RAPID BAYESIAN IDENTIFICATION OF SPARSE NONLINEAR DYNAMICS FROM SCARCE AND NOISY DATA

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ABSTRACT

We propose a fast probabilistic framework for identifying differential equations governing the dynamics of observed data. We recast the SINDy method within a Bayesian framework and use Gaussian approximations for the prior and likelihood to speed up computation. The resulting method, Bayesian-SINDy, not only quantifies uncertainty in the parameters estimated but also is more robust when learning the correct model from limited and noisy data. Using both synthetic and real-life examples such as Lynx-Hare population dynamics, we demonstrate the effectiveness of the new framework in learning correct model equations and compare its computational and data efficiency with existing methods. Because Bayesian-SINDy can quickly assimilate data and is robust against noise, it is particularly suitable for biological data and real-time system identification in control. Its probabilistic framework also enables the calculation of information entropy, laying the foundation for an active learning strategy.

1 Introduction

The pursuit of direct model equation discovery has been an ongoing and significant area of interest in scientific machine learning. The popular *sparse identification of nonlinear dynamics* (SINDy) framework [1] offers a promising approach to extract parsimonious equations directly from data. SINDy's promotion of parsimony by sparse regression allows for the identification of an interpretable model that balances accuracy with generalizability, while its simplicity leads to a relatively efficient and fast learning process compared to other machine learning techniques. The framework has been successfully applied in a variety of applications, such as model idenficiation in plasma physics [2], control engineering [3, 4], biological transport problems [5], socio-cognitive systems [6], epidemiology [7, 8] and turbulence modelling [9]. Furthermore, its remarkable extendibility has attracted a range of modifications, including the adaptation to discover partial differential equations [10], the extension to libraries of rational functions [11], the integration of ensembling techniques to improve data efficiency [12] and the use of weak formulations [13, 14] to avoid noise amplification when computing derivatives from discrete data.

One major difficulty in using scientific machine learning methods in fields such as biophysics, ecology, and microbiology, is that measured data from these fields is often noisy and scarce. This limitation renders many data-intensive machine learning techniques infeasible. Learning speed is also a major requirement in other real-time applications, such as control. Although SINDy is less data-intensive and faster than most machine learning techniques, it is nonetheless susceptible to noise when the dataset is scarce. In some cases, SINDy may have difficulties recovering accurate and

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parsimonious models if the data is too scarce and noisy. To address the challenge of fast and robust learning in a noisy and scarce dataset, we present Bayesian-SINDy. This method combines sparse Bayesian learning and SINDy to extract parsimonious governing equations directly from the data. It accounts for the uncertainty in the data and the estimated parameters when selecting parsimonious models, but does so without the need for expensive sampling computations such as Markov Chain Monte Carlo (MCMC), making it particularly suitable for real-time applications in which fast computation is required.

To appreciate the potential improvement of SINDy's performance in scarce and noisy data using sparse Bayesian learning, one must first understand the principles of SINDy. Essentially, SINDy approaches equation discovery as a sparse dictionary learning problem. First, it constructs a dictionary consisting of all potential candidate functions of the input data. Then, SINDy employs sparse regression to identify the most pertinent functions that describe the dynamics and to eliminate the unnecessary ones, yielding parsimonious dynamical equations that retain only the relevant terms. The original study [1] established the Sequentially Thresholded Least Squares (STLS) algorithm as a method for sparse regression with minimal computational cost. STLS assumes that all relevant terms possess coefficients exceeding a certain threshold, allowing for the elimination of irrelevant terms with coefficients below this threshold. Hence, the imposed threshold is effectively a hyperparameter one can tune to control how aggressively sparsity is promoted. Besides STLS, there are other regularisation and relaxation techniques that promote sparsity during regression, such as LASSO [15] and SR3 [16]. Nevertheless, nearly all sparse regression techniques contain hyperparameters that require fine-tuning. To properly select hyperparameters, the Pareto front is commonly employed as a heuristic method to ascertain the appropriate degree of sparsity. However, interpreting the front can be ambiguous, especially in cases where the front lacks a distinct elbow point. The later introduction of Akaike/Bayesian information criterion (AIC/BIC) in the model selection process [17] provides certain remedies to the problem, but they only work well in the high data limit [18].

Another prevalent problem with existing SINDy-like algorithms is their ability to handle scarce and noisy data. While SINDy-based approaches are effective at identifying equations from synthetic data with little noise, real-world measurements are typically scarce and prone to significant measurement error. Large noise in the data presents several challenges to SINDy. First, measurement noise corrupts the precise recovery of time derivatives because conventional finite difference approximations can amplify noise by the order of the sample rate. This problem can be mitigated by using a weak formulation [13, 14]. However, this approach may also introduce bias to the data, as we will illustrate in §2.3. Second, excessive noise in the data can impede STLS's ability to select the most appropriate model. STLS' poor performance in noisy and scarce data is partly due to the fact that it does not account for the uncertainty in the data, even though this uncertainty can be a crucial factor to consider when choosing a model. Furthermore, the performance of STLS deteriorates when the number of data points is lower than a certain threshold. As we will demonstrate later in §4.1, by adopting a Bayesian viewpoint, one can better find the reason for such deterioration and the remedy against it, effectively addressing the constraints posed by scarce and noisy data.

In the Bayesian point of view, in contrast to STLS, models are selected not by the magnitude of the coefficients of each term but by the marginalised likelihood (i.e. evidence), which is proportional to the probability of the selected model given the data. This procedure of selecting the model that maximises the evidence is also known as type-II maximum likelihood and is arguably a better method to evaluate how suitable the model is to describe the data than STLS. Firstly, this method naturally avoids overfitting and promotes sparsity, as the evidence naturally includes an 'Occam factor' that penalises over-parameterised models [19]. Compared to the use of the Pareto front or information criterion to tune hyperparameters, the Bayesian approach provides a more natural way to determine the parsimony of the model according to the level of uncertainty in the data. The Bayesian account of uncertainty in the data also makes it more resilient against large noise. Secondly, model selection in this Bayesian framework is agnostic to the magnitude of the coefficients. As a result, it can effectively acquire terms with small coefficients, which is challenging for STLS. Finally, it is well known that many sparse regression techniques and information criteria can be re-interpreted from a Bayesian perspective [20, 21]. For instance, LASSO can be considered as the maximum a posteriori estimate under a Laplace prior, whereas thresholding is equivalent to empirically removing terms that have little significance to the likelihood. The BIC is rooted in the Bayesian framework and can be regarded as an asymptotic estimate of the marginalised likelihood at the high data limit, whereas the AIC can be seen as an extension of the BIC with a different prior [18].

Therefore, the goal of this work is to demonstrate how a Bayesian perspective can encompasses all preceding techniques and is best suited for extending the current SINDy techniques to function in challenging data environments where data are noisy and scarce. As we shall demonstrate later in §4.1, by grounding SINDy in the Bayesian framework, we can also more rigorously justify how and why SINDy works well in the high data low noise limit.

1.1 Related work and contribution of this work

Sparse Bayesian learning (or regression) is a mature and well-established method in the statistical literature [19, 22, 23]. The dictionary learning technique typically employs a sparsity-inducing prior to promote sparsity. The current 'gold standard' sparsity-inducing priors are the spike-and-slab-typed priors, but earlier work has also investigated the use of hierarchical priors. For example, when one combines a Gaussian prior with variance α^{-1} with a maximum-entropy hyperprior on α (i.e. a scale uniform distribution), the result is a sparsity-inducing student-t true prior that promotes sparsity. A notable early study that utilises this hierarchy prior to promote sparsity is the RVM/SparseBayes package developed by Tippings & Faul [24]. The Relevance Vector Machine (RVM) is a type-II-maximum-likelihood algorithm that was initially created as a probabilistic alternative to the Support Vector Machine. It utilises kernels as the dictionary and aims to maximise evidence while also encouraging sparsity in dictionary learning. This method exploits the fact that the conditional hyperprior given the data can be approximated as a Dirac Delta, while the rest of the probability distributions can be approximated as Gaussians, which allows for analytical tractability. The resulting approach has been demonstrated to be both computationally efficient and robust, not only for classification tasks but also for addressing general sparse regression problems. Due to its Bayesian foundation, it is also resilient to noisy and scarce datasets. Therefore, it is especially appropriate for the objective of this study.

We note that this work is not the first to apply sparse Bayesian regression in the context of equation discovery or system identification [25, 26, 27], nor the first to integrate sparse Bayesian regression with SINDy [28, 4]. For instance, Zhang & Lin [28] incorporated an additional thresholding procedure into the SparseBayes method and applied the technique to extract different ordinary differential equations (ODEs) and partial differential equations (PDEs) from numerical data. They conducted a comparative analysis of their threshold sparse Bayesian regression algorithm with STLS and LASSO, and demonstrated Bayesian regression's superior performance in the presence of noisy data. Recent work by Niven et al. [27] has also attempted to use Bayesian inference for model selection and uncertainty quantification in system identification. However, the synthetic dataset they considered only introduced uncertainty in the time derivatives and not in the state variables themselves, which can oversimplify the challenge of the system identification problem, as we shall demonstrate later in (9). Fuentes et al. [4] substituted STLS with SparseBayes directly as the sparse regression algorithm and employed the technique for system identification. In particular, apart from synthetic data, they have successfully applied this approach to two experimental benchmarks. Nevertheless, the dataset they used has a high sampling rate (i.e. a high data limit). When finite difference was applied, the noise was significantly amplified in their time derivative, resulting in unreasonably large uncertainty in their learning result despite successfully recovering the desired equations.

While the above work uses variants of SparseBayes, which do not require expensive computation, they rely on Gaussian approximations of the likelihood and prior distribution in order to maintain the analytical tractability of the distributions. In contrast, subsequent works [29, 30, 31, 32] use spike-and-slab-typed priors, which, despite the more accurate recovery of the correct model, necessitates costly sampling techniques, such as MCMC [29, 31] or Approximated Bayesian Computation [30] to capture the distribution of the estimated parameter more accurately. The benefit of using these sampling-based techniques lies in their capacity to precisely capture non-Gaussian distributions in the parameter and their ability to integrate sparsity-promoting spike-and-slab-typed prior. However, they come at the expense of expensive computations. A cheaper alternative is to make use of ensembling technique [12] to estimate uncertainty in the model. Recent work [33] has demonstrated the convergence of the bootstrapping-based STLS algorithm to the equivalent Bayesian statistics. Nonetheless, these algorithms remain an order of magnitude more expensive than STLS and the Gaussian approximations used by SparseBayes.

Given that our goal is to seek a fast and computationally inexpensive learning method, this work will follow the earlier work more closely [28, 4] and shall adopt strategies similar to the SparseBayes method. Despite its similarity with the earlier work, we note a few key differences. First, our work highlights the key strength of the Bayesian framework, namely, its superior performance in scarce and noisy data. This is particularly in contrast to [4], whose dataset is in the high data limit. Second, our work compares the performance of the weak formulation and finite difference in obtaining derivatives from scarce and noisy data and highlights the pros and cons of each method, which is in contrast to the finite difference used in earlier work [28, 4]. Lastly, we shall present a new sparse Bayesian regression algorithm, which, despite heavy inspiration from SparseBayes, gives better performance due to its ability to keep track of how noise variance propagates throughout the calculation.

1.2 Structure of the paper

The rest of the paper is structured as follows. In §2, we will provide a concise overview of the SINDy framework. The significance of noise will be emphasised, particularly in terms of its propagation through the collection of potential functions and the computation of temporal derivatives. Then, we will introduce our Bayesian approach, which takes into account the propagation of noise, and briefly compare our Bayesian algorithm with previous studies. We will also

briefly highlight the difficulty of approximating time derivatives using weak formulations when the sampling rate is low, which further reinforces our case for a noise-robust algorithm. In §3, we will showcase the efficacy of the suggested framework by presenting examples such as the Van Der Pol system, the nonlinear oscillator, the Lorenz system, and predator-prey dynamics. The predator-prey dynamics are based on real-life data, highlighting the practicality of the proposed framework in real-world applications. Then, in §4, we will analyse the reasons why the Bayesian approach is more resilient in scarce and noisy data and highlight some limitations of the proposed algorithm. In particular, in §4.1, we will explain why SINDy works in the high data limit from a Bayesian perspective, but not as well as its Bayesian counterpart in the low data limit. Finally, we will discuss how the improved data efficiency and the probability framework create new potential innovations in active learning.

2 Methodology

2.1 SINDy and tracing noise propagation

Suppose we have a system of interest with state variables $\hat{\mathbf{x}} = [\hat{x}_1(t), \ \hat{x}_2(t) \ ... \ \hat{x}_D(t)] \in \mathbb{R}^{1 \times D}$ with dimension D, governed by the dynamical equation

$$\dot{\hat{\mathbf{x}}} = \frac{d}{dt}\hat{\mathbf{x}} = f(\hat{\mathbf{x}}) \tag{1}$$

with some unknown function $f: \mathbb{R}^D \to \mathbb{R}^D$. The system was observed \tilde{N} times at $t = t_1, \ t_2, \ ...t_{\tilde{N}}$ and the state variables are measured, resulting in the dataset

$$\mathbf{X} = \begin{bmatrix} \mathbf{x}_{1} & \mathbf{x}_{2} & \dots & \mathbf{x}_{D} \end{bmatrix} = \begin{bmatrix} x_{1}(t_{1}) & x_{2}(t_{1}) & \dots & x_{D}(t_{1}) \\ x_{1}(t_{2}) & x_{2}(t_{2}) & \dots & x_{D}(t_{2}) \\ \vdots & \vdots & \ddots & \vdots \\ x_{1}(t_{\tilde{N}}) & x_{2}(t_{\tilde{N}}) & \dots & x_{D}(t_{\tilde{N}}) \end{bmatrix} \in \mathbb{R}^{\tilde{N} \times D}.$$
 (2)

Due to measurement error, the time series data might be contaminated by some noise ϵ_x , such that

$$X_{ij} = x_j(t_i) = \hat{x}_j(t_i) + \epsilon_{x,ij}. \tag{3}$$

For simplicity, here we assume that the noise $\epsilon_x \in \mathbb{R}^{\tilde{N} \times D}$ consists of $\tilde{N} \times D$ independent and identically distributed random variables with Gaussian distributions, with zero means and variance $\sigma_x^2 \in \mathbb{R}^{\tilde{N} \times D}$.

The goal of SINDy is to discover f from the observed time series data X. SINDy postulates that f can be written as a linear combination of a library of candidate functions. For example, a commonly used library is the polynomial library

$$\boldsymbol{\Theta}(\hat{\mathbf{x}}) = \begin{bmatrix} 1 & \hat{x}_1 & \hat{x}_2 & \hat{x}_1^2 & \hat{x}_1\hat{x}_2 & \hat{x}_2^2 & \hat{x}_1^3 & \ldots \end{bmatrix},$$

or when applied to the observed dataset,

$$\Theta(\mathbf{X}) = \begin{bmatrix} \mathbf{1} & \mathbf{x}_1 & \mathbf{x}_2 & \mathbf{x}_1^2 & \mathbf{x}_1\mathbf{x}_2 & \mathbf{x}_1^2 & \mathbf{x}^3 & ... \end{bmatrix} \in \mathbb{R}^{\tilde{N} \times M}$$

in which we assume the library consists of M columns of possible candidates. Here, the product and power operations $\mathbf{x}^n \mathbf{y}^m = [x_i^n y_i^m]$ are applied elementwise.

If f can be described by a few terms in the library of candidate functions, then the parameter vector w that fulfils

$$\dot{\hat{\mathbf{x}}}_c = \mathbf{\Theta}(\hat{\mathbf{x}})\mathbf{w} \tag{4}$$

will be sparse. The aim of SINDy is to recover this sparse parameter vector \mathbf{w} through regression. If f cannot be fully described by a linear composition of library candidate functions, but can be approximated by it, then SINDy finds a parsimonious approximation to f through sparse regression.

However, in practice, the time derivative is usually not measured as part of the dataset. To recover the time derivative from the time-series data X, we assume there exist some linear operators L_I and $L_{\partial t}$ such that

$$L_I \dot{\mathbf{X}}_i \approx L_{\partial t} \mathbf{X}$$
, and therefore $L_{\partial t} \mathbf{X} \approx L_I f(\mathbf{X})$ (5)

in the absence of noise, i.e. when $\epsilon_x = 0$. Note that L_I and $L_{\partial t}$ have dimensions $\hat{N} \times N$. Further discussion on these operators for recovering time derivatives will follow in §2.3.

Plugging (3) and (5) into (4), we rewrite the regression as

$$L_{\partial t}\mathbf{X} = L_I\mathbf{\Theta}(\mathbf{X})\mathbf{w} + \boldsymbol{\epsilon}. \tag{6}$$

Here, we see that the noise is transformed by the operations on ϵ_x , giving rise to the total noise $\epsilon = \epsilon_{\Theta} + \epsilon_{\partial t}$, which consists of $\epsilon_{\partial t}$, the noise arising from the time derivative, and ϵ_{Θ} , the noise arising from the nonlinear library. Since $L_{\partial t}$ is a linear operator on \mathbf{X} , assuming $L_{\partial t}^2$ represents element-wise squaring of the matrix representing $L_{\partial t}$, the variance of $\epsilon_{\partial t}$ is

$$\sigma_{\partial t}^2 = L_{\partial t}^2 \sigma_x^2 \tag{7}$$

and $\epsilon_{\partial t} \sim \mathcal{N}(0, \sigma_{\partial t}^2)$ remains Gaussian. However, $\Theta(\mathbf{X})$ consists of nonlinear functions of \mathbf{X} , so the resulting distribution of ϵ_{Θ} is non-Gaussian and probably not tractable analytically. Nonetheless, one can trace the variance of each nonlinear function through the algebra of random variables, giving rise to a library of variance,

$$\boldsymbol{\sigma}_{\Theta}^{2}(\mathbf{x}, \boldsymbol{\sigma}_{x}^{2}) = \begin{bmatrix} \boldsymbol{\sigma}_{1}^{2} & \boldsymbol{\sigma}_{x_{1}}^{2} & \boldsymbol{\sigma}_{x_{2}}^{2} & \boldsymbol{\sigma}_{x_{1}^{2}}^{2} & \boldsymbol{\sigma}_{x_{1}x_{2}}^{2} & \boldsymbol{\sigma}_{x_{2}^{2}}^{2} & \boldsymbol{\sigma}_{x_{3}^{3}}^{2} & \dots \end{bmatrix}$$
(8)

where each column is associated with each column in the library of candidate functions. Here, we further approximate that the resulting distribution from each column (function) in $\Theta(\mathbf{X})$ are mutually independent, and independent of $\dot{\mathbf{X}}$. Under these assumptions, the variance of the total noise ϵ can be written as

$$\sigma^2 = L_I^2 \sigma_{\Theta}^2(\mathbf{x}, \sigma_x^2) \mathbf{w}^2 + L_{\partial t}^2 \sigma_x^2, \tag{9}$$

where $\mathbf{w}^2 = [w_{ij}^2]$ is the elementwise square of \mathbf{w} and L_I^2 the elementwise square of L_I .

Therefore, according to (9), the noise variance σ^2 depends on the parameter w. This dependency was omitted in previous work [28, 27, 4], but this work will address this dependency in the regression step. It is also worth reiterating that even though we have used the algebra of random variables to keep track of the noise variance, the final distribution of ϵ^2 is expected to be non-Gaussian. Nonetheless, we shall later approximate it as a Gaussian distribution in order to preserve analytical tractability.

For simpler notation, we define the dependent variable in (6) as

$$\mathbf{t} = L_{\partial t} \mathbf{X} \in \mathbb{R}^{N \times D} \tag{10}$$

and the dictionary, which is the library of functions of the independent variable X, as

$$\mathbf{D}(\mathbf{X}) = L_I \mathbf{\Theta}(\mathbf{X}) = [\mathbf{d}_1 \quad \mathbf{d}_2 \quad \mathbf{d}_3 \quad \dots \quad \mathbf{d}_m \quad \dots \quad \mathbf{d}_M] \in \mathbb{R}^{N \times M}$$
(11)

Substituting the redefined variables t and D into (6), the sparse regression problem becomes

$$\mathbf{t} = \mathbf{D}\mathbf{w} + \boldsymbol{\epsilon} \tag{12}$$

in which $\epsilon \sim \mathcal{N}(0, \boldsymbol{\beta}^{-1})$ is approximated to be a Gaussian noise with zero mean and elementwise variance $\boldsymbol{\beta}^{-1} = \boldsymbol{\sigma}^2$. Here, we use the inverse noise variance $\boldsymbol{\beta}^{-1}$ instead of $\boldsymbol{\sigma}^2$ to facilitate easier notation in the next section.

2.2 Fast Bayesian Regression under the Gaussian assumption

In some fields, such as control engineering, the Bayesian approach is avoided because accurate calculations of the statistical distributions are analytically intractable, forcing one to resort to expensive Monte Carlo simulations or other sampling techniques. In particular, when the dictionary $\mathbf{D}(\mathbf{X})$ consists of nonlinear functions of \mathbf{X} , the distribution of each entry in $\mathbf{D}(\mathbf{X})$ is probably non-Gaussian. Nonetheless, if we accept the Gaussian approximation in the previous section, then the likelihood can be approximated as Gaussian. Combining the Gaussian likelihood with a Gaussian prior, one can maintain analytical tractability throughout the calculation. This is effectively how SparseBayes obtained its fast computational efficiency.

In the following section, we will introduce the Bayesian view of regression and derive our regression method alongside SparseBayes. For pedagogical reasons, in this section we will focus on the regression of a single state variable, i.e. a single column in \mathbf{t} (and \mathbf{w}). One can easily extend the method to multiple dimensions later by repeating the regression for each state variable.

A Bayesian view of regression is as follows. In the Bayesian framework, instead of seeking \mathbf{w} that best fits (12), one seeks the posterior distribution of \mathbf{w} conditioned on the dataset (\mathbf{D}, \mathbf{t}) , which can be calculated using the Bayes rule

$$p(\mathbf{w}|\mathbf{t}, \mathbf{D}, \boldsymbol{\beta}, \boldsymbol{\alpha}) = \frac{p(\mathbf{t}|\mathbf{D}, \mathbf{w}, \boldsymbol{\beta})p(\mathbf{w}|\boldsymbol{\alpha})}{p(\mathbf{t}|\mathbf{D}, \boldsymbol{\beta}, \boldsymbol{\alpha})}.$$
(13)

Here, $p(\mathbf{t}|\mathbf{D}, \mathbf{w}, \boldsymbol{\beta})$ is the likelihood and $p(\mathbf{w}|\boldsymbol{\alpha})$ is the prior distribution with the hyperparameter $\boldsymbol{\alpha} \in \mathbb{R}^M$, elements of which define the inverse of the prior variance of each element in \mathbf{w} . The evidence $p(\mathbf{t}|\mathbf{D}, \boldsymbol{\beta}, \boldsymbol{\alpha}) =$

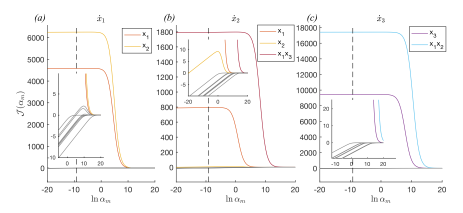


Figure 1: Contribution of the m-th term in the model to the evidence at given prior variance α_m . Data are generated from a Lorenz system, with noise $\sigma_x=0.25$ added to 125 data points. Terms in the model are highlighted by their respective colours, whereas terms that should not be in the model are grey. The inset shows the zoomed-in picture of the same graph near $\mathcal{J}(\alpha_m)=0$. Notice that terms that should be included have dramatically positive $\mathcal{J}(\alpha_m)$, except for x_2 in (b), which has a relatively small but positive contribution. Also, in the inset of (a), some grey lines are slightly above zero in small regions of α_m , but remain negative when α_m is small enough. For our method, α_m is fixed at a small value, as shown by the black dashed line.

 $\int p(\mathbf{t}|\mathbf{D},\mathbf{w},\boldsymbol{\beta})p(\mathbf{w}|\boldsymbol{\alpha})d\mathbf{w}$ normalises the posterior distribution. If we assume that the prior is Gaussian with variance $\boldsymbol{\alpha}^{-1}$, i.e. $p(\mathbf{c}|\boldsymbol{\alpha}) \sim \mathcal{N}(\mathbf{0},\alpha^{-1})$, and the likelihood can be approximated as Gaussian under the assumption of $\epsilon \sim \mathcal{N}(0,\boldsymbol{\beta}^{-1})$, then all of the above distributions are Gaussians and can be written out analytically. In particular, the log-evidence can be written as [24]

$$\mathcal{J}(\mathbf{t}|\mathbf{D},\boldsymbol{\beta},\boldsymbol{\alpha}) = -\frac{1}{2}(\ln|2\pi\mathbf{C}| + \mathbf{t}^T\mathbf{C}^{-1}\mathbf{t}). \tag{14}$$

where

$$\mathbf{C} = \mathbf{B}^{-1} + \mathbf{D}\mathbf{A}^{-1}\mathbf{D}^{T} = \mathbf{B}^{-1} + \Sigma_{m=1}^{M} \frac{1}{\alpha_{m}} \mathbf{d}_{m} \mathbf{d}_{m}^{T} \in \mathbb{R}^{N \times N},$$
(15)

where **A** is a diagonal matrix with entries $A_{mm} = \alpha_m$, **B** is a diagonal matrix with entries $B_{nn} = \beta_n$, and $|\mathbf{C}|$ denotes the determinant.

Now, during regression, one can perform model selection by selecting the columns \mathbf{d}_m in \mathbf{D} that should be included in the model while removing columns that should not be in the model. In STLS, columns are removed by setting the corresponding w_m to zero if it is smaller than a certain threshold. In a Bayesian framework, models are compared by the evidence, so we seek to maximise the log-evidence $\mathcal{J}(\mathbf{t}|\mathbf{D},\boldsymbol{\beta},\boldsymbol{\alpha})$ by removing columns from \mathbf{D} . In the calculation in (15), this is equivalent to setting the corresponding elements in $\boldsymbol{\alpha}$ to infinity. Now, it is interesting to isolate how each element α_m and the corresponding column \mathbf{d}_m changes $\mathcal{J}(\mathbf{t}|\mathbf{D},\boldsymbol{\beta},\boldsymbol{\alpha})$. By defining

$$\mathbf{C}_{-m} = \mathbf{C} - \frac{1}{\alpha_m} \mathbf{d}_m \mathbf{d}_m^T, \tag{16}$$

and using the Sherman-Morrison formula, we isolate the contribution of the m-th term $\mathcal{J}(\alpha_m)$ from the evidence without m-th term $\mathcal{J}(\alpha_{-m})$, i.e. $\mathcal{J}(\mathbf{t}|\mathbf{D},\boldsymbol{\beta},\boldsymbol{\alpha})=\mathcal{J}(\alpha_{-m})+\mathcal{J}(\alpha_m)$, and find that

$$\mathcal{J}(\alpha_m) = \frac{1}{2} \left(\ln \alpha_m - \ln \left(\alpha_m + \mathbf{d}_m^T \mathbf{C}_{-m}^{-1} \mathbf{d}_m \right) + \frac{(\mathbf{d}_m^T \mathbf{C}_{-m}^{-1} \mathbf{t})^2}{\alpha_m + \mathbf{d}_m^T \mathbf{C}_{-m}^{-1} \mathbf{d}_m} \right). \tag{17}$$

Figure 1 shows the typical $\mathcal{J}(\alpha_m)$ for the correct terms that should be included in a Lorenz system (colored) and the incorrect terms that should not be in the equations (c.f. (23)) when the algorithm is trying to learn them from noisy data of the system. Asymptotically, $\mathcal{J}(\alpha_m)$ always approaches zero when $\alpha_m \to \infty$, because it is equivalent to not including the m-th term in the model. Meanwhile, as $\alpha_m \to -\infty$, $\mathcal{J}(\alpha_m)$ approaches negative infinity, which is equivalent to setting prior and evidence vanishingly small. Note that, for all incorrect terms, the corresponding $\mathcal{J}(\alpha_m) \leq 0$ for all $\alpha_m \in \mathbb{R}$, implying that the inclusion of the term will always decrease evidence. Meanwhile, for the correct terms, $\mathcal{J}(\alpha_m) > 0$ in a large range of α_m and $\mathcal{J}(\alpha_m)$ inevitably exhibits a peak at

$$\alpha_m = \frac{(\mathbf{d}_m^T \mathbf{C}_{-m}^{-1} \mathbf{d}_m)^2}{(\mathbf{d}_m^T \mathbf{C}_{-m}^{-1} \mathbf{t})^2 - \mathbf{d}_m^T \mathbf{C}_{-m}^{-1} \mathbf{d}_m}.$$
(18)

Therefore, one can determine if a term should be included in the model by observing if $\mathcal{J}(\alpha_m) > 0$ for some α_m , which can be determined by $(\mathbf{d}_m^T \mathbf{C}_{-m}^{-1} \mathbf{t})^2 - \mathbf{d}_m^T \mathbf{C}_{-m}^{-1} \mathbf{d}_m > 0$. However, it should be noted that $\mathcal{J}(\alpha_m)$ depends on \mathbf{C}_{-m} , which changes according to the existence of other terms in the model. As other columns $\mathbf{d}_{i\neq m}$ are added or subtracted from the model, $\mathcal{J}(\alpha_m)$ for the column \mathbf{d}_m will need to be recomputed.

The SparseBayes algorirthm exploits this observation and devises an iterative algorithm that optimises each element in α to maximise $\mathcal{J}(\alpha_m)$, and removes incorrect terms that do not contribute to evidence positively, thereby promoting sparsity. Another interpretation is that this algorirthm effectively makes the Gaussian prior a sparsity-promoting spike-and-slab prior. Since the algorithm always seeks to maximise $\mathcal{J}(\alpha_m)$ for each α_m , incorrect terms will have the corresponding α_m optimised to $\alpha_m \to \infty$, effectively making the prior a spike at zero, while terms that should be included in the model will have a finite α_m value, which is effectively a slab prior. As for β , the SparseBayes algorithm accepts a prescribed value σ^2 for all data points, i.e. $\mathbf{B} = \sigma^{-2}\mathbf{I}$. Alternatively, the algorithm can optimise a single σ^2 value for maximum evidence, but it does not accept σ^2 varying across different time points.

Here, we present a novel Bayesian regression approach that also maximises evidence, but differs from SparseBayes in the optimisation of α and β . Phenomenologically, we noticed that in some cases, incorrect terms have $\mathcal{J}(\alpha_m)$ values exceeding zero in some small regions of α , which might be attributed to incidental matching between spurious terms and noise. SparseBayes often mistakenly captures these spurious terms, usually with small coefficients. Zhang & Lin [28] addressed this problem by incorporating a thresholding procedure following the regression performed by the SparseBayes algorithm. In contrast, this study introduces a technique that circumvents the optimisation process of α entirely. Instead, based on the observation that $\mathcal{J}(\alpha_m)$ for the correct terms is always positive for a large range of α_m , we set α to a certain low value, ensuring that $\mathcal{J}(\alpha_m)$ remains positive for the correct terms, while becoming negative for the incorrect ones. The value of α can often be approximated by expert prior knowledge or physical scaling argument, and it represents our prior knowledge of the system. Often the value of α is not critical, as long as it is small enough to ensure that $\mathcal{J}(\alpha_m)$ remains positive for the correct terms. In another viewpoint, α is effectively a tunable hyperparameter in our proposed method.

Another distinguishing feature of our method compared to SparseBayes and previous works [28, 27, 4] is its treatment of the noise inverse variance β . Previous methods assumed that the noise level is the same across all data points, but this assumption may impede the algorithm's ability to learn the optimal outcome. Instead, (15) showed a more precise description of the noise variance in the context of learning dynamic equations from time-series data. The formulation requires the estimated parameter w *a priori*, but its value remains unknown until after the regression. Hence, we propose an algorithm for an iterative technique to determine β by considering the w-dependent component of equation (9) as a correction. Initially, we conduct regression under the assumption that $\mathbf{w} = 0$, which is consistent with the prior. Following each regression, we modify the value of $\beta = \sigma^{-2}$ based on the value of \mathbf{w} from the last iteration and equation (9). Subsequently, we iterate the regression process until the modification to β falls below a specific numerical threshold. Generally, convergence can be achieved within 3-5 iterations.

In order to find the optimal model, specifically the model with maximum evidence, we suggest using a Greedy algorithm. Starting from the full model comprising M terms, we assess the potential increase in evidence by eliminating each individual term. Subsequently, we choose M-1 terms that yield the greatest increase in evidence, and iterate this procedure until any further increase in evidence is impossible. The technique of our proposed method is outlined in algorithm 1, which combines our search strategy to maximise evidence with the aforementioned noise iteration.

The computational complexity of this algorithm scales with the number of terms in the library M by $\mathcal{O}(M^2/2)$. Although this is slower than the STLS method, it is still much faster than a brute-force search $(\mathcal{O}(2^M))$ and any sampling technique. Furthermore, our elimination strategy is more robust than the iterative procedure employed by SparseBayes. In the SparseBayes algorithm, the search begins with a single term and additional terms are later added based on the prospective rise in evidence. Although this technique can be more computationally efficient, its inclusion of both addition and deletion of terms may result in occasional entrapment in a loop where collinear terms are repeatedly added and eliminated. Our elimination technique, in contrast, ensures convergence within $M^2/2$ iterations.

Algorithm 1: Greedy algorithm for evidence maximisation (with noise-parameter iteration)

```
input: Library D(X), dynamics t, library noise variance L_I \sigma_{\Theta}^2, dynamics noise variance \sigma_{\partial t}^2, prior variance
                inverse \alpha
   output: Learnt parameter w
1 initialization;
2 repeat
         for ith column in D do
               Set \hat{\mathbf{D}} that includes all columns in \mathbf{D} except the current ith column;
              Perform ridge regression \min_{\mathbf{w}}(\frac{1}{2}||\boldsymbol{\beta}(\mathbf{t} - \mathbf{D}\mathbf{w})||_2^2 + \frac{1}{2}||\boldsymbol{\alpha}\mathbf{w}||_2^2) to get \mathbf{w}, using \boldsymbol{\beta} = \boldsymbol{\sigma}_{\partial t}^{-2} as the inverse
                 noise variance and the prescribed \alpha as inverse prior variance;
               while changes in w is greater than a threshold do
                     Update \beta using (9), with w from the previous iteration;
                    Perform ridge regression \min_{\mathbf{w}}(\frac{1}{2}||\boldsymbol{\beta}(\mathbf{t} - \mathbf{D}\mathbf{w})||_2^2 + \frac{1}{2}||\boldsymbol{\alpha}\mathbf{w}||_2^2) to update \mathbf{w}, using the updated \boldsymbol{\beta} as the
                       inverse noise variance and the prescribed \alpha as inverse prior variance;
               end
               Calculate and store the evidence \mathcal{J}(\mathbf{t}|\tilde{\mathbf{D}}, \boldsymbol{\beta}, \boldsymbol{\alpha}) using (14);
```

Comparing derivative approximation using weak formulation and finite difference

until further removal of columns does not increase evidence;

Select the column that yields the greatest evidence after removal, and remove this column in **D**;

One of the major challenges in applying SINDy-type algorithms to learn governing equations from data is approximating temporal (for learning ODEs) and spatial (for learning PDEs) derivatives using discrete data in the presence of noise. Conventional finite differences can greatly amplify noise, rendering the dynamics unrecoverable for the STLS algorithm. To overcome this problem, one can employ the weak formulation [13, 14]. This formulation uses Galerkin projection and a compact and smooth test function $\phi(t)$ to "transfer" the derivatives on the data to the test function. For example, the Galerkin projection allows the time derivative on the data $\hat{x}^{(k)}(t)$ to be expressed as

$$\int_{a}^{b} \hat{x}^{(k)}(t)\phi(t)dt = (-1)^{k} \int_{a}^{b} \hat{x}(t)\phi^{(k)}(t)dt,$$

provided that $\phi(t)$ and its derivatives up to $\phi^{(k)}(t)$ have compact support in $t \in [a, b]$. As a result, any noise present in the data is not magnified, leading to a substantial enhancement in the performance of the subsequent regression.

In practice, when applying the method to a discrete dataset X, both finite difference and weak formulation can be written as linear operations on X, i.e. as matrices multiplied to X, resulting in the formulation in (5). In fact, both methods result in banded matrices

$$L_{\partial t} = \begin{bmatrix} a_{-n} & a_{-n+1} & \dots & a_{n-1} & a_n & 0 & \dots & 0 \\ 0 & a_{-n} & a_{-n+1} & \dots & a_{n-1} & a_n & \dots & 0 \\ \vdots & \ddots & \ddots & \ddots & \ddots & \ddots & \vdots \\ 0 & \dots & 0 & a_{-n} & a_{-n+1} & \dots & a_{n-1} & a_n \end{bmatrix}$$
(19)

and

3

4

5

8

9

10 11

12

$$L_{I} = \begin{bmatrix} b_{-n} & b_{-n+1} & \dots & b_{n-1} & b_{n} & 0 & \dots & 0 \\ 0 & b_{-n} & b_{-n+1} & \dots & b_{n-1} & b_{n} & \dots & 0 \\ \vdots & \ddots & \ddots & \ddots & \ddots & \ddots & \vdots \\ 0 & \dots & 0 & b_{-n} & b_{-n+1} & \dots & b_{n-1} & b_{n} \end{bmatrix}.$$

$$(20)$$

For an equispaced central difference method of order 2n, $b_0=1$, and $b_i=0$, $\forall i\neq 0$, and $a_i=\hat{a}_i/\Delta t$, where Δt is the size of the timestep in the regularly sampled time-series data, and \hat{a}_i the coefficients for the central difference. For example, when n=2, $[\hat{a}_{-2} \dots \hat{a}_{2}] = [1/12, -2/3, 0, 2/3, -1/12]$. Meanwhile, the weak formulation gives

$$a_n = w_n \dot{\phi}_n$$
 and $b_n = w_n \phi_n$

where $w_{-n}...w_n$ gives the quadrature for the integration and ϕ_i and $\dot{\phi}_i$ are the test function and its time derivative at t = (a+b)/2 + 2i(b-a)/n.

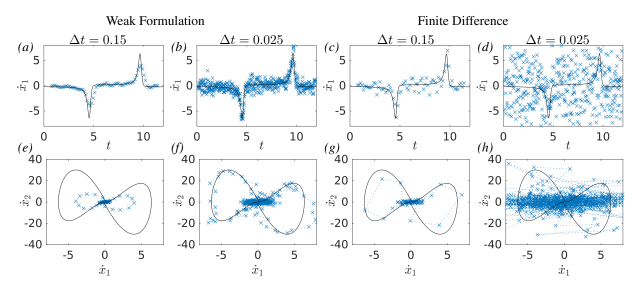


Figure 2: Comparison of time derivatives in the Van Der Pol system recovered from noisy data (blue crosses) against the true system (black line), using (a,b,e,f) weak formulation with 9 points and test function $\phi=(t^2-1)^2$ and (c,d,g,h) 8th order (9 points) central difference: (a-d) the time derivative of x_1 over time t; (e-h) the phase diagram on the $\dot{x}_1-\dot{x}_2$ plane. While the noise $\sigma_x=0.1$ and the number of points used to calculate each derivative n+1=9 are fixed, the sampling frequency of the dataset varies between $\Delta t=0.1$ in (a,c,e,g) and $\Delta t=0.025$ in (b,d,f,h). Hence, the time window over which the weak formulation is applied varies proportionally to Δt . Note that in (a), the weak formulation causes significant artificial smoothing due to a large time window, leading to an inaccurate approximation of the phase diagram in (e).

Note that both $L_{\partial t}$ and L_I are rectangular matrices with dimension $\tilde{N} \times N$, where \tilde{N} is the number of measurements in the original dataset, whereas $N \leq \tilde{N}$ is the number of data after computing the derivatives. $\tilde{N} - N = 2n$ number of data points are usually filtered away near the two ends of the data series due to a lack of data beyond the measured point to compute the corresponding derivative. Also, notice that the scaling of $|L_{\partial t}|:|L_I|$ is of order $\mathcal{O}(1/\Delta t)$ in finite difference, whereas that of weak formulation is of order $\mathcal{O}(1)$. This is why, when applied to (5), finite difference amplifies noise, while weak formulation does not.

However, from a signal processing standpoint, the weak formulation can be described as a convolution operator that performs low-pass filtering [13]. This filtering process can remove high-frequency signals and perhaps result in misidentification of the model. This is best demonstrated by the phase diagram of the Van der Pol system (figure 2), where the dynamic features a sudden and rapid change. When the sampling rate is high, the weak formulation effectively captures time derivatives without significantly amplifying noise (figure 2(b, f)) compared to the finite difference method (figure 2(d, h)). However, at a low sampling rate where the integral window is unavoidably large, the weak formulation's low-pass filtering characteristic leads to noticeable artificial smoothing in the data (figure 2(a, e)). By comparison, the finite difference method remains reasonably unbiased despite the significant amplification of the noise variance (figure 2(c, q)).

Therefore, the decision between weak formulation and finite difference involves a compromise between attenuating noise and maintaining signal. The finite difference method does not attenuate any signal from a spectrum standpoint, but it amplifies noise by the magnitude of the sampling rate. Hence, it is more suitable for data sampled at a low sampling rate. Conversely, the weak formulation is a proficient method for attenuating noise and mitigating noise amplification, but it may also suppress signals at high frequencies. It is suitable for data that is sampled at a rate significantly higher than the Nyquist frequency of the signal. In practice, the sampling rate is often limited by the capabilities of the measuring hardware, and problems that push the limit of the equipment are often sampled at a frequency just above the Nyquist frequency of the relevant dynamics. As a rule of thumb, weak formulation usually requires at least 6-12 data points in the integral. Hence, practically, it can only resolve dynamics 6-12 times slower than the sampling frequency. By improving model selection performance in the high noise regime, the Bayesian reformulation presented in this work improves the overall chances of recovering the correct model even when the sampling rate is close to the Nyquist frequency of the dynamic, i.e. in the low-data regime.

3 Examples

This section will showcase the noise robustness and data efficiency of our Bayesian-SINDy algorithm in recovering the desired equations precisely in comparison with the original SINDy algorithm [1] that is based on Sequentially Thresholded Least Squares (STLS). We will also compare our Bayesian-SINDy algorithm with SparseBayes [24] to show the improvements our Bayesian-SINDy has brought to the Bayesian approaches. Four examples will be used to compare the three algorithms, in which we will consider a polynomial library $\Theta(\mathbf{X})$ up to third order. To compare the three algorithms and to ensure the result is not skewed by individual runs, we will vary the sampling frequency and noise variance for the first three synthetic examples, and iterate the learning process 10000 times to compare the percentage of runs that each algorithm has successfully recovered exactly all the terms in original equation without extra spurious term, i.e. success rate of recovering the original equations. In this work, the use of success rate as the metric is justified as we are focused on the most parsimonious and generalisable equations, which are the original equations, precisely. Furthermore, to avoid miscounting models with the same sparsity pattern but substantially different coefficients, the success rate is only counted when the model coefficient error (MCE = $||\mathbf{w} - \hat{\mathbf{w}}||_2/||\hat{\mathbf{w}}||_2$, where $\hat{\mathbf{w}}$ is the truth) is below 25%. As for the fourth example, we will learn the dynamical equations from real-world measurements. Although a ground-truth model is absent, historical uses of the dataset [34, 35, 31] have shown that we should be expecting the Lotka-Volterra equations.

We should note that in this section, we are not aiming to claim the Bayesian-SINDy to be the most effective method in recovering the correct dynamical equations from data, nor is SINDy (STLS) the most effective methodology among many SINDy extensions. As we shall discuss later in §4.2, we expect more computationally-intensive sampling-based methods to be more robust. However, by comparing SINDy (STLS) with an equivalently simple and computationally efficient method derived from the Bayesian framework, we will demonstrate that a Bayesian approach is more robust in noisy and low data scenario, and thereby encourage a Bayesian perspective to other extensions of SINDy. The comparison with SparseBayes, meanwhile, demonstrates the improvement we have brought to our learning algorithm. To compare the algorithms properly, we have also taken steps to ensure they are performing the best they can with the appropriate hyperparameters. Details of the hyperparameters used for each algorithm and in each example are listed in the appendix.

3.1 Van der Pol oscillator

In this example, we consider the Van der Pol oscillator system given by

$$\dot{x}_1 = x_2, \quad \dot{x}_2 = bx_2(1 - x_1^2) - x_1,$$
 (21)

in which we use the values b=4 and the initial conditions $\mathbf{x}=[2,0]$. A Gaussian measurement noise with distribution $\mathcal{N}(0,0.1^2)$ or $\mathcal{N}(0,0.2^2)$ is added to the data, sampled over the interval $t\in[0,12]$. Figure 3 shows a typical example of the data provided to the algorithms and the estimation by SINDy (STLS), Bayesian-SINDy and SparseBayes, while figure 4 shows the success rates of the each algorithm using both an 8th-order (9 points) central difference scheme and a weak formulation scheme with 9 points and $\phi=(t^2-1)^2$.

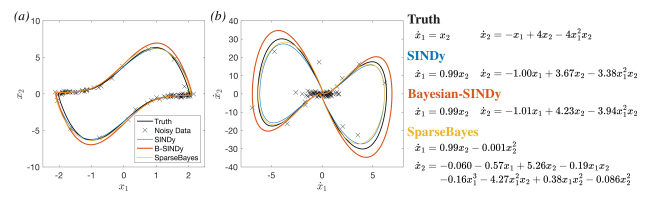


Figure 3: The Van der Pol system (21) with $\beta=4$ (black line) and the system estimated by SINDy (STLS, blue), Bayesian-SINDy (red) and SparseBayes (RVM, yellow) given the noisy data of the system (black crosses). The right panel shows the equations learnt, while (a) shows the phase diagram in x_1-x_2 space, and (b) shows the equivalent in $\dot{x}_1-\dot{x}_2$ space. Finite difference is used to obtain $\dot{\mathbf{x}}$. The noise level is $\sigma_x=0.1$ and the data are sampled at $\Delta t=0.15$ for $t\in[0,12]$.

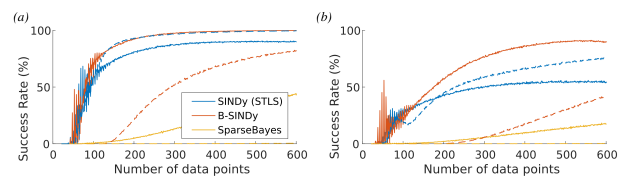


Figure 4: The percentage of runs that successfully identifies the Van der Pol oscillator equations from data using SINDy (STLS, blue), Bayesian-SINDy (red) and SparseBayes (yellow) after 10000 iterations against the number of data points $\tilde{N}=12/\Delta t$. Solid lines show results from finite differences and dashed lines show results from weak formulations. The noise levels are (a) $\sigma_x=0.1$ and (b) $\sigma_x=0.2$. Note that the oscillation at low \tilde{N} is an artefact of stroboscopic effect.

At both noise levels, Bayesian-SINDy shows robust performances in recovering the Van der Pol equations using finite difference despite the noise amplification (c.f. figure 2h). Although the weak formulation helps SINDy recover the equations when noise is low, its success rate remains not as high as Bayesian-SINDy when noise is high. Furthermore, in the high-noise case, the weak formulation causes SINDy's performance to drop below that of finite differences. This is due to the low-pass filtering property, as anticipated by our discussion in §2.3 and illustrated in figure 2. Meanwhile, SparseBayes performs consistently worse than the other two algorithms. The poor performance of SparseBayes is due to the lack of facilities to account for noise variation over time, which we will further explain in §3.2.

One may notice that the success rate is highly oscillatory when \tilde{N} is low. This is due to a stroboscopic effect at low sampling rates. In a stiff system such as the Van der Pol oscillator, the majority of the information about the system is concentrated within the few data points at which the dynamics spike (i.e. when $\dot{\mathbf{x}}$ is far from $\mathbf{0}$ in figures 2(e-h) and 3(b)). The rest of the data points simply repeat information about the plateau region, where $\dot{\mathbf{x}}$ is close to zero. With only two spikes within the time interval, successful recoveries of the equations are highly sensitive to whether the spikes are sampled sufficiently in the data. With a continuously changing sampling rate and fixed time interval, the stroboscopic effect implies that the spike might not be sampled enough at some sampling rates, leading to oscillation in the resulting success rate. Later in §4.3, we will show how to drive the algorithms to preferentially learn from more informative data points and improve data efficiency.

3.2 Cubic oscillator

In this example, we consider the damped nonlinear oscillator given by

$$\dot{x}_1 = ax_1^3 + bx_2^3, \qquad \dot{x}_2 = cx_1^3 + dx_2^3,$$
 (22)

in which we use the values a=-0.1, b=-2, c=2, d=-0.1, and the initial conditions $\mathbf{x}=[1,0]$. Data are sampled over a fixed time interval $t\in[0,5]$ at various rates, and a Gaussian measurement noise with distribution $\mathcal{N}(0,0.005)$ or $\mathcal{N}(0,0.01)$ is added to the data. To recover the time derivatives, here we use the weak formulation with 7 data points, with a test function $\phi=(t^2-1)^4$.

Figure 5 depicts a typical instance of the data provided to the algorithms and the resulting learning outcome of Bayesian-SINDy, and figure 6 summarises the success rate of each algorithm in recovering (22). Here, Bayesian-SINDy has shown better performance in all scenarios examined. SparseBayes is also resilient to noise, but it is less data-efficient than Bayesian-SINDy. In contrast, SINDy (STLS) exhibits a decline in performance when the noise variance increases.

One reason Bayesian-SINDy performs better than SparseBayes is that it properly accounts for the noise variance according to (9). As shown in figure 7, the noise variance varies significantly over time due to the contribution from the cubic terms. However, SparseBayes presumes a constant (inverse) noise variance β over time and hence cannot account for the time-varying contribution from the cubic terms. In contrast, the noise iteration procedure in Bayesian-SINDy captures the contribution of σ_{Θ}^2 to σ^2 well, thereby improving Bayesian-SINDy's performance in the low-data limit.

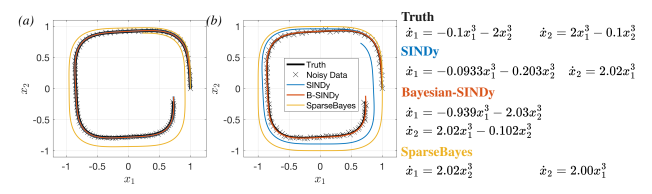


Figure 5: The phase diagram of the cubic nonlinear oscillator system (22, black line) and the system estimated by SINDy (STLS) (blue), Bayesian-SINDy (red) and SparseBayes/RVM (yellow) given the noisy data (crosses) of the system. The noise level are (a) $\sigma_x=0.005$ and (b) $\sigma_x=0.01$ and the data are sampled at $\Delta t=0.05$ for $t\in[0,5]$. The right panel shows the model equations from each algorithm when $\sigma_x=0.01$. In (a), the prediction by SINDy and B-SINDy is so close to truth their differences are practically indiscernible.

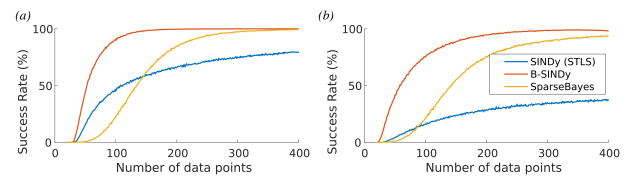


Figure 6: The percentage of runs that successfully identifies the cubic nonlinear oscillator equations from data using SINDy (STLS, blue), Bayesian-SINDy (red) and SparseBayes (yellow) after 10000 iterations against the number of data points $\tilde{N} = 5/\Delta t$. The noise levels are $(a) \sigma = 0.005$ and $(b) \sigma = 0.01$.

3.3 Lorenz system

In this example, we consider the Lorenz system given by

$$\dot{x}_1 = s(x_2 - x_1), \qquad \dot{x}_2 = rx_1 - x_2 - x_1x_3, \qquad \dot{x}_3 = x_1x_2 - bx_3,$$
 (23)

in which we use the values s=10, b=8/3, r=28, and the initial conditions $\mathbf{x}=[-1,6,15]$. Data are sampled over the time interval $t\in[0,2.5]$, and a Gaussian measurement noise with distribution $\mathcal{N}(0,\sigma_x^2)$ is added to the data,

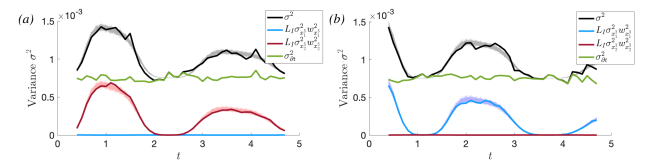


Figure 7: Cubic oscillator: breakdown of contributions to the total variance of noise σ in the dynamics of (a) x_1 and (b) x_2 by each term in (9). Thick lines show the variance computed from the statistics of 1000 iterations, and lighter lines (shaded area) show the bound of the theoretical calculation using (9) from each iteration.

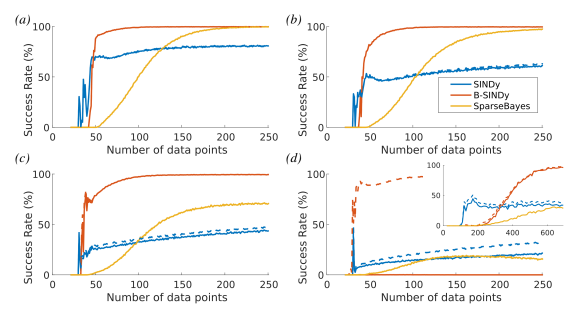


Figure 8: The percentage of runs that successfully identifies the Lorenz system equations from data sampled between $t \in [0, 2.5]$ using SINDy (STLS, blue), Bayesian-SINDy (red) and SparseBayes (yellow) after 10000 iterations against the number of data points $\tilde{N} = 2.5/\Delta t$. The noise levels are (a) $\sigma = 0.025$, (b) $\sigma = 0.05$, (c) $\sigma = 0.1$ and (d) $\sigma = 0.2$. The dashed line shows the success rate of recovering the model without the x_2 term in the equation for \dot{x}_2 . The inset in (d) shows the success rate when the time interval is extended to $t \in [0, 10]$.

with σ_x varying between 0.025 to 0.2. Here, we use a central finite difference with 13 data points to recover the time derivatives.

Figure 8 shows the success rates of the three algorithms. When noise is low, SINDy shows moderate success in recovering the dynamical equations, but performance degrades quickly as noise increases. Meanwhile, both Bayesian-SINDy and SparseBayes show more robustness in learning the correct model as noise increases, and Bayesian-SINDy maintains better performance at the low-data limit.

However, in figure 8(d), in which the noise is $\sigma=0.2$, none of the three algorithms can recover the model exactly. Both SINDy (blue) and SparseBayes (yellow) can only recover the full model sporadically and showed little consistency in the model they produced. In contrast, Bayesian-SINDy consistently recovers a model with only the x_2 term in equation for \dot{x}_2 missing (as shown by the dashed red line). The missing x_2 term in \dot{x}_2 is particularly difficult to learn because it is highly collinear with the existing x_1 term in the model. To distinguish the two would require more data in regions in which the two differ significantly, but, as shown in figure 9, the given signal is highly obscured by noise and does not sample those regions sufficiently in the short time interval $t \in [0, 2.5]$. In other words, there is simply not enough information in the small time window to infer the x_2 term in the equation for \dot{x}_2 . From the evidence-maximisation perspective, this difficulty is reflected in figure 1, where $\mathcal{J}(\alpha_m) > 0$ only for a small range of α_m . In practice, when the data is too low or too noisy, the information content in the data may not be enough to infer the full dynamics. In such cases, one inevitably has to increase the amount of data available by considering a long time interval, say, $t \in [0, 10]$. As demonstrated in the inset of figure 8(d), Bayesian-SINDy can eventually recover the full dynamics most of the time, while SINDy remains less robust in this high noise regime. This lack of information, as depicted here in figure 9, also hints at the possibility of further improving data efficiency if we are allowed to sample data in the most informative state. We will discuss this information-theoretical approach further in §4.3.

3.4 Lotka-Volterra equation from Lynx-Hare population data

In this example, we apply the three algorithms to model the population dynamics of Lynx and Hare in Canada, using pelt trade data collected by the Hudson Bay Company between 1900 and 1920 [34, 35]. Assuming the pelts of these two species traded by the Company are proportional to the true population, it may be anticipated that these two populations will approximately adhere to the Lotka-Volterra model for predator-prey dynamics. In fact, this dataset has been used as empirical evidence to support the Lotka-Volterra equations, and has become increasingly popular as a real-life benchmarking dataset among statisticians and data scientists [36, 12, 31]. The primary challenge posed by this dataset

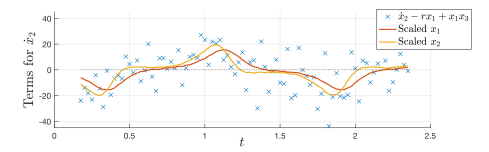


Figure 9: Learning the dynamics of $\dot{x}_2 - rx_1 + x_1x_2$ in the Lorenz system: the crosses show the values of $\dot{x}_2 - rx_1 + x_1x_3$ from the data, which should match x_2 (red line). However, x_1 (yellow line) is highly collinear with x_2 , and it is not obvious which term should fit the data better. Both x_1 and x_2 are scaled to best match the data $\dot{x}_2 - rx_1 + x_1x_3$. Here, $\Delta t = 0.025$ ($\tilde{N} = 100$) and $\sigma = 0.2$.

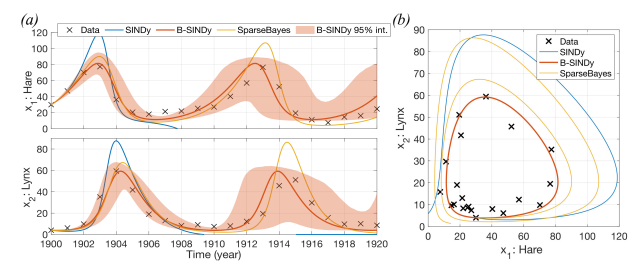


Figure 10: SINDy (STLS, blue), Bayesian-SINDy (red) and SparseBayes (yellow) applied to the lynx-hare population data based on the pelts trade by the Hudson Bay Company between 1900 and 1920 [34, 35]. (a) Hare (upper) and Lynx (lower) populations over time. Solid lines show the model reconstruction from the parameters estimated by each algorithm, and the shaded area depict the 95% confidence interval given by the ensemble reconstruction using the parameter uncertainty estimated by Bayesian-SINDy. (b) Phase diagram of the Lynx and Hare populations.

is its limited number of data points - specifically, 21 annual data points for each population, resulting in a total of 42 data points. Due to the low sampling rate, the finite difference method is more appropriate, but it will, as one expects, also amplify the noise in the derivative, which we will then mitigate by relying on the noise robustness of the Bayesian framework. The weak formulation was also tried, but the low-pass filtering effect filtered vital signals in the data, leading to subpar learning performance across all methods.

The data is shown in figure 10, where x_1 corresponds to Hare populations, and x_2 corresponds to Lynx populations. Given the data is not a direct measurement of the population, it is expected to be highly uncertain. However, the level of uncertainty is not quantified. The lack of uncertainty quantification is not a major challenge for SINDy or SparseBayes, as SINDy does not consider this factor, while SparseBayes can optimise the noise variance through evidence maximisation. Our Bayesian-SINDy algorithm, on the other hand, requires a value for σ_x .

To address the difficulty, we ran a hyperparameter sweep on σ_x , assuming $\alpha=10^{-2}$, and chose the value of $\sigma_x\approx 2.7$ by evidence maximisation. Although this technique is similar to the evidence maximisation process in SparseBayes, a notable distinction is that our procedure does not assume a uniform σ for all data points. Instead, we calculate it using equation (9) and algorithm 1. The parameters estimated are displayed in table 1, where the Lotka-Volterra model is successfully recovered by Bayesian-SINDy, while the other two struggle to learn a consistent model. Uncertainties in the estimated parameters, also shown in table 1, are also easily recovered from the the posterior distribution. Figure 10 also shows how the uncertainty in the estimated parameter propagates through the dynamics, assuming the same initial

Table 1: Coefficients and their uncertainty (one standard deviation) estimated from the lynx-hare data using SINDy (STLS), Bayesian-SINDy and SparseBayes. The library consists of all combinations of x_1 and x_2 up to third order, but the table only shows terms that have non-zero coefficients from one of the algorithms. The non-zero terms in the Lotka-Volterra model are highlighted in yellow.

	$\dot{x}_1 : 1$	$\dot{x}_1:x_1$	$\dot{x}_1:x_2$	$\dot{x}_1:x_1x_2$
SINDy (STLS)	-9.92	0.82	0.31	-0.036
Bayesian-SINDy	0	0.53 ± 0.0043	0	$-0.026 \pm 6.1 \times 10^{-6}$
SparseBayes	0	0.56 ± 0.0045	0	$-0.027 \pm 5.3 \times 10^{-6}$
	$\dot{x}_2 : 1$	$\dot{x}_{2}:x_{2}$	$\dot{x}_2: x_1x_2$	$\dot{x}_2: x_2^3$
	_	2 2	** Z * ** I ** Z	** <u>Z</u> · ** <u>Z</u>
SINDy (STLS)	-1.26	-0.92	0.027	0
SINDy (STLS) Bayesian-SINDy	-1.26 0			$ \begin{array}{c} 0 \\ 0 \\ 0 \\ 3.6 \times 10^{-5} \pm 1.7 \times 10^{-9} \end{array} $

condition. Notably, our method was able to estimate the parameter uncertainties without expensive sampling, unlike previous techniques [31].

4 Discussion

4.1 Scaling of likelihood and Occam factor with N and β

To better understand why the Bayesian-SINDy, and Bayesian approaches in general, are more robust than frequentist approaches such as STLS at selecting the correct model in the noisy and scarce data limit, we compare the two approaches under a unifying Bayesian framework. From the Bayesian viewpoint, frequentist approaches seek to maximise the log-likelihood, which, under the premise of Gaussian likelihood, can be expressed as

$$\mathcal{J}(\mathbf{t}|\mathbf{D}, \mathbf{w}, \boldsymbol{\beta}) = -\frac{N}{2} \ln 2\pi - \frac{1}{2} \boldsymbol{\beta} \cdot \boldsymbol{\beta} - \frac{1}{2} \boldsymbol{\beta} \cdot (\mathbf{t} - \mathbf{D}\mathbf{w})^{2}.$$
 (24)

Meanwhile, our Bayesian approach aims to maximise the log-evidence

$$\mathcal{J}(\mathbf{t}|\mathbf{D},\boldsymbol{\beta},\boldsymbol{\alpha}) = \mathcal{J}(\mathbf{t}|\mathbf{D},\mathbf{w},\boldsymbol{\beta})|_{\mathbf{w}=\boldsymbol{\mu}} + \mathcal{J}(\mathbf{w}|\boldsymbol{\alpha})|_{\mathbf{w}=\boldsymbol{\mu}} + \frac{1}{2}\ln\left(\det(\boldsymbol{\Sigma})\right), \tag{25a}$$

which is composed of three components: the determinant of the covariance matrix

$$\Sigma = (\mathbf{A} + \mathbf{D}^T \mathbf{B} \mathbf{D})^{-1} \in \mathbb{R}^{M \times M}, \tag{25b}$$

the log-likelihood $\mathcal{J}(\mathbf{t}|\mathbf{D},\mathbf{w},\beta)$ and the log-prior $\mathcal{J}(\mathbf{t}|\mathbf{D},\mathbf{w},\beta)$ at the maximum a posteriori (MAP) point. The MAP point refers to the parameter values $\mathbf{w}=\mu$ that maximise the posterior. It should be noted that in equation (24), the log-likelihood scales linearly with the magnitude of the inverse noise variance β and the number of data points N. On the other hand, the remaining components in the log-evidence, also known as the Occam factor, serve as a penalty for models that have an excessive number of parameters. The Occam factor scales with the number of terms M in the model and only has a partial and weak $\ln N$ scaling with N.

Figure 11 demonstrates how the scaling of the log-likelihood and the Occam factor plays out in the model selection process. The left column of figure 11 shows the contribution of the log-likelihood (blue) and the Occam factor's (red) to the log-evidence across all 1024 possible models for \dot{x}_3 in the Lorenz system when a smaller, second-order polynomial library containing ten terms is considered. For each model, the barcode plot below show the library terms that are non-zero, with the non-zero terms x_3 and x_1x_2 in the true model are coloured green. The models are sorted by log-evidence, demonstrating a Bayesian perspective on model selection, where the model with maximum evidence is selected as the most parsimonious model for the data. For the same dataset, the right column of figure 11 sort the model based on log-likelihood calculated from the maximum likelihood estimate (MLE), i.e. $\mathbf{w} = \mu_{\text{MLE}}$, which can be obtained by ordinary least squares. (Note that when N and β is large, the MLE and MAP estimates of μ converge.) In contrast to the Bayesian perspective on the left, the right column reflects a frequentist perspective where the model with the best fit is selected as the best model for the data. As one would expect, the MLE-based model selection always select the model with all ten terms. To induce sparsity, the frequentist approach STLS introduces a empirical thresholding process. The idea is to remove small coefficient terms that doesn't contribute much to the likelihood, thereby achieving a compromise between maximising likelihood and sparisty.

When the data is abundant and the noise is minimal (i.e. $N||\beta|| \gg M$), models that include the correct terms have significantly higher likelihood and are easily distinguishable from the other models when sorted by likelihood. This is

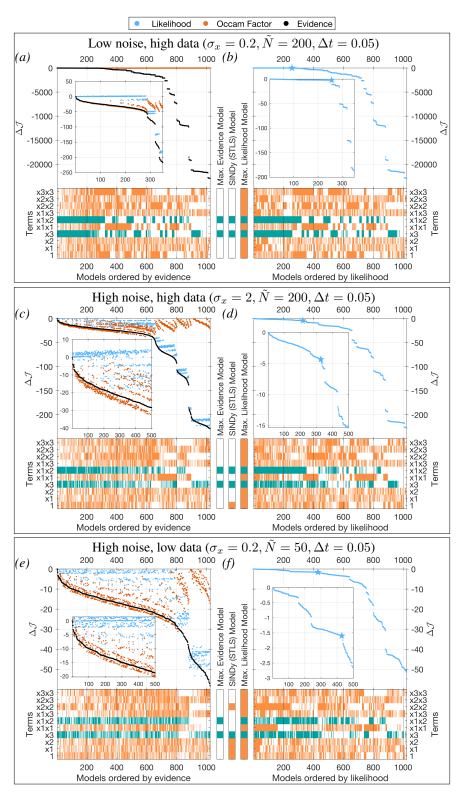


Figure 11: Difference in log-evidence, log-likelihood and the Occam factor (i.e. log-evidence minus log-likelihood) compared to the highest-ranked model of all possible models for \dot{x}_3 in the Lorenz system. Models are ranked by (a,c,e) evidence and (b,d,f) likelihood. Insets show a zoomed plot of the first few hundred models, and the barcode plots below show the non-zero terms in each model, marked as green (correct) or orange (incorrect). The correct model is always ranked highest by evidence in (a,e,i), and indicated with a star in (b,d,f). Meanwhile, the model selected by highest likelihood contain all the terms, and the models selected by SINDy are shown in the middle. Here, (a-b) $\sigma_x=0.2$ and $\tilde{N}=200$, (c-d) $\sigma_x=1$ and $\tilde{N}=200$, and (e-f) $\sigma_x=1$ and $\tilde{N}=50$.

demonstrated by the continuous green band in the barcode plot in 11(b). Phenomenologically, this signifies that all models containing the appropriate terms exhibit comparable levels of likelihood. STLS leverages this clear separation in likelihood and only eliminate irrelevant terms that do not affect the likelihood much in its thresholding process, thereby effectively constraining its search for the most sparse model within this region of large likelihood.

However, as the amount of data is lowered (i.e. smaller N), or when the noise increases (i.e. smaller $||\beta||$), the distinct separation of the high likelihood region from the other possible models becomes less evident. In these scenarios, models lacking the relevant terms are more likely to fit the data incidentally, as reflected by the discontinuous green band in barcode plot of figure 11(d,f). STLS may, thus, converge on an incorrect model. Alternatively, STLS may not converge to the most sparse model since there is a non-negligible chance for irrelevant terms also to have coefficients larger than the threshold when N or β decreases.

Conversely, figure 11(a, c, e) demonstrates that the Occam factor assumes greater significance in model selection when noise is large and data is scarce. The Occam factor serves two crucial functions in model selection. First, it penalises models with too many terms, as it scales with M. Second, it quantifies the variance of the estimated parameter, and discards terms that have small variances in their coefficients, as this indicates that the corresponding model is not robust (or too sensitive to small changes in the parameter values). In performing the above two functions, the rigorously derived Occam factor excludes terms that incidentally fit the data, ensuring the robustness of the estimated model.

4.2 Limitation and use cases of Bayesian-SINDy

Although Bayesian-SINDy has shown greater resilience than the original STLS-based SINDy in accurately identifying the proper dynamical equations from noisy and scarce datasets, there is still much room for improvement.

For example, one might find the noise level in the nonlinear (cubic) oscillator example in §3.2 low compared to other examples. In fact, none of the three algorithms were able to recover the correct equations upon further increases in noise level. This is because, in the presence of highly nonlinear cubic terms, the noise distribution can become highly skewed and deviate from a Gaussian distribution upon an increasing noise-to-signal ratio. As we have stressed in (9) and in §2.2, the Gaussian distribution is only strictly valid if the noise is truly Gaussian and the model consists of linear terms only. However, it serves as a good approximation in most cases, as shown empirically by the other examples. Alternatively, when the noise distributions are no longer tractable or well approximated by analytical methods, one can resort to sampling-based techniques to better capture the accurate noise distribution. For example, recent work by Hirsh et al. [31] has demonstrated the possibility of further improving noise robustness and data efficiency by accurately capturing distributions through sampling techniques such as MCMC. However, this improvement comes at the expense of costly sampling. For example, it took days of computation to recover the Lotka-Volterra equation from the Hudson Bay Lynx-Hare population dataset using the MCMC-based method in [31], compared to the sub-second computation by Bayesian-SINDy.

Consequently, the choice of which approach to adopt depends on the specific application and circumstances. If computing time and cost are not significant constraints, sampling-based methods such as those suggested by [30] or [31] should give more accurate models than Bayesian-SINDy. Nonetheless, Bayesian-SINDy is a valuable tool for conducting rapid preliminary investigations. In situations that demand rapid learning, such as system identification in control, using a sampling-based method would not be practical. Instead, Bayesian-SINDy could be a cost-effective approach to extracting symbolic models from noisy and limited data in a fast and reliable manner.

4.3 Application of Bayesian-SINDy in active learning

One way to further improve data efficiency in the Bayesian framework is to optimise experimental design. Lindley [37] established the approach to optimal design by choosing, from all possible data points (labelled 'experiments' in [37]), the one that maximises the information content. This information content is quantified by a utility function akin to Shannon's information entropy, which is naturally defined by the probabilistic description in the Bayesian framework. However, evaluating the information entropy usually involves integration in high-dimensional spaces, which can be expensive in a sampling-based technique. For example, in a parameter inference problem, the information entropy is defined by the parameter posterior. Evaluating this information entropy would require integration over the entire parameter space, the cost of which scales exponentially with the number of parameters.

This computational complexity is further exacerbated if an information-based approach analogous to Lindley's is applied to the model selection problem. The information entropy H of interest here is

$$H = -\int p(\mathbf{t}|\mathbf{D}, \boldsymbol{\beta}, \boldsymbol{\alpha}) \ln (p(\mathbf{t}|\mathbf{D}, \boldsymbol{\beta}, \boldsymbol{\alpha})) d\mathbf{t},$$
 (26)

which is based on the evidence $p(\mathbf{t}|\mathbf{D}, \boldsymbol{\beta}, \boldsymbol{\alpha})$. In a sampling-based technique, the evaluation of $p(\mathbf{t}|\mathbf{D}, \boldsymbol{\beta}, \boldsymbol{\alpha})$ already requires the integration of the posterior in the parameter space. The evaluation of H, then, would have required further integration in the \mathbf{t} -space, which would be prohibitively expensive if a sampling technique were used. Fortunately, the expensive evaluation can be avoided if we accept the Gaussian approximation used throughout this work. Then, the analytical tractability of the Gaussian distribution allows for H to be rewritten explicitly as

$$H = \frac{N}{2}(1 + \ln(2\pi)) + \frac{1}{2}\ln|\mathbf{C}|,\tag{27}$$

after substituting (14-15) into (26). Here, C is the covariance matrix of the evidence.

However, in contrast to Lindley's approach, which seeks to maximise information content (analogous to the negative of H), here we seek to maximise the information entropy H over all the possible data we can sample. This might seem counter-intuitive, but unlike the parameter inference problem, in which one seeks to minimise uncertainty in the parameters, in our model selection problem, one seeks to minimise certainty of the current model so as to encourage the exploration for a better model. This exploration maximisation through entropy is analogous to entropy maximisation in reinforcement learning.

Optimising data efficiency has, therefore, become an exercise of sampling data that maximises the covariance C. To illustrate the potential improvement this may bring, here we consider again the Van der Pol oscillator and the Lorenz system as examples. Instead of using the entire dataset, here, each new data point is selected by maximising gain in H given the estimated model from the existing dataset. Then the newly selected point is assimilated, and the model is updated with the new dataset. This process is repeated, and the number of data points required to recover the full model is recorded.

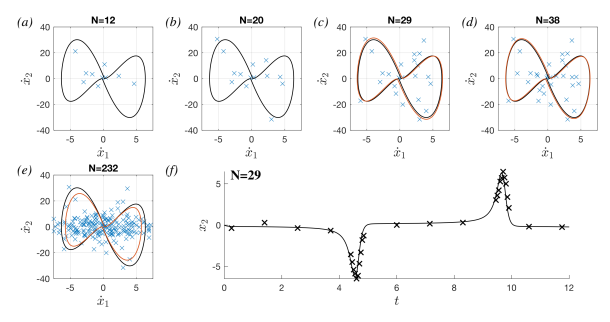


Figure 12: Recovery of the Van der Pol model by sequential data assimilation, where each new data point is selected by maximising information gain. Blue crosses in (a-e) show the data sampled in the phase diagram of $\dot{\mathbf{x}}$, and black lines show the original dynamic without noise. At N=29 (panel (d)), the full model is recovered, and the red dashed line shows the prediction by the recovered model. Panel (e) shows the full dataset with N=232 points. Panel (f) shows the data sampled on x_2-t space when N=29, which are concentrated at the region when the dynamic spikes.

In the Van der Pol example, we have briefly mentioned that the majority of the system's information is concentrated during the short time when the dynamics $\dot{\mathbf{x}}$ is far from the origin (see §3.1). Here, figure 12 shows how effective the information gain H is in guiding the learning algorithm to concentrate sampling in the more informative regions, thereby recovering the full system with less data. The full system is recovered with only N=29 data points, an order of magnitude lower than the full dataset. Notice that most data are sampled around the spikes in figure 12(f), showing that most information about the system is indeed concentrated in those regions.

As for the Lorenz system, §3.3 has shown that the x_2 term in \dot{x}_2 is particularly difficult to learn due to high noise that obscurs the dynamics and a lack of information to distinguish x_2 from x_1 . Here, figure 13 shows how H preferentially

samples regions with more information to distinguish between the two terms, thereby significantly reducing the overall data required to recover the full model.

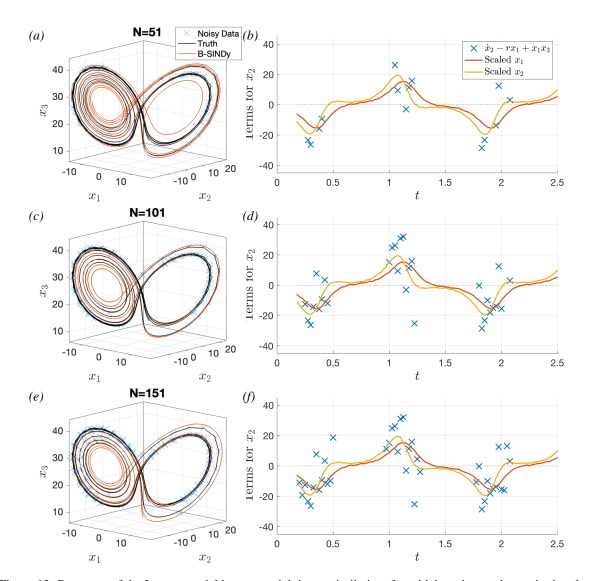


Figure 13: Recovery of the Lorenz model by sequential data assimilation, for which each new data point is selected by maximising information gain. In (a,c,e), blue crosses show the data sampled, and black lines and red dashed lines show the original dynamic without noise and the model prediction, respectively. Similar to figure 9, panels (b,d,f) show the residue signal from the dynamics of \dot{x}_2 , which should be fitted by x_2 . Here, blue crosses show the data sampled, which is concentrated at the time where x_2 is the most distinguishable from x_1 . At N=51 and N=101 (panels (a-d)), the Lorenz model is partially recovered, with the exception of the x_2 term in \dot{x}_2 . At N=151 (panels (e-f)), the full model is recovered.

In real-life applications such as system identification for control, one can imagine using H as part of an objective function for optimal control. By balancing the objective of learning more about the dynamics governing the system (i.e. exploration) with the objective of achieving certain target state(s) (i.e. exploitation), one can develop the current framework into a fully automated active learner that can learn about the system and drive it to a certain state in a very rapid manner.

We also note that, as well as our current definition of H, there exists a whole family of strategies in optimal experimental design theory to optimise model selection in the most data-efficient manner, yet its application within the SINDy framework remains largely unexplored due to a lack of probabilistic interpretation of the learning result of SINDy. We

hope that by grounding SINDy in the Bayesian framework, we can encourage further exploration for more optimal strategies to learn about the system in the most data-efficient manner.

5 Conclusion and future work

In this work, we have presented a unifying Bayesian framework to compare Bayesian-SINDy with the original STLS-based SINDy. By grounding STLS in the Bayesian framework, we revealed how STLS can be a robust model selection method in the high-data low-noise limit due to the dominant contribution of the likelihood to the evidence. However, in the low-data high-noise limit, the Occam factor becomes more important, and Bayesian-SINDy's ability to penalise overfitting becomes crucial. The Bayesian framework also provides a more rigorous justification for selecting more parsimonious models than SINDy. Instead of selecting models by heuristically tweaking hyperparameters towards the Pareto front, the Bayesian framework formally defines the Occam factor, resulting in a more principled approach to model selection. This results in the Bayesian-SINDy algorithm, which provides a more concrete theoretical foundation for current SINDy-based approaches and a more robust framework for further extensions. Along with other recent publications highlighting the Bayesian perspective on SINDy, such as [33]'s demonstration of the convergence of Ensemble-SINDy to Bayesian statistics, we believe that many current SINDy extensions can benefit from the Bayesian framework to improve robustness in noisy and scarce datasets.

To demonstrate how rigorously defined Occam factor helps model selection in the low data regime, we substituted STLS with sparse Bayesian learning to create the Bayesian-SINDy algorithm. Notably, in all examples considered, the Bayesian-SINDy algorithm was able to recover the intended equations most of the time in hundreds of data points or fewer, demonstrating its data efficiency. We have also compared Bayesian-SINDy to SparseBayes, which is the foundation of previous Bayesian extensions to SINDy, and demonstrated the importance of considering how noise propagates through the nonlinear candidate functions and the variation of noise level over time.

Although Bayesian-SINDy has significantly improved data efficiency, its resilience to noise can be further improved. In particular, the cubic oscillator example has revealed the shortcomings of the Gaussian approximation, which limited the noise level Bayesian-SINDy can tolerate. Nevertheless, the Gaussian approximation is found to be broadly valid in other examples, resulting in robust learning performance in real-life data such as the Lynx-Hare population dynamics. To improve the proposed method further, one can seek a more accurate approximation of the noise distribution as it propagates through the library.

Aside from improving the approximation of the noise distribution, much work can be done to expand the current methodology. For example, while this work has primarily focused on retrieving desired equations from homogeneous datasets where data are synthesized from the same unknown parameters, our methodology can also combine multiple datasets that share the same underlying dynamics but with different unknown parameters. The original SINDy-based method can train parameterised models from datasets of different parameters if the parameters are known or measured as part of the dataset. However, fixing or knowing all parameters in real-world data is typically difficult or impossible to achieve. For example, in modelling cell motility, a population of cells may share a common model for their motility, yet each cell might be parameterised differently and it is difficult to accurately measure each parameter before learning the model. Because the evidence is derived by marginalising the parameters, it is independent of the specific parameter value and provides a mathematical framework to combine multiple datasets with varying unknown parameters.

Another transformative application of Bayesian-SINDy is its easy extension to quantify the information content in each data point, as we have briefly demonstrated in §4.3. By considering the information gain per data point, one can optimise the experimental design to maximise the information gain per data point, thereby significantly improving data efficiency. This is particularly useful in real-life applications such as system identification in control, where the goal is to learn about the system in the most data-efficient manner.

Lastly, it would also be interesting to benchmark Bayesian-SINDy with other recent advancement in SINDy, such as the ensembling technique [12] and the denoising method based on automatic differentiation [38]. In particular, recent work has established open standard for benchmarking different sparse system identification methodologies [39]. While we do not expect the Bayesian-SINDy to be the most noise-robust or most accurate in all scenarios, its relatively cheap computational cost and excellent data efficiency make it more suitable for real-time applications such as model identification in control.

Appendix A. Selection of hyperparameters values

Section 3 compares the learning performances of SINDy (STLS), Bayesian-SINDy and SparseBayes in four examples. To ensure a fair competition between algorithms, we tuned the hyperparameters of each algorithm to maximise their performance in each given example. The resulting hyperparameter values used in each example are summarised in the tables 2 and 3.

	Prior Variance α^{-1} (B-SINDy)	Weak Formulation	Finite Difference
Van der Pol	10^{2}	$\phi = (t^2 - 1)^2$, 9 points	8th order
Cubic Oscillator	1	$\phi = (t^2 - 1)^4$, 6 points	
Lorenz	25^{2}	, , ,	12th order
Lynx-Hare	10^{2}		8th order

Table 2: Hyperparameters used in each example in §3.

For SINDy (STLS), the threshold parameter λ is tuned in increments of the first significant figure to maximise the overall success rate at each noise level in the synthetic examples and tuned to recreate the Lotka-Volterra equation with the least number of spurious terms in the Lynx-Hare example (table 3). For Bayesian-SINDy, the choice for the prior variance α^{-1} is chosen to be of similar order to the expected parameter value. The values of the prior variance represent prior knowledge of the order of magnitude of the parameter value. In practice, this rough knowledge of the parameters' order of magnitude can usually be obtained from simple scaling arguments or physical intuition. Meanwhile, the SparseBayes algorithm has no hyperparameter that requires tuning, as both α and β are optimised by the algorithm.

	Noise level σ_x	Threshold Parameter λ
Van der Pol	0.1	0.4
	0.2	0.4
Cubic Oscillator	0.005	0.04
	0.01	0.06
Lorenz	0.025	0.2
	0.05	0.3
	0.1	0.4
	0.2	0.5
Lynx-Hare	N/A	0.025

Table 3: Optimal values for thresholding parameter λ in SINDy (STLS) in each example in §3.

The choices between weak formulation and finite difference are made according to the sampling frequency of the data compared to the typical frequency of the data. As per the discussion in §2.3, the weak formulation is preferred if the sampling frequency is found to be suitably higher than the typical frequency of the dynamics. Note that in practice, the application of weak formulation or finite difference should be decided according to the signal in the data before the learning takes place. Here, the weak formulation is used in the Cubic Oscillator example because it does not significantly distort the signal. In other examples, the finite difference is chosen because crucial high-frequency signals are filtered out when the weak formulation is used.

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