2 + \left\| \sum_{i=1}^{t-1} \psi_{\text{LDS}}[i] - \sum_{i=1}^{t-1} \psi_{\text{SF}}[i] \right\|^2 \leq 2\varepsilon.$$

Therefore, matching the outputs of the LDS and the STU within small error forces each impulse response of the LDS to be close to the corresponding impulse response of the SF approximation, which generalizes to any input sequence bounded as above. We thus conclude that matching outputs implies matching impulse responses.

A.7 Learning a Symmetric LDS with and without noise

Figure 1 compares SpectraLDS to other system-identification methods on the task of learning an arbitrary symmetric LDS with and without noise. The LDS signal has hidden dimension 256, input and output dimension 5, and maximum eigenvalue magnitude 0.99 with Gaussian initialization. Each step provides a Gaussian input sequence of length 100 with variance $1/d_{in}$ and the final output. For the learning with noise experiment, Gaussian noise with 0.5 variance was added to the hidden state at each step, and Gaussian noise with 5 variance was added to the output. Each architecture was trained with the default PyTorch RMSProp optimizer configuration, except for the LDS, which required a lower learning rate to converge. The Ho-Kalman matrices are recomputed every 20 steps for computational ease, while the other methods were updated every step, and the Ho-Kalman parameters are the maximum for an input of length 100 (that is $T_{\text{horizon}} = 99$, $T1 = 49$, $T2 = 49$, $\text{state_dim_est} = 48$). The shaded region in the figure shows the 95% confidence interval over 8 runs.

A.8 Fit of the Spectral Filters with a Linear Dynamical System (State Dimension 80)

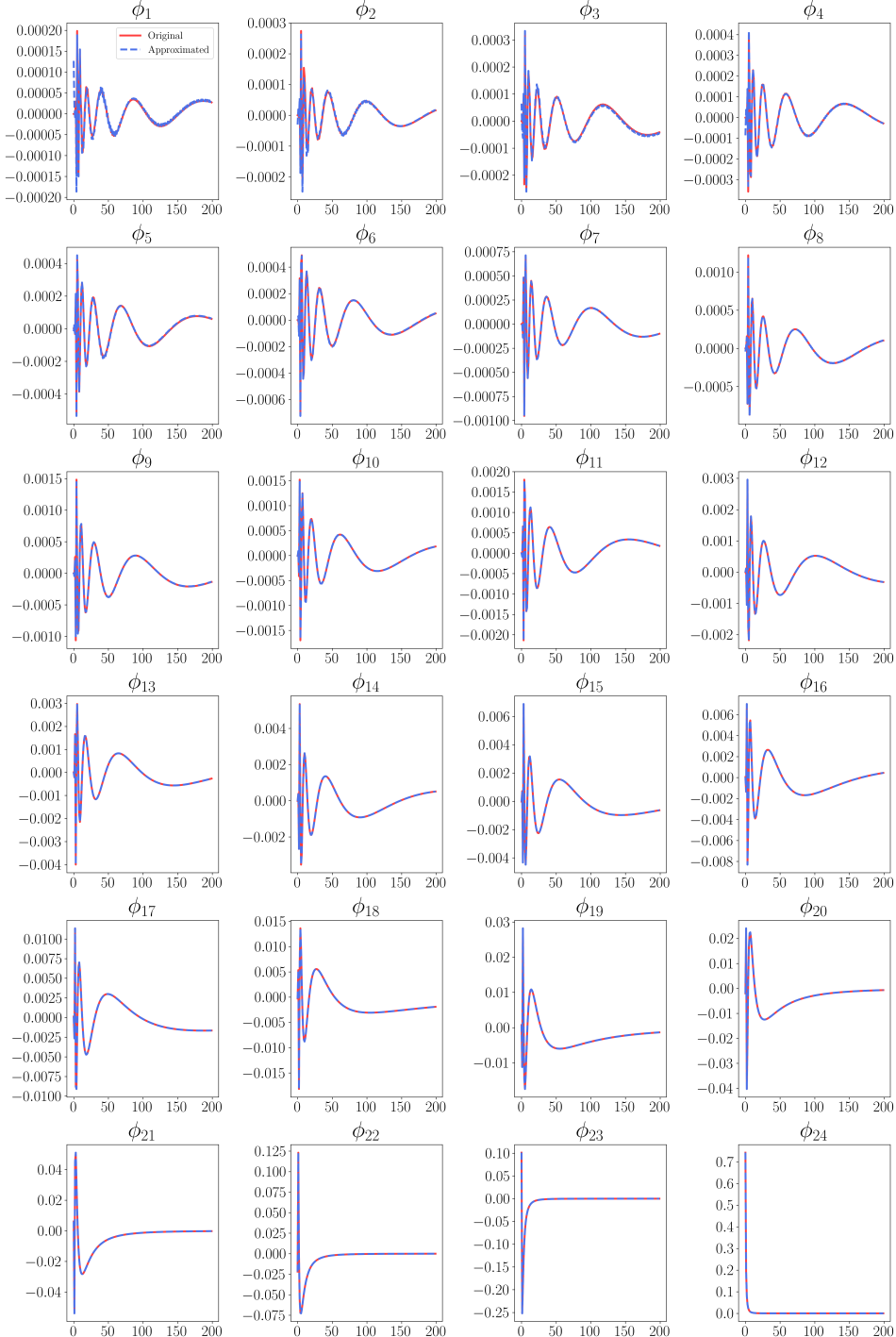


Figure 7: Visualization of the positive spectral filters and their approximation using an LDS with state dimension 80, as obtained via Algorithm 3. The figure illustrates that even with a low-dimensional state, the LDS can accurately fit the spectral filters, confirming the efficacy of our distillation process.

A.9 State Dynamics of the Linear Dynamical System

We train a linear dynamical system with state dimension 80 to fit the Spectral Filters using Algorithm 3 and examine the eigenvalues of the resulting system in Figure 8. The reconstruction error of the spectral filters for the trained LDS is 1.23×10^{-12} . We transform the eigenvalues to visualize the full range of the values as they approach 1 and differentiate the negative and positive eigenvalues as blue and red (e.g. 0.97 and -0.97 would both map to ≈ 3.5 , although the first would join a red column and the second blue).

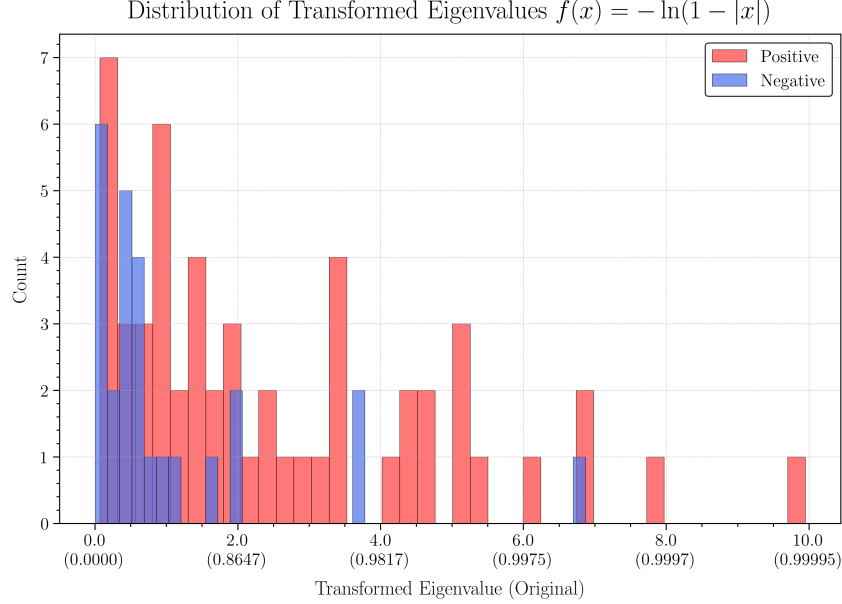


Figure 8: Distributions of Eigenvalues for LDS trained to predict first 24 Spectral Filters.

For practical efficiency, we can compute all $U_{t,1}^+, \dots, U_{t,k}^+, U_{t,1}^- \dots U_{t,k}^-$ simultaneously with a single LDS parameterized by $\left\{ \begin{bmatrix} \widetilde{M} \Gamma & \mathbf{0} \\ \mathbf{0} & \widetilde{M} \Gamma \end{bmatrix}, \begin{bmatrix} A & \mathbf{0} \\ \mathbf{0} & -A \end{bmatrix}, \mathbf{1}_{2h} \right\}$. This LDS has a state dimension of $2hm$, where each of the m input dimensions have a hidden state updated by $\begin{bmatrix} A & \mathbf{0} \\ \mathbf{0} & -A \end{bmatrix}$. The hidden state dimension differs from $2h$ as we treat m as a batch dimension, resulting in $Bu_t \in \mathbb{R}^{2h \times m}$ rather than \mathbb{R}^{2h} . This mimics an LDS acting independently across each of the m dimensions, equivalent to treating m as a batch-axis, and this is the analog to how each of the k spectral filters convolve along each input dimension independently. The above Figure 8 showcases the eigenvalues of matrix A , whereas for practical deployment, the state matrix is $\begin{bmatrix} A & \mathbf{0} \\ \mathbf{0} & -A \end{bmatrix}$.

A.10 Reconstruction Error of the Spectral Filters by Initial LDS State Dimension

We demonstrate the efficacy of Algorithm 3 in fitting the spectral filters with a low initial LDS state dimension. We generate 61784 1D-LDSs, with A 's eigenvalue distribution similar to Figure 6. For each LDS, we then fit it with an STU model with $k = 24$ filters and without an autoregressive term. This provides a large set of LDS impulse and STU impulse pairs, which allows us to relate the spectral filters to the LDS impulses. As described in Algorithm 3, we choose an initial subset of these LDSs, perform the matrix inversion procedure described in Algorithm 2 and measure the error. We greedily add additional LDS impulses, which corresponds to increasing h , to minimize the loss until we reach a desired state dimension. In Figure 9, we ablate the starting subset size for this procedure and measure the loss. Before running this procedure, we filter out any LDS-STU pairs where the STU did not learn the LDS to within a certain threshold. This figure does not include the final gradient descent step used to improve the fit, which typically leads to 30% to 50% improvements. Figure 7 and Figure

[\[8\]](#) leverage an LDS with state dimension 80 and error $1.23 \cdot 10^{-12}$. In this procedure we only aim to fit the k positive spectral filters. To fit both the k positive spectral filters and k negative spectral filters, we can achieve the same error with double the state dimension, as described in Appendix [A.9](#)

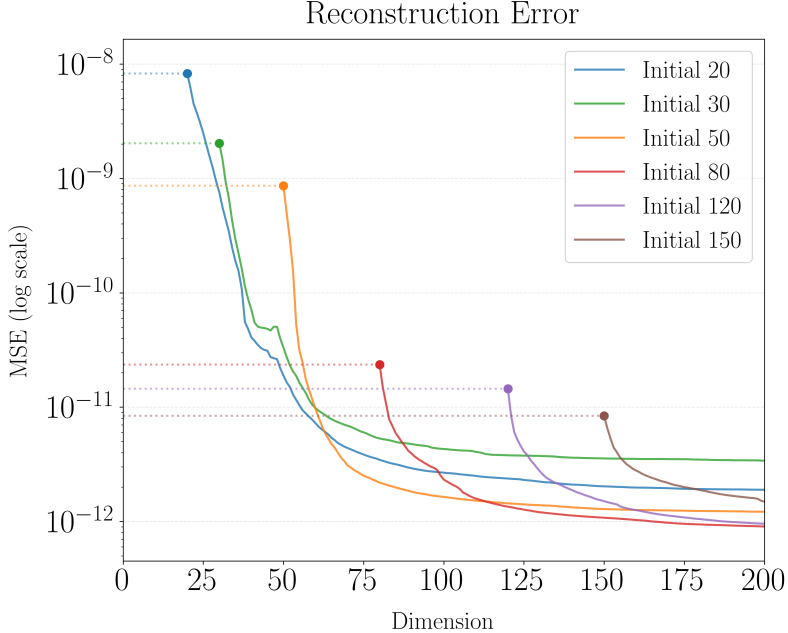


Figure 9: Reconstruction error of spectral filters as a function of LDS state dimension. Each curve starts from a different initial subset size of LDS impulses (chosen by best sample of that size out of 10^6 trials) and is then expanded greedily in single-dimensional increments until reaching a final state dimension of 200.

A.11 Experiments on Synthetic Tasks

We evaluate SpectraLDS against a strong baseline on a synthetic task, reporting results in Table [3](#) and Figure [10](#). A symmetric linear dynamical system with input dimension d_{in} and state dimension d_h is initialized by drawing all entries from a standard normal distribution, after which the update matrix A is rescaled so its largest eigenvalue has magnitude $1 - \delta$. We then distill a SpectraLDS model from an STU model trained with AdaGrad (learning rate 1.0) for 2000 steps; each step minimizes the mean-squared-error (MSE) between the STU output and the ground-truth LDS output over batches of 32 sequences of length seq_len . For the baseline, we fit a randomly initialized symmetric LDS with AdaGrad (learning rate 0.0001) under the same loss. To keep training times practical, most baseline runs use a shorter sequence length, and several runs were stopped early due to computational limits, marked with an asterisk. **When the state dimension satisfies $d_h \geq 1000$, baseline training with a learning rate of 0.01 becomes unstable and often diverges, whereas the STU remains stable even with a learning rate of 1.0.**

Performance Results											
Type	Len.	Delta	d_in	d_h	Avg MSE	MSE Std	Step 10 Loss	Step 100 Loss	Runs	Time (s)	Time Std (s)
LDS GD	8192	1×10^{-2}	10	100	2.15×10^{-2}	1.32×10^{-3}	3.49×10^{-1}	1.43×10^{-1}	4*	2153.12	14.16
LDS GD	8192	1×10^{-3}	10	100	2.53×10^{-2}	3.18×10^{-3}	4.53×10^{-1}	1.80×10^{-1}	3*	2147.61	18.64
LDS GD	8192	1×10^{-4}	10	100	3.03×10^{-2}	6.09×10^{-3}	5.99×10^{-1}	2.29×10^{-1}	3*	2150.73	28.05
LDS GD	1024	1×10^{-2}	10	100	9.54×10^{-3}	0.00×10^0	1.18×10^{-1}	6.72×10^{-2}	1*	248.33	0.00
LDS GD	1024	1×10^{-2}	10	1000	1.90×10^{-5}	5.93×10^{-6}	5.25×10^{-4}	3.39×10^{-5}	5	252.24	1.64
LDS GD	1024	1×10^{-3}	10	100	7.34×10^{-3}	1.02×10^{-3}	1.06×10^{-1}	4.97×10^{-2}	2*	248.68	0.13
LDS GD	1024	1×10^{-3}	10	1000	2.40×10^{-5}	1.71×10^{-6}	4.00×10^{-4}	3.71×10^{-5}	5	250.24	0.87
LDS GD	1024	1×10^{-4}	10	100	7.72×10^{-3}	2.52×10^{-4}	1.13×10^{-1}	5.31×10^{-2}	2*	248.32	0.38
LDS GD	1024	1×10^{-4}	10	1000	2.01×10^{-5}	4.53×10^{-6}	5.29×10^{-4}	3.53×10^{-5}	5	250.58	1.40
LDS GD	1024	1×10^{-5}	10	100	6.77×10^{-3}	6.43×10^{-4}	1.01×10^{-1}	4.84×10^{-2}	5	250.80	0.17
LDS GD	1024	1×10^{-5}	10	1000	1.94×10^{-5}	4.92×10^{-6}	4.79×10^{-4}	3.39×10^{-5}	5	251.51	0.67
SpectraLDS	8192	1×10^{-2}	10	100	4.21×10^{-4}	1.78×10^{-4}	1.33×10^{-2}	4.60×10^{-4}	5	42.14	0.02
SpectraLDS	8192	1×10^{-2}	10	1000	1.48×10^{-5}	1.96×10^{-6}	1.23×10^{-2}	5.68×10^{-5}	5	48.11	1.39
SpectraLDS	8192	1×10^{-3}	10	100	3.59×10^{-4}	1.67×10^{-4}	1.27×10^{-2}	4.02×10^{-4}	5	42.15	0.02
SpectraLDS	8192	1×10^{-3}	10	1000	1.44×10^{-5}	4.05×10^{-6}	1.52×10^{-2}	5.81×10^{-5}	5	47.40	0.03
SpectraLDS	8192	1×10^{-4}	10	100	3.18×10^{-4}	9.33×10^{-5}	1.30×10^{-2}	3.61×10^{-4}	5	42.10	0.01
SpectraLDS	8192	1×10^{-4}	10	1000	1.88×10^{-5}	2.68×10^{-6}	1.21×10^{-2}	6.17×10^{-5}	5	48.07	1.36
SpectraLDS	8192	1×10^{-5}	10	100	4.23×10^{-4}	2.40×10^{-4}	1.38×10^{-2}	4.60×10^{-4}	5	42.13	0.01
SpectraLDS	8192	1×10^{-5}	10	1000	1.51×10^{-5}	4.74×10^{-6}	1.36×10^{-2}	5.94×10^{-5}	5	47.40	0.02
SpectraLDS	8192	1×10^{-6}	10	100	2.68×10^{-4}	7.13×10^{-5}	1.16×10^{-2}	3.07×10^{-4}	5	42.12	0.01
SpectraLDS	8192	1×10^{-6}	10	1000	1.29×10^{-5}	2.05×10^{-6}	1.32×10^{-2}	5.29×10^{-5}	5	47.43	0.03
SpectraLDS	8192	1×10^{-7}	10	100	4.36×10^{-4}	1.16×10^{-4}	1.35×10^{-2}	4.79×10^{-4}	5	42.12	0.02
SpectraLDS	8192	1×10^{-7}	10	1000	1.43×10^{-5}	3.18×10^{-6}	1.44×10^{-2}	5.61×10^{-5}	5	47.38	0.02

Table 3: SpectraLDS performance on learning synthetic linear dynamical systems with maximum eigenvalue $1 - \delta$ against a strong baseline.

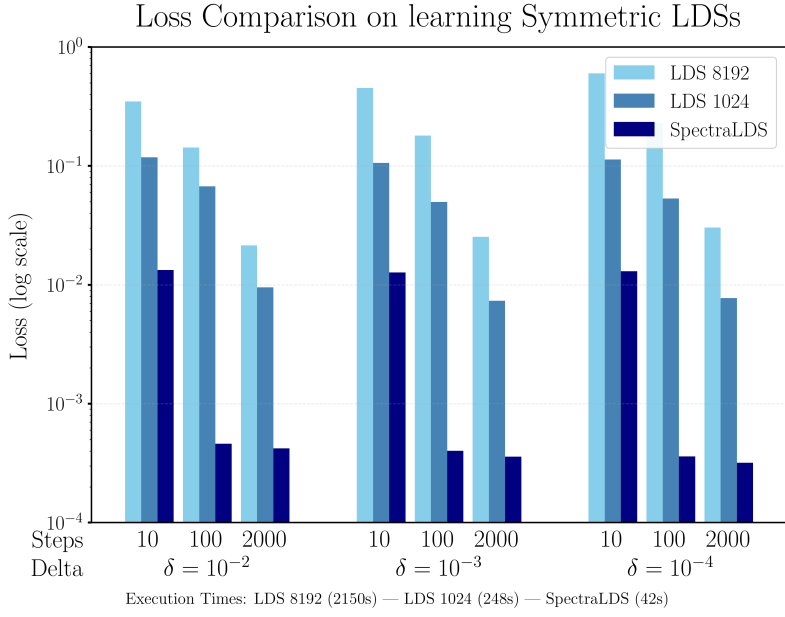


Figure 10: Comparison of learning performance on synthetic LDS tasks between SpectraLDS and a gradient-descent-updated LDS baseline. The plot shows the loss at different training steps under varying δ , input/output dimensions, and sequence lengths. SpectraLDS consistently achieves lower loss with significantly reduced runtime, especially at larger output dimensions. Full experimental details are provided in Table 3.

A.12 Layer Speeds of SpectraLDS and STU

We benchmark the inference speed of a single SpectraLDS layer across several state dimensions and compare it with two accelerated schemes for computing the STU convolution: Epoched Future Fill [1] and an STU that employs the tensor-dot approximation [22]. The SpectraLDS layer itself is produced by applying STU-to-LDS distillation to the tensor-dot STU. For the timings reported in Table 4, each model is evaluated on L (Seq. Len.) autoregressive convolutions on inputs of dimension 128, and the mean and standard deviation over five runs are recorded. All STU variants use a filter

length of 1,048,576 to accommodate the longest sequences. Although every architecture begins in the linear-scaling regime, only SpectraLDS continues to scale favorably as the sequence length increases. The resulting layer-speed comparison is visualized in Figure 11; benchmarks of SpectraLDS layers embedded in large language-model architectures appear in Appendix A.13.

Seq. Len.	Inference Time Performance (ms)									
	STU Future Fill		STU Tensor-Dot		STU		SpectraLDS (SD 100)		SpectraLDS (SD 800)	
	Mean	Std	Mean	Std	Mean	Std	Mean	Std	Mean	Std
32	30.86	1.20	31.87	6.43	134.96	22.13	21.27	3.86	18.98	0.73
64	36.52	0.72	28.92	0.42	123.16	2.54	22.40	0.14	22.27	0.29
128	49.87	0.52	40.20	6.63	135.94	2.28	29.49	0.18	29.72	0.58
256	75.73	0.35	45.98	1.12	162.74	2.90	43.54	0.27	43.62	0.47
512	136.67	7.90	70.24	0.61	211.66	2.99	71.40	0.56	73.01	0.88
1024	234.10	6.99	122.89	1.19	317.91	4.79	127.52	1.25	128.98	0.97
2048	441.09	3.69	226.47	8.02	525.65	5.09	241.16	4.43	252.58	8.78
4096	863.81	8.72	426.43	7.96	936.94	6.53	472.15	8.40	477.96	8.42
8192	1856.63	191.17	837.46	2.03	1772.30	20.46	921.55	10.23	929.06	9.76
16384	3377.29	33.16	1812.17	16.47	3434.19	30.83	1839.63	25.21	1846.04	17.43
32768	7139.35	515.40	4286.86	24.74	6721.83	19.65	3620.84	43.71	3686.90	26.45
65536	13485.25	234.22	11614.43	28.04	13478.53	89.72	7181.13	35.96	7362.73	78.96
131072	27252.83	240.34	36427.10	14.56	26703.77	323.97	14356.92	74.91	14649.56	162.32
262144	63437.26	123.25	117796.38	268.48	55775.49	275.16	28573.69	119.09	29156.42	113.55
524288	177502.89	651.96	365168.31	460.23	142896.29	621.72	57083.23	404.91	58027.04	276.39
1048576	576129.57	1425.12	1145862.83	1805.50	451607.11	721.73	114044.79	1215.07	115999.77	670.78

Table 4: Autoregressive Inference Time (ms) across model architectures (5 runs).

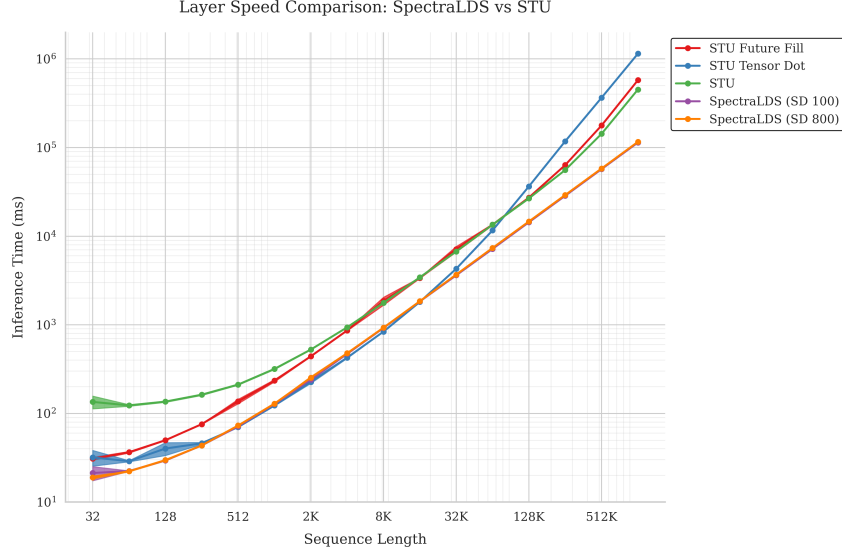


Figure 11: Autoregressive Inference Time (ms) across model architectures.

A.13 FlashSTU Ablations

A.13.1 Implementation Architecture Details

To perform the FlashSTU performance evaluations with STU-T, we employ the architecture depicted in Figure 12. We test token generation efficiency with a hybrid architecture, alternating between layers using STU-T and sliding window attention (SWA), and we additionally test with each layer using STU-T only. Each layer consists of RMS-Norm, followed by STU-T or SWA, followed by RMS-Norm, and then followed by an MLP. Inputs are tokenized by the o200k_base tokenizer and the FlashSTU model begins with an embedding layer, which is weight-tied to the output unembedding layer. To start generation, we add special tokens such as <|endoftext|> and <|endofprompt|>.

The sliding window attention layers leverage Flash Attention v2 [8][7] and ALiBi position embeddings [33]. The tested model has input dimension 896 and 12 layers, which has 550.31 million parameters for the hybrid model, and 535.99 million parameters for the STU-only model. All layers are run in bfloat16 except for the LDS distilled layers, which require float64 precision. The Flash STU-T leverages the STU with tensor-dot [22] approximation rather than the base STU layer for faster inference, and thus we perform the STU-to-LDS distillation on the STU with tensor-dot approximation. For tests with generation length up to 131072, we leverage STU filter length of 131072. For the generation length of 262144, we leverage an STU filter length of 262144. For each runtime measurement, we first run a warmup generation, before reporting the mean of two generations of that length. All benchmarks include only inference time and not model setup or filter computation costs. Additionally, each MLP layer has hidden dimension $12 \times$ the input dimension (MLP expansion factor). Other configuration details are identical to those in Table 7

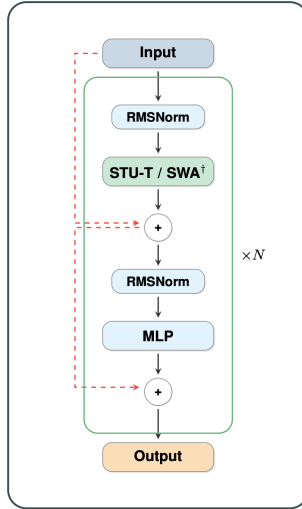


Figure 12: FlashSTU architecture [22].

A.13.2 Implementation Efficiency

Using the setup of Fig. 12, we time autoregressive generation for sequence lengths 4,096–262,144 tokens under two architectures—(i) a hybrid network that interleaves STU-T and sliding-window attention layers and (ii) an STU-only network in which every layer is STU-T. Tables 5–6 report mean runtimes (over two runs after a warmup) for SpectraLDS with state dimensions 100–800 alongside the STU-T with naive convolutions (Base STU) and the Epoch FutureFill method. We note three main results. First, SpectraLDS runtimes grow nearly linearly with sequence length and are virtually independent of the chosen state dimension. Second, while the STU-T with naive convolutions is competitive at 4–8 k tokens, it becomes progressively slower, so that SpectraLDS is $\approx 2\times$ faster by 65 k tokens and over $4\times$ faster at 262 k tokens in the STU-only setting (and $2\times$ faster in the hybrid setting). Third, Epoch FutureFill narrows the gap at medium lengths but is still outpaced by SpectraLDS beyond 131 k tokens and, in the STU-Only architecture, exhausts memory (OOM) at 262 k tokens, whereas SpectraLDS completes the run. Together these results demonstrate that SpectraLDS delivers the most favorable long-context scaling and remains robust across model hyper-parameters.

Seq. Len.	LDS SD 100	LDS SD 200	LDS SD 400	LDS SD 800	Base STU	FutureFill
4096	20.34	20.71	20.40	20.40	18.91	19.92
8192	40.48	41.32	40.69	40.82	38.09	38.39
16384	80.78	82.90	80.95	81.63	76.49	74.89
32768	161.43	163.92	162.13	163.30	164.70	141.16
65536	323.40	327.31	323.90	325.92	389.72	290.68
131072	646.66	653.96	648.99	651.35	1014.67	666.24
262144	1588.81	1639.24	1591.50	1536.20	3113.45	2498.89

Table 5: Hybrid Model Runtime (seconds) for generation across SpectraLDS with different state dimensions and Baseline and Epoched FutureFill implementations

Seq. Len.	LDS SD 100	LDS SD 200	LDS SD 400	LDS SD 800	Base STU	FutureFill
4096	14.04	13.87	13.87	13.74	12.14	19.35
8192	28.15	27.74	27.67	27.42	24.84	35.86
16384	55.71	55.51	55.34	54.73	57.32	67.96
32768	110.85	110.88	110.57	108.68	149.12	135.75
65536	220.11	221.55	221.13	215.00	429.20	329.91
131072	439.87	442.65	442.04	428.36	1352.40	952.70
262144	1104.97	1101.76	1132.18	1107.49	4672.60	OOM

Table 6: STU-Only Runtime (seconds) for generation across SpectraLDS with different state dimensions and Baseline and Epoched FutureFill implementations

A.14 Details of STU Model In Experiments

We summarize in Table 7 all relevant details for the FlashSTU model used in the language evaluations in Table 2. The distilled LDS layer used in the language benchmarking experiments was obtained by Algorithm 3 and has a state dimension of 160, incorporating both positive and negative spectral components. The weights for the distilled model were directly transferred from the FlashSTU model described below. The FlashSTU architecture is further described in Appendix A.13 and graphically shown in Figure 12.

Model Architecture		
	Description	Flash STU
Parameter Count	Total number of parameters	340M
Embedding Dimension	Dimensionality of embedding space	1024
Number of Heads	Attention heads	4
Number of Layers	Transformer + STU layers	12
ALiBi Attention	Attention scores modification using linear biases	No
RoPE Theta	RoPE scaling factor for rotary embeddings	10,000
Sliding Window Size	Sliding window attention context lookback size	512
Sequence Length (Training)	Input sequence length during training	4,096
Sequence Length (Inference)	Input sequence length during inference via position interpolation	131,072
Vocabulary Size	Size of the model's vocabulary	200,064
MLP Expansion Factor	Expansion factor in MLP layers	4
Bias	Use of bias terms in linear layers	No
Dropout	Dropout rate	0.0
Number of Filters	Number of filters	24
Training and Optimization		
Epochs	Number of training epochs	1
Global Batch Size	Number of tokens processed per step	524,288
Micro Batch Size	Batch size per GPU	8
Gradient Accumulation Steps	Number of steps before performing a gradient update	8
Warmup Steps	Number of warmup steps	1,907
Evaluation Period	Evaluation frequency (steps)	50
Max Grad Norm	Maximum gradient norm for clipping	1.0
Optimizer Configuration		
Optimizer	Optimizer type	AdamW
Learning Rate Schedule	LR scheduling strategy	Linear decay with warmup
Max Learning Rate	Maximum learning rate	4.0×10^{-4}
Min Learning Rate	Minimum learning rate	4.0×10^{-5}
Torch Dtype	Data type for PyTorch tensors	<code>bfloat16</code>
Betas	Optimizer betas	(0.9, 0.999)
Epsilon	Optimizer epsilon	1.0×10^{-8}
Weight Decay	Weight decay factor	1.0×10^{-2}
AMSGrad	Use AMSGard variant	No
Fused	Use fused optimizer	Yes
Optimization Techniques		
Activation Checkpointing	Enable activation checkpointing	Yes
Use Flash FFT	Enable Flash FFT	No
Use Tensordot Approx.	Enable tensordot approximation	Yes
Use Attention	Enable attention mechanism	Yes
Softcap	Softcap threshold	50.0
Torch Compile	Enable Torch compile optimization	Yes

Table 7: Model and training configuration details for the 340M Flash STU model.

Benchmarks were conducted with task-specific few-shot configurations. For each task, if the few-shot value is not set to -1, that number of examples is provided during evaluation; otherwise, the few-shot setting is left unchanged. Specifically, the few-shot settings are as follows: HellaSwag: 0 shots, MMLU: 5 shots, PIQA: 0 shots, BoolQ: 0 shots, Winogrande: -1, Commonsense_QA: 7 shots, OpenBookQA: -1, ARC-Easy: -1, ARC-Challenge: -1. The model employs `bfloat16` precision except for the LDS computations, where it requires `float64` precision, and a global batch size of 3 examples per task. All computations were performed on a single H100 GPU.

A.15 Example Generation

The following is an example completion from the FlashSTU 340M model after LDS distillation with temperature 1.0:

Prompt: After school each day, Deidre Smith would look over the essays her students submitted at the end of class and, without fail, each day Alex would surprise her. Today, he wrote about

Generated Text: After school each day, Deidre Smith would look over the essays her students submitted at the end of class and, without fail, each day Alex would surprise her. Today, he wrote

about 40 essays from his freshman class, all of which were collected over the past three years for *The Book of Sentiments: What Students Should Do When Reading Literature* – First published in 1962, this book was very well received. What people who read this book would not immediately accept or appreciate the author's thought-provoking message of what it means to read literature.

What exactly are you doing with your newfound ability to read? You are reading? What emotions do you feel at the time? What do you do with your newfound ability to read? These are just a few examples of how much reading students think they can gain from the experience. It is no accident, then, that each and every level of student will have difficulty with the text in this book. Students are given a choice: choose an assignment that you are comfortable with and one that you would like to read more deeply and critically. As the task to read becomes more difficult and the pressure for it is much greater, students will struggle to put their best minds together when reading literature. To help students overcome this hurdle, I have devised a book that makes the task a bit easier by providing all the information they need from a reader. If a reader is too afraid to read, I have designed a page that provides guidance from the textbook that will...