Less greedy equation learning: Balancing interpretability and expressivity through Bayesian model selection

Abstract

In the field of equation learning, exhaustively considering all possible combinations derived from a basis function dictionary is infeasible. Sparse regression and greedy algorithms have emerged as popular approaches to tackle this challenge. However, the presence of strong collinearities poses difficulties for sparse regression techniques, and greedy steps may inadvertently exclude important components of the true equation, leading to reduced identification accuracy. In this article, we present a novel algorithm that strikes a balance between comprehensiveness and efficiency in equation learning. Inspired by stepwise regression, our approach combines the coefficient of determination, $R^2$, and the Bayesian model evidence, $p(y|M)$, in a novel way. Through three extensive numerical experiments involving random polynomials and dynamical systems, we compare our method against two standard approaches, four state-of-the-art methods, and bidirectional stepwise regression incorporating $p(y|M)$. The results demonstrate that our less greedy algorithm surpasses all other methods in terms of identification accuracy. Furthermore, we discover a heuristic approach to mitigate the overfitting penalty associated with $R^2$ and propose an equation learning procedure solely based on $R^2$, which achieves high rates of exact equation recovery.

1 Introduction

Uncovering the underlying laws governing a system is essential for understanding its behavior, and mathematical equations serve as a concise representation of these laws. Equipped with such equations, predictions can be made, and valuable insights can be derived analytically. The pursuit of inferring these governing equations directly from observations has a long history, dating back to Johannes Kepler’s deduction of planetary motion laws in 1609. In modern times, significant progress has been made in automated equation inference using machine learning techniques, a discipline commonly referred to as ‘equation learning’ or ‘symbolic regression.’

In contrast to deep learning, equation learning focuses on maximizing model expressivity while minimizing complexity to ensure interpretability. This delicate balance between interpretability and expressivity constitutes the central challenge of equation learning. Other challenges include stable feature selection, solvability of learnt equations, computational feasibility, and handling limited data. Greedy algorithms and regularization techniques applied to regression models built from basis functions are commonly employed to address these challenges.

Greedy algorithms, like stepwise regression, explore the model space iteratively by evaluating scores for individual models. Although these methods are computationally efficient, they tend to significantly reduce the size of the considered model space, often leading to the exclusion of the true model from consideration. On the other hand, regularization transforms the search into an optimization problem, where all candidate models serve as points in the objective function landscape. However, finding the global minimum that corresponds to the true model is not guaranteed, and local minima may only accidentally lead to the true model. Collinearities among basis functions further complicate optimization algorithms due to the jagged nature of the optimization space.

In this work, we tackle the aforementioned challenges of equation learning by enhancing the stepwise regression approach in a less greedy way and exploring multiple score functions. Our method begins with an elimination
process that employs the computationally inexpensive coefficient of determination, $R^2$, in order to assess almost all individual candidate models belonging to a complexity class. Selecting the best models from that process enables subsequent model selection based on the Bayesian model evidence. The model evidence reflects the probability of a model being the true model for the given data, making it an ideal criterion for penalizing overfitting and addressing the central challenge of equation learning: achieving the optimal balance between interpretability and expressivity. By employing specifically tuned conjugate priors within an empirical Bayes framework, we circumvent the computationally demanding estimation of the evidence and instead derive it analytically.

To evaluate our approach, we employ artificially generated data from known models. This choice allows us to assess whether our strategy can uncover the ground truth model, serving as a testament to its ability to strike the delicate balance between expressivity and interpretability in realistic scenarios involving complex, analytically challenging models. We compare our method against two standard techniques, least absolute shrinkage and selection operator (LASSO) and least-angle regression (LARS), as well as four state-of-the-art methods available in the PySINDy package: sparse relaxed regression (SR3), forward regression orthogonal least squares (FROLS), sequentially thresholded least squares (STLSQ), and best subset selection via mixed-integer optimized sparse regression (MIOSR). Additionally, we propose evidence-based bi-directional stepwise regression (SR) for equation learning using our tuned model evidence. We evaluate the identification accuracy of approximately 80 scenarios for each system and assess the forecasting accuracy based on 100 initial values for each scenario and system, amounting to a total of around 8,000 tests. Our findings demonstrate that our proposed methods outperform other approaches in terms of identifying the correct model and can achieve competitive forecasting accuracy.

Our paper is organized as follows. We begin with a short overview of existing method in Section 2, and introduce our approach and the methods we compare to in Section 3. In Section 4 we present our numerical results, which we discuss in Section 5. We conclude in Section 6 and provide more details of our approach in the appendix.

2 Related work

The literature on equation learning can be roughly divided into three approaches: evolutionary algorithms, tree and neural network representations, and regularized regression. A well-known example of evolutionary algorithms is EUREQA, a commercial software used for symbolic regression Dubčáková (2011); Stoutemyer (2013). A recent open source alternative to EUREQA also using evolutionary algorithms is the python package pySR Cranmer (2023). Tree representations construct equations by combining basic operations and are then employed to minimize regularized objective functions Vaddireddy et al. (2020), using posterior sampling with sparsity-promoting priors Jin et al. (2019), or through mixed-integer linear programming Neumann et al. (2020). Similarly, neural network architectures have been used to represent equations, where network nodes are replaced by expression building blocks Martius & Lampert (2016); Sahoo et al. (2018); Werner et al. (2021). These neural networks are trained with a regularized minimizer applied to objective functions Rackauckas et al. (2020). Another approach, which allows for the incorporation of physics-informed properties such as symmetries, has also been developed Udrescu & Tegmark (2020).

These methods find applications in complementing deep neural networks to enhance generalization Arabshahi et al. (2018) and reducing the data requirement for training Yang et al. (2021). They are also utilized in the analysis of spatio-temporal biological data Nardini et al. (2020) and uncovering complex ecosystem dynamics Chen et al. (2019).

The above approaches are highly non-linear and typically involve complex optimization algorithms. A simpler approach is based on linear regression models, where features are replaced by basis functions derived from observed data. Regularized regression ensures sparse weight estimates on the basis functions Hastie et al. (2009), resulting in concise mathematical expressions. One prominent and widely used method for sparse regression is the LASSO with $\ell_1$-regularization Tibshirani (1996). The advantage of $\ell_1$-regularization is the convexity of the objective functions, which allows for efficient optimization. However, as we will discuss later, while sparse regression performs well with independent features in reconstruction tasks, the correlations
introduced by basis function expansion can lead to detrimental instability in equation learning. Recent advancements of LASSO, such as SR3, can be found in Zheng et al. (2019); Tibshirani & Friedman (2020).

Greedy algorithms like SR, FROLS, and STLSQ have proven to be more successful than LASSO in equation learning. The latter two algorithms are implemented in the Python package PySINDy, which is designed for the sparse identification of nonlinear dynamics (SINDy) Brunton et al. (2016). SINDy, initially proposed in Brunton et al. (2016) using STLSQ, has seen numerous extensions, including applications to partial differential equations Rudy et al. (2017), improved noise robustness through automated differentiation Kaheman et al. (2020), supplemented with a-posteriori (MAP) estimates Niven et al. (2020), re-weighted ℓ1-regularization Cortiella et al. (2021), relaxed regularization Champion et al. (2020), and most recently, best subset selection using MIOSR Bertsimas & Gurnee (2023).

In Bayesian linear regression, sparsity-promoting priors are used instead of ℓ0-regularization for equation learning Nayek et al. (2021). Thresholded sparse regression also employs sparsity-promoting priors Zhang & Lin (2018; 2021). By restricting basis functions to quadratic order, the linear structure of the models allows for deterministic results even in the case of the more challenging ℓ0-regularization Schaeffer et al. (2018).

3 Less greedy equation learning

The goal of equation learning is to find a function \( f(x) \) that accurately represents the relationship between the input variables \( x \) and the output variable \( y \). In the context of regression models, we consider a dataset consisting of \( N \) observations, where the inputs are organized in a design matrix \( X \) and the corresponding outputs are modeled by a random variable \( Y \). We can express the relationship between \( x \) and \( y \) as follows:

\[
Y = f(x) + \sigma Z, \tag{1}
\]

where \( x \) is a row of \( X \) (a feature vector), \( Z \) is a standard normal random variable, and \( \sigma^2 \) is the variance of the noise term. Our goal is to represent the unknown function \( f(x) \) using a basis function expansion: We assume that \( f(x) \) can be represented as a linear combination of \( p \) basis functions \( k_n(x) \),

\[
f(x) = \sum_{n=1}^{p} w_n k_n(x), \tag{2}
\]

where \( w_n \) denotes the weights associated with each basis function. We can construct a basis function matrix \( K \), with elements \( K_{nj} = k_n(x_j) \). The ordinary least squares (OLS) estimates for the weights \( w_n \) are given by Montgomery et al. (2012):

\[
\hat{w} = (K^TK)^{-1}K^Ty, \tag{3}
\]

where \( y \) represents a vector consisting of \( N \) samples of the output variable \( Y \). With the estimated weights \( \hat{w} \), we can make predictions \( \hat{y} = K\hat{w} \).

Common choices for the basis functions \( k_n(x) \) involve products and powers of the individual features \( x_j \). For example, we can have \( k_1(x) = x_1^2 \), \( k_2(x) = x_1x_2 \), \( k_3(x) = x_1x_3 \), and so on. By imposing restrictions on the maximum power of individual factors and the total number of factors in each \( k_n(x) \), we can control the total number \( p \) of basis functions. Equation learning involves selecting a small subset of \( k_n(x) \) for which the corresponding weights \( w_n \) are estimated, while setting the remaining weights to zero.

Once a choice of basis functions \( k_n(x) \) is made, the actual learning of the model using equation equation 3 is straightforward. However, the challenging part lies in the selection of the appropriate basis functions. On one hand, we require the model to have enough flexibility (expressivity) to minimize bias, ensuring that it captures the underlying relationship between \( x \) and \( y \). On the other hand, we want to avoid overfitting, which can lead to high variance in predictions. This trade-off between bias and variance guides the determination of the number of non-zero weights, denoted as \( m \), or equivalently, the model size. In addition to finding the appropriate model size, we also aim to identify the “correct” set of basis functions \( k_n(x) \), which corresponds to recovering the true \( f(x) \) from data generated by equation equation 1.
3.1 Regularized regression

A common way to avoid overfitting is to use regularization,

\[
\hat{w} = \arg \min_{w \in \mathbb{R}^p} \left[\|Kw - y\|_2^2 + \lambda\|w\|_q\right],
\]  

(4)

where \(\|w\|_q = |\sum_n |w_n|^q|^{1/q}\), and \(\lambda\) is the Lagrange parameter that sets the strength of the \(\ell_q\) penalty. The standard, sparsity-promoting choice is \(q = 1\) which is known as the LASSO (Tibshirani 1996). The choice \(q = 0\), for which \(\|w\|_0 = \sum \delta_{w_n,0}\) is the number of non-zero weight estimates, is often called best subset selection and requires specialized optimization algorithms (Zhu et al. 2020; Hastie et al. 2020), like MIOSR (Bertsimas & Gurnee 2023).

A relaxed regularization like SR3 can be introduced by letting the penalty act on an auxiliary variable \(u\), where distance of \(u\) to the actual weights is controlled by an additional regularization term, e.g. \(\|w - u\|_q\) (Zheng et al. 2019).

3.2 Model selection

Taking a different route, sparse solutions of equation 3 may be realized by model selection. A computationally cheap criteria for model selection is the coefficient of determination,

\[
R^2 = \frac{y^T K (K^T K)^{-1} K^T y}{y^T y},
\]  

(5)

here in a simplified form for standardized \(y\) (Montgomery et al. 2012).

However, \(R^2\) is known for its lack of overfitting penalty as it would just increase with decreasing sparsity. Alternatives like adjusted versions of \(R^2\) exist to incorporate an overfitting penalty, but here we make use of the Bayesian model evidence

\[
p(M) \propto p(y|M) = \int d\sigma \int dw p_\sigma(y|w, \sigma, M) p_\sigma(w, \sigma),
\]  

(6)

where a selection of basis functions \(k_n(x)\) defines a model \(M\) and a likelihood distribution \(p_\sigma(y|w, \sigma, M)\) via equation 1 and equation 2. For a conjugate prior \(p_\sigma(w, \sigma)\), the marginalization above can be done analytically and \(p(y|M)\) is known exactly, as detailed in appendix A.

The use of \(p(y|M)\) is often motivated by its excellent overfitting penalizing properties, which can be ascribed to the fact that \(p(y|M)\) is proportional to the probability \(p(M)\) of the model \(M\) being the true model for the data \((y, X)\) (Murphy 2012). The downside of using \(p(y|M)\) is the imperative to specify \(p_\sigma(w, \sigma)\) even in cases of scarce prior knowledge, which can have a significant impact on \(p(y|M)\). Here we use the empirical Bayes method to fix \(p_\sigma(w, \sigma)\) and exploit the normality property of linear regression which implies that estimates \(\hat{w}\) are normally distributed with the mean given by the true values of \(w\) and the variance given by

\[
\delta^2 = \frac{(y - \hat{y})^T(y - \hat{y})}{N - p}
\]  

(7)

The details of this procedure is included in appendix A.

3.3 Stepwise regression (SR)

A standard greedy algorithm to select basis functions \(k_n(x)\) is stepwise regression in which a criterion like adjusted \(R^2\), \(F\)-statistics or other information criteria (e.g. AIC, BIC) are used (Montgomery et al. 2012). Here we promote the evidence \(p(y|M)\) as criterion, which is rarely used for SR due to its intricacies in terms of prior selection and computational cost (Hastie et al. 2009). Our stepwise procedure is bi-directional, that is, we start with an empty model and select the \(k_n(x)\) that maximizes \(p(y|M)\), add a second \(k_n(x)\) maximizing \(p(y|M)\), and so on (forward selection). Once \(p(y|M)\) cannot be increased further by adding more \(k_n(x)\)
Figure 1: Schematic representation of algorithm LG-\(R^2\) and LG-\(p\mathcal{M}\). Starting point is the set of candidate models built from \(p\) basis functions \(k_n(x)\). For fixed model size \(m\), \(R^2\) is computed for all \(\binom{p}{m}\) candidate models. Incrementing \(m\), top models in terms of \(R^2\) are collected, from which basis functions \(k_n(x)\) are rated based on counts of \(k_n(x)\) in these top models. Typically, incrementing \(m\), new \(k_n(s)\) with high rates are found and selected for the inferred model \(\mathcal{M}^*\) by algorithm LG-\(R^2\) while \(m\) is smaller than the true model size. The iteration in \(m\) therefore terminates if no new \(k_n(s)\) are selected for \(\mathcal{M}^*\). Basis functions \(k_n(x)\) that are hardly selected are not considered for building candidate models in the next \(m\)-step. The top models selected by \(R^2\) are scored again by \(p(\mathcal{M})\) of which the best model is the output of algorithm LG-\(p(\mathcal{M})\).

3.4 Less greedy (LG)

A non-greedy algorithm would consider all \(2^p - 1\) combinations of \(k_n(x)\) which obviously is computationally infeasible. However, since we are interested in parsimonious models, we may choose a small model size \(m\) and explore all \(\binom{p}{m}\) possible combinations of \(k_n(x)\) within that budget. For a fixed model size, overfitting
penalty is not important, and we may use $R^2$ which is particularly cheap to compute for standardized data, c.f. equation \[5\]. In this way, we can single out the best models in terms of $R^2$ for different budgets $m$.

To find the best model size $m$, we employ the overfitting penalty property of the model evidence $p(y | \mathcal{M})$. Starting with $m = 1$, we compute $p(y | \mathcal{M})$ for the best models selected by $R^2$, and continue to do so for increasing $m$ until a stopping criterion is reached.

As a stopping criterion, we may use the first decrease of $p(y | \mathcal{M})$, but we propose a heuristic criterion purely based on $R^2$ which proved superior in our study. The $R^2$ criterion we propose builds on the observation that the true $k_n(x)$ have a tendency to be consistently selected in the top $R^2$ models. We therefore keep increasing $m$ until no new $k_n(x)$ is selected consistently, and then build the inferred model from those consistently selected terms. Since ${n \choose m}$ still becomes very large for larger model sizes, we add an additional pruning step, in which all $k_n(x)$ that have consistently not been selected are removed from the basis expansion.

The stopping criterion and the pruning step classifies our procedure as a greedy algorithm. However, due to the exhaustive search for each considered $m$, we consider a drastically larger model space than other greedy algorithms and are as such significantly less greedy.

While this is a heuristically developed method, it is motivated by the reasonable assumption that the maximization of $R^2$ is dominated by the true $k_n(x)$. Therefore, as soon as we look at the $R^2$ values of models one term larger than the true model, the procedure is forced to randomly select one extra term in addition to the consistently selected true terms.

For a direct comparison of this rather heuristic selection method with an established method, we also select the model that maximizes $p(y | \mathcal{M})$ out of the top models selected by $R^2$ across all model sizes. To the first method solely based on $R^2$ we refer to as LG-$R^2$, to the second method using a final selection via $p(y | \mathcal{M})$ we refer to as LG-$p(\mathcal{M})$. We illustrate our procedure in figure 1. More details, explicit algorithms, and an illustration of the claims made can be found in appendix C.

It is worth noting that our procedure can also be used in cases where $p(y | \mathcal{M})$ needs to be estimated by computationally costly evidence estimators, as we effectively reduce the pool of candidate models to just a few.

### 3.5 Model class

Apart from the equations that can directly be written in the form of equation \[2\], a prominent application of equation learning is the learning of dynamical systems,

\[
\dot{x}(t) = f(x(t)), \tag{8}
\]

where $\dot{x}(t)$ denotes the time derivative of $x(t)$. To map this problem to equation \[2\] the response variable can be computed from finite differences $y_i = \frac{y(t_{i+1}) - y(t_i)}{\Delta t}$ for a fixed time step $\Delta t$.

A restriction for the regression models to stay linear in its parameters is that parameters of basis functions may only enter as weights $w$. Basis functions like $e^{ax}$, $\ln(a + x)$, $\cos ax$, $x^a$, $\frac{1}{(a + x)^m}$, ... with internal parameter $a$ are not suitable.

This restriction might seem quite limiting. On the other hand, the function $f(x(t), w)$ defining a dynamical system typically is linear in its parameters $w$. The reason for that is that these functions often reproduce when differentiated, which can be used to eliminate these functions, retrieving the standard linear form in equation \[2\] Some special functions like Bessel, Hankel, Struve and Meijer functions are even defined as solutions of differential equations linear in their coefficients. In general, by considering the differentiated response variable, $y \rightarrow \frac{dy}{dx} \sim \frac{y(x_{i+1}) - y(x_i)}{x_{i+1} - x_i}$, if necessary to higher order, we can learn a surprisingly broad class of equations relating $y$ and $x$, even relations that do not exist in closed form.

### 4 Numerical experiments

We compare all introduced methods (LASSO, LARS, LG-$R^2$, LG-$p(\mathcal{M})$, SR, SR3, FROLS, STLSQ, MIOSR) in three numerical experiment. For LASSO and LARS, we use the PYTHON package scikit-learn [Pedregosa](https://scikit-learn.org/).
Table 1: Equation learning techniques used in this study building on regression models as in equations 1, 2.

<table>
<thead>
<tr>
<th>Acronym</th>
<th>Full name</th>
<th>Description</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>LASSO</td>
<td>Least absolute shrinkage and selection operator</td>
<td>Regularized regression as in equation 4 for $q = 1$.</td>
<td>Tibshirani (1996).</td>
</tr>
<tr>
<td>LARS</td>
<td>Least angle regression</td>
<td>Forward stepwise regression using correlations between basis functions $k_n$ and residuals.</td>
<td>Efron et al. (2004).</td>
</tr>
<tr>
<td>LG–$R^2$</td>
<td>Less-greedy $R^2$ elimination</td>
<td>For an increasing model size $m$, the $R^2$ score is calculated for all candidate models. Based on the number of times basis functions $k_n$ contribute to the models with largest $R^2$, kernel functions $k_n$ are rated and the lowest rated $k_n$ are excluded. Once no new highly rated $R^2$ are found, the iteration in $m$ terminates and the model with largest $R^2$ is returned.</td>
<td>This work, appendix C.</td>
</tr>
<tr>
<td>LG–$p(M)$</td>
<td>Less-greedy Bayesian model selection</td>
<td>Same as LG–$R^2$, but a selection of models is returned based on maximal Bayesian model evidence $p(M)$.</td>
<td>This work, appendix C.</td>
</tr>
<tr>
<td>SR</td>
<td>Bi-directional stepwise regression</td>
<td>Iterated forward and backward selection using the model evidence Bayesian model evidence $p(M)$ as score.</td>
<td>Hastie et al. (2009), this work for $p(M)$, appendix B.</td>
</tr>
<tr>
<td>SR3</td>
<td>Sparse relaxed regression</td>
<td>The regularization is put on an auxiliary variable $u$ and an extra distance term like $|w-u|_q$ is added to equation 4.</td>
<td>Zheng et al. (2019).</td>
</tr>
<tr>
<td>FROLS</td>
<td>Forward regression orthogonal least-squares</td>
<td>Forward selection stepwise regression with score given by equation 4 for $q = 2$.</td>
<td>Billings (2013).</td>
</tr>
<tr>
<td>STLSQ</td>
<td>Sequentially thresholded least squares</td>
<td>Backward selection stepwise regression using results of regularized regression for $q = 2$ in equation 4.</td>
<td>Brunton et al. (2016).</td>
</tr>
<tr>
<td>MIOSR</td>
<td>Best subset selection via mixed-integer optimized sparse regression</td>
<td>Formulation of regularized regression equation 4 for $q = 0$ as a mixed-integer linear program and specialized algorithms.</td>
<td>Bertsimas &amp; Gurnee (2023).</td>
</tr>
</tbody>
</table>

et al. (2011), for LG–$R^2$, LG–$p(M)$ and SR we use our own implementation, and for SR3, FROLS, STLSQ, MIOSR we use the python package PySINDy (de Silva et al. (2020), Kaptanoglu et al. (2022). As mixed-integer optimizer for MIOSR we used gurobi with an academic license Gurobi Optimization, LLC (2023). For all methods from scikit-learn and PySINDy we used 5-fold cross validation and 3 refinement steps to determine optimal hyperparameters for each application separately. Our own methods, LG–$R^2$ and LG–$p(M)$, are not very sensitive to hyperparameters and we worked out the universally best values which, in contrast to the methods we compare to, were used for all applications. The SR method we implemented comes without hyperparameters. Table 4 gives an overview over all considered methods. More details can be found in appendix C.
Figure 2: Identification accuracy of learning 100 random polynomials measured in terms of the difference $N_{\text{diff}}$ between the number of true terms found and wrong terms found. We generated polynomials with 2, 3 and 4 terms (left to right), which also constitutes the highest possible value for $N_{\text{diff}}$ and is indicated by a gray horizontal line. The percentages above this line indicate how often all exact terms and no wrong terms have been recovered.

In all experiments, the data is artificially generated, with the advantage that we know the true model. In the first experiment, we assess the identification accuracy of 100 random polynomials. Since PySINDy is tailored to learning dynamical systems and cannot directly be applied to learning polynomials, we omitted those methods in this experiment. In the second and third experiment the methods are applied to two (chaotic) dynamical systems, where, for the sake of readability, we omitted LARS due to its similar performance compared to LASSO.

In all experiments we have three features $\mathbf{x} = (x_1, x_2, x_3)$, and we used a basis function expansion of polynomials where the power of individual factors was limited to a maximum of 4, and the combined power of terms $k_n(\mathbf{x})$ to a maximum of 6. For instance, $k(\mathbf{x}) = x_1^2 x_2 x_3^2$ would be a valid basis function, while $k(\mathbf{x}) = x_1^5 x_2$ would be excluded for exceeding the individual power limit of 4, as would $k(\mathbf{x}) = x_1^4 x_2 x_3^3$ for exceeding the combined power limit of 6. The resulting feature dimension of $K$ is $p = 72$.

The results of each experiment are illustrated statistically by boxplots, where the box indicates the interquartile range, the whiskers extend by a factor 1.5 beyond the box, and outcomes exceeding the whiskers are shown as individual symbols. The median is shown as a darker horizontal line, the mean as a circle.

4.1 Random polynomials

We randomly generated 100 polynomials with 2, 3, and 4 non-zero weights respectively. We restricted the terms of the polynomials to have a maximum collective power $M_2 = 4$, where individual features are restricted to maximum power $M_1 = 2$. The non-zero weights are randomly selected with equal probabilities and their values are uniformly sampled from the set $[-4, -1] \cup [1, 4]$. We generated artificial data $\mathbf{X}$ for each polynomial by sampling from normal distributions with means randomly selected from the interval $[-20, 20]$ and standard deviations such that 5% of their probability mass overlap respectively. For the polynomials sized 2, 3, and 4 we generated $N = 20$ and $N = 65$ and $N = 95$ datapoints, respectively. Plugging $\mathbf{X}$ into equation 1 with $f(x_i)$ given by the random polynomial, and corrupting the output with normal noise with standard deviation $\sigma = 0.01$, we generate data for the response variable $y_i$.

The statistics of the identification accuracy for the 100 polynomials are shown in figure 2.

4.2 Lorenz system

As a first dynamical system to test our method, we use the chaotic Lorenz system defined as

$$\dot{x}(t) = \epsilon (y(t) - x(t)),$$

$$\dot{y}(t) = x(t) (\rho - z(t)) - y,$$

$$\dot{z}(t) = x(t)y(t) - \beta z(t).$$  \hspace{1cm} (9)
The parameters are fixed to its standard values $\epsilon = 10$, $\rho = 28$ and $\beta = 8/3$. As initial condition, we use $(x_0 = -8, y_0 = 8, z_0 = 27)$. We obtain between $N = 200$ and $N = 10000$ data points by solving equation 9 numerically for timestep widths between $\Delta t = 0.001$ and $\Delta t = 0.1$. We corrupt the solutions with normal noise levels between $\sigma = 0.001$ and $\sigma = 0.1$. Forcing the simulation time to be larger than $T = 2$, we thus have 80 different scenarios with varying $N$, $\Delta t$, $\sigma$ and $T$. We apply the equation learning methods to all scenarios to obtain some statistics on identification accuracy and mean-absolute error (MAE).

We measure the identification accuracy as the number of correctly identified equations comprising the model equation 9. The MAE is obtained by solving the learnt dynamical system equations numerically for randomly selected initial values, and comparing the result with the solution we get solving the true equations 9 for the same initial condition.

To further increase the sample size for the statistical evaluation of the methods, and to exclude the possibility to have a particular initial condition that suits a certain method by chance, we sample 100 initial values from the Lorenz attractor and solve each learnt model from all scenarios and equation learning models for the same 100 initial values.
In figure 3 we show the statistical results of our experiment. To emphasize the goal to parsimoniously learn models, we also plot the statistics of the MAE multiplied by the size $N_{\text{model}}$ of the learnt model. In addition to the overall statistics, we also show the dependency on $N$ and $\Delta t$ (splitting the data up in terms of $\sigma$ or $T$ did not reveal any insights).

4.3 Rabinovich-Fabrikant equations

![Figure 4: Statistical results from models learnt from data created by solving the Rabinovich-Fabrikant system equation (10).](image)

As a second dynamical system, we use the Rabinovich-Fabrikant equations

\[
\begin{align*}
\dot{x}(t) &= y(t)(z(t) - 1 + x(t)^2) + \gamma x(t), \\
\dot{y}(t) &= x(t)(3z(t) + 1 - x(t)^2) + \gamma y(t), \\
\dot{z}(t) &= -2z(t)(\alpha + x(t)y(t)).
\end{align*}
\] (10)

We choose the parameter values $\alpha = 0.14$ and $\gamma = 0.1$, and the initial conditions $(x_0 = -1.5, y_0 = 0, z_0 = 1)$. We consider 84 scenarios comprising values $N = 1000$ to $N = 16000$, $\Delta t = 0.001$ to $\Delta t = 0.1$, and $\sigma = 0.0001$ and $\sigma = 0.01$, where we enforce $T \geq 5$. We solve equation (10) numerically and corrupt the solutions with noise as for the Lorenz system.

The results are shown in figure 4 in the same fashion as for the Lorenz system.

The PYTHON code to reproduce these results have been included in the supplementary material.
5 Discussion

In this section, we discuss the results of the numerical experiments shown in the previous section.

We begin with the learning of the random polynomials and stress the difficulty of this task, which consists in various aspects: i) Randomly generated polynomials are not hand-picked examples where some fine-tuning is possible. ii) The artificially generated data was not tested to represent the polynomial uniquely. iii) Even if an inferred polynomial deviating from the true polynomial would describe the data well, it does not contribute positively to the identification accuracy.

With these aspects in mind, it is quite remarkable that overall many of the polynomials could be recovered, as shown in figure 2. Particularly striking is the success rate of LG-$R^2$ solely based on $R^2$, whereas the LASSO and LARS clearly overfit in terms of model size. However, the overall accuracy clearly declines with increasing number of non-zero terms. This can be explained with a higher chance of terms being selected spuriously leading to too early or too late stopping of exploring the required number of terms. Also the risk of erroneously removal of true terms from $K$ increases. We are confident that more refined stopping and removal criteria can overcome these inaccuracies. In combining several criteria, we see an opportunity to improve the accuracy even further.

The Lorenz system has between 2 and 3 terms per equation, and as such was learnt quite successfully, as the first row of figure 3 shows. The highest identification accuracy is again achieved by LG-$R^2$, the LASSO was not able to identify any equation, and of the SINDy methods the relaxed regularized regression SR3 and the best subset-selection using mixed integer optimization MIOSR performed best. In terms of MAE, the LASSO is among the best, but requires significantly larger models is shown by $N_{model} MAE$. Most of the methods are relatively robust against $N$ and $∆t$. Interestingly, $N_{model} MAE$ worsens with more datapoints in the case of MIOSR but improves for the LASSO, signifying that $ℓ_0$ regression “sees” more in the data than there is if given enough data, while $ℓ_1$ requires more data to produce smaller models (c.f. equation 4). Regarding robustness against larger timesteps, it turns out that MIOSR is particularly sensitive with deteriorating performance for smaller $Δt$, as well as FROLS and the LASSO to a lesser extent.

The Rabinovich-Fabrikant system turned out to be particularly difficult to learn. The identification accuracy for the LG methods, the SR using $p(\mathcal{M})$ and the relaxed regularized SR3 was on average just below 1 equation, while the other methods failed to identify any equation in all scenarios. No method was able to learn the complete system of equations correctly. Interestingly, the MAE was still reasonable, in particular the LASSO performed well. Splitting up in $N$ and $Δt$ do not provide any additional insight in particular and are just shown for completeness.

A particular problem in learning dynamical systems is that there is no guarantee that the learnt models are solvable. In cases where the numerical solutions failed, we excluded the results from the statistics and kept a record of how often this happened for the various equation learning methods. For the Lorenz system, LASSO, SR3 and STLSQ failed at about 1%, MIOSR at about 20%, and the other methods always produced solvable systems. This is a little different for the Rabinovich-Fabrikant system, where all methods produced unsolvable models between about 1% and 15%, with STLSQ the most problematic followed by SR3 and LG-$R^2$.

6 Conclusions

We extensively tested our methods LG-$R^2$, LG-$p(\mathcal{M})$ and the bi-directional SR employing $p(\mathcal{M})$ against the standard methods LASSO and LARS, as well as state-of-the-art methods SR3, FROLS, STLSQ and MIOSR. To our knowledge, we are the first to explore equation learning based on an exhaustive candidate model evaluation outperforming existing state-of-the-art methods in terms of identification accuracy, and at least on equal terms regarding forecast quality.

A disadvantage of our less greedy approach is the higher computational cost and the absence of clear time complexity measures. However, a direct advantage is that the model evaluation is trivially parallelizable. Also the little amount of data needed for high success rates is striking – tests on two sets of random polynomials where even done with less datapoints than rows in $K$. Testing candidate models individually also allows for great flexibility when it comes to constraints or conditions on models such as solvability, as well as eliminating
the risk of getting stuck in local minima of an objective function. It also allows combining several criteria for model and feature selection, in particular complementing existing methods in an independent way with the potential of synergy effects.

The observation that learnt chaotic model comprising all true terms plus a few extra terms can decrease the MAE has an interesting implication worth exploring in a future work: It seems possible to learn correction terms from data that lead to a better forecast horizon than the true model itself.

Finally, we contributed towards the utilization of the Bayesian model evidence $p(M)$ in equation learning. Here, we benefit from using a conjugate prior for which $p(M)$ can be computed analytically, and showed in our numerical experiments that our choice of empirical prior is well suited for the tasks considered here. However, in general, one would like to have the freedom to select any prior which may entail particularly computationally costly evidence estimators. Performing the less greedy search with $R^2$ (and possibly other criteria) can boil down the number of candidate models to a feasible number, an approach planned to be explored in a future work.

Considering these possibilities and the promising identification accuracy achieved, we hope to open a new avenue of equation learning.

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(after acceptance)

References


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A Linear regression

In this section, we show how linear regression can be applied to equation learning, which sets the basis of this work. Details to regression can be found in Montgomery et al. (2012).

Starting point are \( N \) observations \((y_i, x_{ij})\), where the index \( i \) denotes data points, and the index \( j \) denotes features. We assume that the response (dependent) variable \( y \) is given as a function of explanatory (independent) variable \( x \),

\[
y = f(x) + \sigma z, \tag{11}
\]

where \( f(x) \) defines the model, \( z \) is a standard normal random variable, and \( \sigma^2 \) is the variance of the noise term. The explanatory observations are often organized in terms of a design matrix \( X \), with features in columns and datapoints in rows, \( X_{ij} = x_{ij} \).

We assume that \( f(x) \) can be given in terms of \( p \) basis functions \( k_n(x) \),

\[
f(x) = \sum_{n=1}^{p} w_n k_n(x), \tag{12}
\]

with weights \( w_n \), known as basis function expansion. Similar to the design matrix, we can define a basis function design matrix \( K \) with elements \( K_{in} = k_n(x_i) \).

The ordinary least squares (OLS) estimates for \( w_n \) are known to be

\[
\hat{w} = \arg \min_{w \in \mathbb{R}^p} \| Kw - y \|_2^2 = (K^T K)^{-1} K^T y \tag{13}
\]

with the \( q \)-norm

\[
\| w \|_q = \left[ \sum_n |w_n|^q \right]^{1/q}. \tag{14}
\]

Predictions based on the OLS estimates are then given by \( \hat{y} = K \hat{w} \).

\[
\hat{y} = K \hat{w}. \tag{15}
\]

From the normality of linear regression, it is known that the estimates \( \hat{w} \) follow a normal distribution with the mean given by the true values for \( w \) and the variance given by \( \hat{\sigma}^2 = \frac{(y - \hat{y})^T (y - \hat{y})}{N-p} \). We will use these properties later to empirically define the prior in the Bayesian description.

In view of equation 13, once a choice of \( k_n(x) \) is made, the actual learning of the model is straight forward. The difficult part is the choice of \( k_n(x) \): on the one hand we require sufficient expressivity of the model to minimize bias, on the other hand we want to avoid overfitting to minimize variance of predictions. This bias-variance trade-off essentially dictates the number of \( k_n(x) \), i.e. the effective dimension of feature space or complexity. Apart from the appropriate model size, we also seek the "correct" \( k_n(x) \), in the sense that the true \( f(x) \) is recovered from data generated by equation 11.

A common approach to avoid overfitting is regularization,

\[
\hat{w} = \arg \min_{w \in \mathbb{R}^d} \left[ \| Kw - y \|_2^2 + \lambda \| w \|_q \right], \tag{16}
\]

where \( \| w \|_q = \left[ \sum_n |w_n|^q \right]^{1/q} \), and \( \lambda \) is the Lagrange parameter that sets the strength of the \( \ell_q \) penalty. Common choices for \( q \) are \( q = 2 \) (Ridge regression), \( q = 1 \) (the standard, sparsity promoting choice known as the LASSO), or combinations such as elastic net. A special case of regularization is \( q = 0 \), for which \( \| w \|_0 = \sum_n \delta_{w_n,0} \) is the number of non-zero weight estimates – the regression procedure with this penalty is often called best subset selection and requires specialized optimization algorithms.
A standard measure for goodness of fit is the coefficient of determination, \( R^2 \), which relates the variance explained by the prediction \( \hat{y} \) to the variance of the response variable \( y \),

\[
R^2 = 1 - \frac{(y - \hat{y})^T (y - \hat{y})}{(y - \bar{y})^T (y - \bar{y})}
\]  

(17)

Assuming standardized \( y \), \( R^2 \) can be simplified to

\[
R^2 = 1 - \frac{y^T y - y^T K \hat{w} - \hat{w}^T K^T y + \hat{w}^T K^T K \hat{w}}{(y - \bar{y})^T (y - \bar{y})}
\]  

(18)

\[
= 1 - \frac{y^T y - y^T K \hat{w}}{(y - \bar{y})^T (y - \bar{y})}
\]  

(19)

\[
= 1 - \frac{y^T y - \hat{w}^T y}{(y - \bar{y})^T (y - \bar{y})}
\]  

(20)

\[
= 1 - \frac{y^T y}{y^T y}
\]  

(21)

\[
= \frac{y^T K (K^T K)^{-1} K^T y}{y^T y}
\]  

(22)

where we plugged in equation 15 in the first line, in the 2nd line we used that \( K^T y = K^T K \hat{w} \), in the 3rd line that \( y^T K \hat{w} = \sum_{jk} y_j K_{jk} \hat{w}_k \), \( \sum_{jk} \hat{w}_k K_{jk} y_j = \hat{w}^T K^T y \), in the 4th line we assumed that the mean \( \bar{y} = 0 \) due to centering to zero, and in the last line we used \( \hat{w} = (K^T K)^{-1} K^T y \). \( R^2 \) is essentially a quadratic form for \( y \) where the coefficient matrix is \( K \) multiplied with its pseudo-inverse. For sparse weight estimates \( \hat{w} \), \( R^2 \) is extremely efficient to compute.

A value of \( R^2 \) close to 1 signifies good predictions \( \hat{y} \). However, it is well known that \( R^2 \) can always be brought closer to 1 by increasing the number of features in the model, thus essentially lacking any overfitting penalty if used naively.

For the purpose of adequate model selection, it is helpful to formulate regression in a Bayesian setting. The main step to this end is defining the likelihood distribution for \( y \). In the simple case of equation 11, \( y \) is normally distributed,

\[
p_n(y|w, \sigma) = N(\mu, \sigma),
\]  

(23)

where the mean vector is given by \( \mu_i = f(x_i) \). Maximization of the log-likelihood reproduces the OLS result equation 13. Encoding existing information on the weights \( w_n \) as a prior distribution \( p_{pr}(w, \sigma) \), Bayes’ formula implies for the posterior distribution

\[
p_{ps}(w, \sigma|y) = \frac{p_n(y|w, \sigma) p_{pr}(w, \sigma)}{p(y)},
\]  

(24)

where the normalization factor \( p(y) \) is known as the evidence or marginal likelihood.

The distributions in equation 24 also depend on the choice of model \( M \) given by the representation of \( f(x_i) \) in terms of \( K \). Adding \( M \) as a condition, and rewriting Bayes’ formula on the level of models,

\[
p_{ps}(M|y) = \frac{p(y|M) p_{pr}(M)}{p(y)},
\]  

(25)

we see that the evidence \( p(y) \) is in fact the model-likelihood \( p(y|M) \) and as such proportional to the model-posterior (assuming constant model-prior \( p_{pr}(M) \) for simplicity). Therefore, if we could maximize \( p(y|M) \) over \( M \), we would in fact identify the model \( M \) that most likely explains the observations \( y \).

From equation 24 it follows that the evidence is given by

\[
p(y|M) = \int d\sigma \int d\omega p_n(y|w, \sigma, M) p_{pr}(w, \sigma),
\]  

(26)
which in general is not solvable analytically, and also poses a particularly tough numerical challenge [1999]. Fortunately, by choosing \( p_{\text{pu}}(w, \sigma) \) conjugate to \( p_{\text{li}}(y|w, \sigma, \mathcal{M}) \), the integral becomes solvable analytically. The conjugate prior, however, is not necessarily the sensible choice from the inference point of view. In fact, making a good choice for the prior is a much debated problem [2015]. Here, we demonstrate that for the purpose of linear equation learning, the conjugate prior is a suitable choice, if hyper-parameters are distilled from data. The question whether other choices for the prior would perform significantly better is left for future research.

The conjugate prior for the likelihood equation [23] is the gamma-normal distribution O'Hagan & Kendall [1994]

\[
p(w, \tau | \mu, M, k, \vartheta) = \frac{\sqrt{\det M}}{(2\pi)^{p/2}\Gamma(k)^2} \tau^{p/2+k-1} e^{-\frac{1}{2} \frac{1}{\tau} (w-\mu)^T M (w-\mu) - \frac{\tau}{\vartheta}}
\]  

(27)

with mean vector \( \mu \) and precision matrix \( M \) for the weights \( w \), and shape \( k \) and scale \( \vartheta \) for the precision \( \tau = 1/\sigma^2 \). Plugging equation [27] and equation [23] into equation [26] and performing the integration, we obtain for the log-evidence per data-point the closed expression

\[
\frac{1}{N} \ln p(y|M) = \frac{1}{2N} \ln \frac{\det M}{\det A} - \frac{1}{2} \ln 2\pi - \left( \frac{1}{2} + \frac{k}{N} \right) \ln \left( \frac{\xi}{2} + \frac{\vartheta}{2} \right)
\]

\[
- \frac{k}{N} \ln \vartheta + \frac{1}{N} \ln \Gamma \left( \frac{N}{2} + k \right) - \frac{1}{N} \ln \Gamma(k)
\]

(28)

with \( A = K^T K + M \), \( b = K^T y + M \mu \), and \( \xi = y^T y + \mu^T M \mu - b^T A^{-1} b \).

\[
A = K^T K + M,
\]

\[
b = K^T y + M \mu,
\]

\[
\xi = y^T y + \mu^T M \mu - b^T A^{-1} b
\]

(29)

(30)

(31)

We detail the calculations in appendix B.

Owing to the normality of linear regression, and from standardizing the data, it is reasonable to assume the following parameters for the prior: For the mean vector, we choose \( \mu = \hat{w} \), and the precision matrix is \( M \) is taken to be diagonal with elements \( \text{diag}(M) = \frac{1-p/N}{y_i^2 - \hat{w}_i^2 K^T y} \), resulting in normal distributions broadened by a factor \( N \) to make the prior more uninformative. The gamma distribution entering equation [27] has the mode \((k - 1)\vartheta\), which we set to 1 due to standardized \( y \). The scale is set to \( \vartheta = 1/2 \) which appears to be broad enough for an uninformative prior.

For completeness, we mention a few more selection criteria used for comparison in this work. The adjusted \( R^2 \) [2012]

\[
R^2_{\text{adj}} = 1 - \frac{N - 1}{N - p - 1} (1 - R^2)
\]

(32)

equips the usual \( R^2 \) with an overfitting penalty. The Akaike information criterion (AIC) measures the loss of information by using the inferred model instead of the (unknown) true model [2012], and similarly but derived from the model evidence equation [34] in the big data limit, follows the Bayesian (Schwarz) information criterion,

\[
\text{AIC} = 2p_{\text{li}}(y|\hat{w}, \hat{\sigma}) - 2p, \quad \text{BIC} = p_{\text{li}}(y|\hat{w}, \hat{\sigma}) - 2p \ln N
\]

(33)

Similarly, but derived from the model evidence equation [34] in the big data limit, is the Bayesian (Schwarz) information criterion,

\[
\text{BIC} = p_{\text{li}}(y|\hat{w}, \hat{\sigma}) - 2p \ln N.
\]

(34)

Apart from the equations that can directly be written in the form of equation [12] a prominent application of linear equation learning is the sparse identification of dynamical systems,

\[
\dot{x}(t) = f(x(t)),
\]

(35)
where $\dot{x}(t)$ denotes the time derivative of $x(t)$. To map this problem to equation 12, the response variable can be computed from finite differences $y_i = \frac{x_{i+1} - x_i}{\Delta t}$ for a fixed time step $\Delta t$.

A restriction for the regression models to stay linear in its parameters is that the parameters of basis functions only enter as weights $w$. Basis functions like $e^{ax}$, $\ln(a + x)$, $x^a$, $\frac{1}{(a + x)^m}$, ..., (36)

with internal parameter $a$ are not suitable.

This restriction might seem quite limiting. On the other hand, the function $f(x(t), w)$ defining a dynamical system typically is linear in its parameters $w$. The reason for that is that functions shown in equation 36 often reproduce when differentiated, which can be used to eliminate these functions, retrieving the standard form shown in equation 12 and equation 11 being linear in parameters. Some special functions like Bessel, Hankel, Struve and Meijer functions are even defined as solutions of differential equations.

In general, by considering the differentiated response variable,

$$y \mapsto \frac{dy}{dx} \simeq \frac{y(x_{i+1}) - y(x_i)}{x_{i+1} - x_i},$$

(37)

if necessary to higher order, we can learn a surprisingly broad class of equations relating $y$ and $x$, even relations that do not exist in closed form. Here, we restrict ourselves to dynamical systems and leave the full exploration of learning in this broad model class for future work.

While many equations with non-linear parameters can be rewritten in linear form by differentiation as explained above, some functions like $x^a$, $\frac{1}{(a + x)^m}$ can only be reduced to fractions or require many differentiations which can give rise to numerical issues. Therefore, if we were able to include fractions in the basis function expansion, we could expand the model class even further. The problem is that basis functions like $\frac{1}{1 + x}$ are divergent for certain values of $x$, and are also quite limiting in their form.

It is, however, possible to modify the regression model to also incorporate fractions. In its simplest form, we may consider

$$y = \frac{\sum_{n=1}^{p} w_n k_n(x) + \sigma z}{\sum_{m=1}^{p} v_m k_m(x)}$$

(38)

with different (sparse) weights $v_m$ but same set of basis functions for the denominator. For the next step, we assume that $k_n(x)$ is part of the numerator, that is $w_1 \neq 0$, and we can rewrite

$$k_1(x) = \sum_{m=1}^{p} \frac{v_m}{w_1} y k_m(x) - \sum_{n=2}^{p} \frac{w_n}{w_1} k_n(x).$$

(39)

In this form, $k_1$ takes the role of the response variable, and we have a second set of basis functions given by $yk_m$. For a given model of this form, the weights also follow deterministically from equation 13 only $w_1$ needs to be determined from a 1-dimensional numerical root-finding algorithm.

A similar idea has been proposed in Kaheman et al. (2020), where a $\ell_0$ regularized objective function needs to be minimized for each possible basis function taking the role of $k_1$ above. Our less greedy strategy naturally includes this procedure as a straight forward possibility, which is planned to be investigated in future work.

### B Exact Bayesian model evidence

The model is given by

$$y_i = \sum_{n} w_n K_{in} + z_i / \sqrt{\tau}$$

(40)
where $K_{in} = K_n(x_i)$ is the basis function design matrix, $w_n$ are the weights, $\tau = 1/\sigma^2$ is the precision, and $z \sim N(0, 1)$. For the whole vector $y$ of $N$ responses, we can use the multivariate normal for the likelihood,

$$p(y | K, w, \tau) = \frac{\tau^{N/2}}{(2\pi)^{N/2}} \exp \left( -\frac{\tau}{2} (y - Kw)^T (y - Kw) \right), \quad (41)$$

where the precision matrix is diagonal with identical $\tau$ on the diagonal. Since the weight parameters $w_n$ enter quadratically, we can rewrite this expression in normal form for $w$,

$$p(y | K, w, \tau) = \frac{\tau^{N/2}}{(2\pi)^{N/2}} \exp \left( -\frac{\tau}{2} S \right) \times \exp \left( -\frac{\tau}{2} (w - \hat{w})^T K^T K (w - \hat{w}) \right) \quad (42)$$

with the residual sum of squares

$$S = (y - K\hat{w})^T (y - K\hat{w}) \quad (43)$$
$$= y^T y - \hat{w}^T K^T y \quad (44)$$
$$= y^T y - \hat{w}^T K(K^T K)^{-1} K^T y \quad (45)$$

The mixed terms cancel after plugging in $K^T y = K^T K \hat{w}$ from the known OLS solution $\hat{w} = (K^T K)^{-1} K^T y$. The above is of the form of a gamma distribution for $\tau$ multiplied with a normal distribution for $w$ conditioned on $\tau$. If we use a prior of the same form, we keep the form for the posterior, and thus have found the conjugate prior.

As a prior for the weights $w$, we choose

$$p(w | \mu, M) = \frac{\tau^{p/2} \sqrt{\det M}}{(2\pi)^{p/2}} \exp \left( -\frac{\tau}{2} (w - \mu)^T M (w - \mu) \right), \quad (46)$$

where $\tau M$ is the precision matrix with $\tau$ split off, and $\mu$ is the mean vector of the multivariate normal prior. Splitting off $\tau$ technical means that specifying $M$ is relative to the unknown $\tau$, but $\tau$ does not need to be known for that, as we integrate over all possible $\tau$ values.

For the posterior, we are interested in the quadratic form involving $w$,

$$(w - \hat{w})^T K^T K (w - \hat{w}) + (w - \mu)^T M (w - \mu) \quad (47)$$
$$= w^T A w - 2 w^T b + c \quad (48)$$

with

$$A = K^T K + M \quad (49)$$
$$b = K^T K \hat{w} + M\mu \quad (50)$$
$$c = \hat{w}^T K^T K \hat{w} + \mu^T M\mu \quad (51)$$
$$= \hat{w}^T K^T y + \mu^T M\mu$$
$$= y^T K(K^T K)^{-1} K^T y + \mu^T M\mu. \quad (52)$$
Put into this form, we can use the Gaussian integral \( \int d^n x \, e^{-\frac{1}{2} x^T A x + b^T x} = \sqrt{\frac{(2\pi)^n}{\det A}} e^{\frac{1}{2} b^T A^{-1} b} \) to marginalize,

\[
p(y \mid \tau) = p(y \mid K, \tau, \mu, M) \quad (53)
\]

\[
= \int d\theta \, p(y \mid K, \theta, \tau) \, p(\theta \mid \mu, M)
\]

\[
= \sqrt{\frac{\tau^{N+p} \det M}{(2\pi)^{N+p}}} e^{-\frac{1}{2} (S+c)} \int d\theta \, e^{-\frac{1}{2} (\theta^T A \theta - 2b^T \theta)}
\]

\[
= \sqrt{\frac{\tau^{N+p} \det M}{(2\pi)^{N+p}}} e^{-\frac{1}{2} (S+c)} \sqrt{\frac{(2\pi)^p}{\tau^p \det A}} e^{\frac{1}{2} b^T A^{-1} b}
\]

\[
= \sqrt{\frac{\tau^N \det M}{(2\pi)^N \det A}} e^{-\frac{1}{2} (y^T y + \mu^T M \mu - b^T A^{-1} b)}.
\]

For the \( \tau \)-integration, we choose the gamma distribution

\[
p(\tau | k, \vartheta) = \frac{1}{\Gamma(k) \vartheta^k} \tau^{k-1} \exp \left( -\frac{\tau}{\vartheta} \right)
\]

as the (conjugate) prior for \( \tau \), and define

\[
\xi = y^T \theta + \mu^T M \theta - b^T A^{-1} b.
\]

The remaining \( \tau \)-integral follows then from \( \int_0^\infty d\tau \, \tau^c e^{-\xi \tau} = c_1^{-c_0-1} \Gamma(c_0+1) \) as

\[
p(y) = p(y \mid K, \mu, M, k, \vartheta)
\]

\[
= \int_0^\infty d\tau \, p(\tau | k, \vartheta) \, p(y \mid \tau)
\]

\[
= \frac{1}{\Gamma(k) \vartheta^k (2\pi)^{N/2}} \sqrt{\frac{\det M}{\det A}} \int_0^\infty d\tau \, \tau^{N/2+k-1} e^{-\tau(\frac{c}{2} + \frac{1}{\vartheta})}
\]

\[
= \frac{\Gamma(N/2+k)}{\Gamma(k) \vartheta^k (2\pi)^{N/2}} \sqrt{\frac{\det M}{\det A}} \left( \frac{z + 1}{z + \vartheta} \right)^{-\frac{N}{2} - k}
\]

and for the log-evidence per data-point we obtain

\[
\frac{1}{N} \ln p(y) = \frac{1}{2N} \ln \frac{\det M}{\det A} - \frac{1}{2} \ln 2\pi - \frac{1}{2} \ln (z + 1)
\]

\[
\quad - \frac{k}{N} \ln \frac{z + 1}{\vartheta} - \frac{k}{N} \ln \vartheta
\]

\[
\quad + \frac{1}{N} \ln \Gamma \left( \frac{N}{2} + k \right) + \frac{1}{N} \ln \Gamma(k).
\]

C Details and illustration of less greedy stepwise regression

To our knowledge, all equation learning approaches apart from stepwise regression include an optimization step in various representation spaces of equations. Here, we propose a strategy that does without any numerical optimization algorithms, and instead considers candidate models individually in an almost comprehensive manner. Since already small dictionaries of basis functions can lead to tremendous numbers of candidate models, a combination of cheap selection criteria and successive reduction of model space with a suitable stopping criterion is required. We demonstrate how the simple criterion \( R^2 \) can be used for such a semi-comprehensive search.

In a first step, a dictionary of basis functions is generated using Algorithm 1. These basis functions consist of all possible products of available features \( x_j \). In these products, the factors are raised to all possible
Algorithm 1 Model ranking $R^2$

1: Function \textsc{DesignMatrix}(X)
2: \textbf{Input:} data $X$ with $N$ datapoints, $l$ features
3: \textbf{Parameters:} maximum degree $M_1$ for individual features, maximum degree $M_2$ for term
4: build all powers $x_{ij}^{m_j}$, $m_j = (1, \ldots, M_1)$ \hfill \triangleright \text{pre-computed for speed, limited to power } M_1
5: initialize counter $p = 0$
6: \textbf{for} all unique $l$-tuples $(m_1, \ldots, m_l)$ \textbf{do}
7: \hspace{1cm} if $\sum_j m_j \leq M_2$ then
8: \hspace{2cm} $p := p + 1$
9: \hspace{2cm} $K_{i,p} := \prod_j x_{:,j}^{m_j}$ \hfill \triangleright \text{Design matrix, candidate models in columns}
10: \hspace{1cm} \textbf{end if}
11: \textbf{end for}
12: \textbf{Return:} Design matrix $K$, shape $N \times p$

The less greedy (LG) strategy we propose is based on this basis function expansion and described by Algorithm 3. It begins by considering all regression models with $r$ non-zero weights $w_j$ (line 9), c.f. equation equation 12. To this end, the auxiliary Algorithm 2 produces a list of models with top $R^2$ values by looping through all candidate models of size $r$. These models are returned as an index matrix $M$, indicating selected terms with a 1 and deselected terms with a 0, where each column stands for a candidate model (lines 5,11). The models are sorted in descending order with respect to $R^2$ (line 13).

Back to Algorithm 3 we successively increase $r$ starting from $r = 1$ (lines 7,22). We found that for a fixed $r$ value, $R^2$ performs particularly well in identifying the best model out of the millions of models (see for instance the left plot in figure 5). To infer a value for $r$ with just $R^2$, we create a feature rating matrix $F$ defined as the weighted counts of terms being selected across $s$ top models, where the weight is given by $R^2$. Based on $F$, we check for terms selected by $R^2$ for two successive model sizes $r$ and $r-1$ (lines 16-20). If for both $r$ and $r-1$ the same terms are selected consistently, we choose these two terms as part of the inferred model and conclude the search.

As for larger values of $r$ the number of candidate models can easily reach hundreds of millions, we implement another strategy to divide out the list of candidate models. If terms have not been selected for two successive model sizes $r$ and $r-1$, we remove these terms from the design matrix $K$ (lines 14-15). In this way, we continuously reduce the model equation space as we go along.

The rationale for this selection and elimination strategy being solely based on $R^2$ is the following: If the true model has $r$ terms, and we are testing all models with $r-2$ terms, then the models with largest $R^2$ will consistently be composed of the $r-2$ terms that contribute the most to explaining the variance of $y$. The other $2$ true terms will be selected sporadically but at least once, terms that have not been selected at all can hence be removed from the candidate models. Testing in the next stage all models with $r-1$ terms, one more term will be consistently selected. The same holds for testing models with $r$ terms, but when testing models with $r+1$ terms, no new term can contribute consistently to explaining more of the variance of $y$. The $R^2$ measure will increase for models for $r+1$ terms, but compared to models with $r$ terms, no extra term will consistently be selected. Therefore, once no new term is selected consistently when incrementing the number $r$ of terms, we may conclude that all contributing terms have been found. An illustration of this strategy can be found in figure 5, where a typical case with three two terms in the true equation is shown.

Since the cheap computation of $R^2$ allows to go through millions of models in a matter of minutes on a standard computer, together with the described less greedy strategy, we are able to consider or exclude all candidate models that can be built from the basis function dictionary.
Algorithm 2 Model ranking $R^2$

1: Function TopRsq($y, K$)
2: Input: response data $y$, design matrix $K$
3: Parameters: number $r$ of terms and $t$ of top models
4: initialize criterion $c$ \hspace{1cm} $\triangleright$ flexible length, to store $R^2$ for candidate models
5: initialize model indices $M$ \hspace{1cm} $\triangleright$ $p$ rows indicating terms part of models, flexible number of columns
6: initialize model number $i = 0$
7: for all $n = (n_1, \ldots, n_p)$ with $n_j \in \{0, 1\}, \sum_j n_j = r$ do
8: \hspace{0.5cm} $i := i+1$, append $c_i$ and $M_{:,i} = 0$
9: \hspace{0.5cm} reduce $K_{\text{red}} := K_{:,n}$ \hspace{0.5cm} $\triangleright$ extract design matrix for selected terms
10: \hspace{0.5cm} determine $R^2$ for $K_{\text{red}}$ and $y$ from equation 22 \hspace{0.5cm} $\triangleright$ each column of $M$ indicates a model with $R^2$ value $c_i$
11: \hspace{0.5cm} store $c_i := R^2$ and $M_{:,i} := n$
12: end for
13: sort columns of $M$ and $c$ according to $c$ (descending) \hspace{0.5cm} $\triangleright$ first columns of $M$ now indicate top models in terms of $R^2$
14: Return: top models $M_{:,t}$ (shape $p \times t$), criterion $c_{:,t}$ (length $t$) \hspace{0.5cm} $\triangleright$ only return the $t$ best models

Figure 5: An example of how candidate models with 3, 4, and 5 terms with largest $R^2$ in each category tend to consistently choose the true terms of the model. The indices of all terms in the dictionary (the basis functions $k_n(x)$) are shown on the left vertical axis. Each candidate model along the horizontal axis is represented by squares indicating which terms make up the respective model. The ground truth model in this example has 3 terms, indicated by two lighter squares on the left and the horizontal lines. The case where the candidate model is the true model is indicated by a vertical line in the middle plot. The $R^2$ value in a logarithmic scale is shown as a line with closed circles, the values are given by the right vertical axis. It can be seen that the true model is chosen by $R^2$ among all models with 3 terms.
Figure 6: In the plot on the left, the $t$ best models in terms of $R^2$ from each of the $r$-sized candidate models in figure 5 have been combined and sorting according to their $R^2$ value. Clearly, the models with more terms are favored over models with fewer terms, and the true model is not selected. The plot on the rights shows the best models now in terms of the Bayesian model evidence $p(y|M)$, where now the true model is selected illustrating the adequate overfitting penalty held by $p(y|M)$.

Algorithm 3 Semi-comprehensive search with $R^2$

1: Function SC-Search($y$, $X$)
2: Input: data $y$, $X$
3: Parameters: maximum number $r_{\text{max}}$ of terms, number $s$ of top models and threshold $c_{\text{min}}$ for feature selection
4: $K, p := \text{DESIGNMATRIX}(X)$
5: initialize feature rating $F$ of shape $p \times r_{\text{max}}$  
6: initialize search := True
7: initialize number of terms $r := 1$
8: while search and $r \leq r_{\text{max}}$ do
9:   $M, c := \text{TopRsq}(y, K, r)$  
10:   append $M$ to $M_{\text{all}}$  
11:   $F_{\cdot r} := \sum_j c_{j} M_{\cdot j}$  
12:   normalize $F_{\cdot r} := F_{\cdot r} / \text{max}(F_{\cdot r})$  
13:   if $r \geq 2$ then
14:      index $i_0 := (F_{\cdot r} + F_{\cdot r-1} = 0)$  
15:      remove $K[, i_0]$ from $K$
16:      index $i_1 := (F_{\cdot r} \geq c_{\text{min}})$
17:      index $i_2 := (F_{\cdot r-1} \geq c_{\text{min}})$
18:      if $i_1 = i_2$ then
19:         search := False  
20:      end if
21:   end if
22:   $r := r + 1$
23: end while
24: Compute criteria \{$p(y|M)$, AIC, BIC, $R^2_{\text{adj}}$\} for $M_{\text{all}}$
25: Return: $M_{\text{all}}$ along with criteria
In a final step, the list of top models from the $R^2$ evaluation can be combined and each tested with other selection criteria like $p(y|M)$, AIC, BIC or $R^2_{\text{adj}}$. The best choice turned out to be the Bayesian model evidence $p(y|M)$ (c.f. [1]). In figure 6 the complexity selection power of $p(y|M)$ is demonstrated.

The hyperparameters of our procedure are the number $s$ of models used to count consistent selection of basis functions $k_n(x)$ together with the threshold $c_{\text{min}}$ the (normalized) count of a feature must exceed to be selected, the number $t$ of the best models in terms of $R^2$ that are combined in a new list of top models for re-evaluation with $p(y|M)$ (or other criteria), and the maximum number of iterations, $r_{\text{max}}$. We found that the universally best values are $s = p/2$, where $p$ is the feature dimension, $c_{\text{min}} = 0.75$, $t = 25$, and $r_{\text{max}} = 8$. 