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Published in:
I E E E Transactions on Signal Processing

Link to article, DOI:
10.1109/78.845935

Publication date:
2000

Document Version
Publisher's PDF, also known as Version of record

Link back to DTU Orbit

Citation (APA):

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Extraction of the Relevant Delays for Temporal Modeling

Cyril Goutte

Abstract—When modeling temporal processes, just like in pattern recognition, selecting the optimal number of inputs is a central concern. In this paper, we take advantage of specific features of temporal modeling to propose a novel method for extracting the inputs that attempts to yield the best predictive performance. The method relies on the use of estimators of generalization error to assess the predictive performance of the model. This technique is first applied to time series processing, where we perform a number of experiments on synthetic data, as well as a real life dataset, and compare the results to a benchmark physical method. Finally, the method is extended to system identification and illustrated by the estimation of a linear FIR filter on functional magnetic resonance imaging (fMRI) signals.

Index Terms—Delay estimation, functional magnetic resonance imaging, generalization error, identification, modeling, time series.

I. OVERVIEW

In this paper, we will be concerned with modeling the future behavior of a system based on its past plus, possibly, some exogenous control signal with application to time series prediction and system identification. Predictive performance depends, among other things, on the design of a proper input or regression vector. With too few inputs, the model does not have sufficient information and is unable to grasp the inner workings of the system, resulting in a large approximation error. On the other hand, a model with irrelevant inputs is overparameterized, which usually results in poor predictive performance, as suggested by the curse of dimensionality [1], [2].

We will focus mainly on time series prediction and later address the problem of system identification. In this context, potential inputs are past values, or delays, of the time series. Our aim is to select delays that are necessary to model the system while discarding unnecessary delays that could harm the overall performance. To our knowledge, the only provably optimal, general input selection method is exhaustive search, which is NP-complete and computationally unfeasible unless the number of inputs is very limited. Our method is related to iterative feature selection techniques used in traditional statistics [3]. It builds on the specificities of temporal processing to provide an original way of selecting potential inputs. The relevance of a candidate delay is assessed directly by its effect on predictive performance that is measured using estimators of generalization error.

The following sections are organized as follows. First, we give a general presentation of feature selection applied to time series modeling from the statistical and physical points of view (Section II). We suggest the use of estimators of generalization error to evaluate the quality of a subset of features. Our extraction of the relevant delays (ERD) method is described in Section III as a principled alternative. The second part of the paper contains a number of experiments conducted on three different datasets. Time series predictions (Section IV) are addressed using the well-known artificial Hénon map and a real-world time series measuring the mean monthly flow of the Fraser river; ERD is then applied to system identification (Section V) using a functional magnetic resonance imaging (fMRI) dataset. We conclude with a discussion of the method and results.

II. FEATURE SELECTION

Let us consider a standard time-series modeling problem; a sequence of measurements \( y(t), 1 \leq t \leq T \) is collected. We wish to predict \( y(t) \) from a set of past values \( y(t-d), d > 0 \).

Extracting the relevant delays consists of finding the set of \( \eta \) delays \( (d_1, d_2, \ldots, d_m) \) such that the input vector

\[
x(t) = [y(t-d_1), \ldots, y(t-d_m)]
\]

(1)
yields the best prediction of \( y(t) \).

A. Physical Approach

The physical approach relies on estimating the embedding dimension of the time series [4]. This is essentially equivalent to finding the set of primary delays, i.e., delays with an explicit influence on the observed values. For example, for a time series generated by \( x(t) = g(x(t-1)) \), \( 1 \) is the only primary delay. As \( x(t) = g(x(t-2)) \), we can also estimate \( x(t) \) using \( x(t-2) \); however, \( 2 \) is a secondary (not a primary) delay.

Several methods have been proposed to estimate the embedding dimension and the embedding space of a time series. Pi and Peterson [5] have introduced the “\( \delta \)-test” in the neural networks literature. He and Asada [6] proposed the use of “Lipschitz quotients” to identify the order of nonlinear input-output systems. An independent but essentially similar method was applied to time series in the signal processing literature [7]. These methods all rely on the assumption that the underlying mapping \( g \) is continuous and reasonably smooth. The existence of a

1 Note that we do not address the problem of obtaining the best sampling frequency, i.e., length of the basic time delay (difference between \( t+1 \) and \( t \)).
continuous mapping between $\mathbf{x}(t)$ and $y(t)$ means that close inputs $\mathbf{x}(u)$ and $\mathbf{x}(v)$ are mapped to close outputs $y(u)$ and $y(v)$. On the other hand, for insufficient input spaces (i.e., missing delays), close inputs can correspond to arbitrarily distant outputs (see Section IV for a practical example). Quantifying this closeness is done either by estimating the probability that two outputs are close given close inputs ($\delta$-test) or by calculating the ratio between output and input distances (Lipschitz quotients).

An important side effect is that all input–output pairs have to be considered, requiring extensive calculation.

Note that the techniques mentioned above are nonparametric, rely on the data alone, and need not specify a given model. This can turn out to be disadvantageous since for a given data set, only one set of relevant delays is selected, regardless of the ability of the model to actually implement the underlying mapping using them. Flexible models such as neural networks should overcome this limitation by their uniform approximation capabilities [8]. However, we will see that in practice, a given model often benefits from the inclusion of secondary delays (cf. Section IV).

### B. Statistical Approach

From a statistical point of view, the extraction of the relevant delays is a special case of feature selection, which is itself a part of the more general problem of analyzing the structure in the data [3]. The statistical approach relies on specifying a (parametric or nonparametric) model $f$ with which we try to estimate the input–output mapping $y(t) = f(\mathbf{x}(t))$. In conventional feature selection, an important assumption is the availability of all necessary variables. Provided that our data are sampled correctly, this assumption is usually satisfied in the case of time series. In the following, delays are positive integers $d_j > 0$, $1 \leq j \leq m$, variables or features are past values of the time series $y(t-d_j)$, and inputs are the vectors $\mathbf{x}(t)$ containing these past values. Finding the relevant delays is then equivalent to finding the input that gives optimal predictive performance. Conventional feature selection relies on three different components:

1) a selection method searching through possible subsets of variables;
2) an evaluation criterion assessing the quality of each subset;
3) a stop condition, which decides whether a satisfactory subset was obtained.

To our knowledge, the only general optimal method for selecting the best features among $F$ is to perform an exhaustive search through all $2^F - 1$ possible subsets of variables. This optimal approach becomes unfeasible for moderate values of $F$. Furthermore, in temporal modeling, the maximum delay and, thus, the total number $F$ of variables, is not known beforehand, but we would typically accept to probe quite far into the past. For monotonous evaluation criteria, the branch and bound algorithm [9] provides an efficient alternative. Unfortunately, the predictive ability is not a monotonous criterion. Common suboptimal alternatives perform an iterative search by regularly increasing or decreasing the number of selected features [3].

**Forward Selection:** Starting from an empty set (no variables), add variables according to the evaluation criterion until the stop condition is reached.

**Backward Elimination:** Starting with a full set (all possible variables), delete one variable at a time according to the evaluation criterion until the stop condition is reached.

**Stepwise Regression:** Alternate between both methods, by, e.g., performing a backward elimination after each inclusion or choosing between adding or deleting variables according to the evolution of the evaluation criterion.

Note that the focus in neural networks research, for example, is almost entirely on backward elimination through various pruning schemes [10]. All these suboptimal methods rely heavily on the evaluation criterion. Typical choices include, for example, the $F$ statistic [3], extensions to nonlinear models, or mutual information [11]. We will now present an alternative to these choices.

### C. Generalization Approach

In the context of nonparametric modeling, our goal is to obtain the best prediction. We will thus use a measure of the predictive abilities as our evaluation criterion. For a model $f$ mapping an input vector $\mathbf{x}(t) = [y(t-d_1) \cdots y(t-d_m)]$ to output $y(t)$, the risk $L(y, f(\mathbf{x}))$ measures the cost associated with estimating the output $y$ observed in $\mathbf{x}$ with the model $f(\mathbf{x})$. Assuming Gaussian noise on the output usually leads to the choice of the quadratic risk, $L(y, f(\mathbf{x})) = (y-f(\mathbf{x}))^2$. The generalization error (or expected risk) associated with model $f$ is defined as the expectation of the risk over the unknown, but fixed, joint input–output distribution

$$G(f) = \int (y-f(\mathbf{x}))^2 p(\mathbf{x}, y) \, d\mathbf{x} \, dy.$$  

This is also known as the prediction error or the integrated squared error. Ideally, the evaluation criterion for a given subset should be the generalization error of the model using this subset as input [12]. However, (2) cannot be used directly because the joint input–output probability is unknown. It will be estimated from the available data $(\mathbf{x}(t), y(t))$, with $t \in \mathcal{D} = \{T_{\text{min}}, \cdots, T_{\text{max}}\}$.

The split-sample (SS) estimator is obtained by replacing the joint distribution $p(\mathbf{x}, y)$ by its empirical estimator on the validation set $\mathcal{V} \subset \mathcal{D}$

$$\hat{G}_{\text{SS}} = \frac{1}{|\mathcal{V}|} \sum_{t \in \mathcal{V}} (y(t) - f^{-\mathcal{V}}(\mathbf{x}(t)))^2$$  

where $|\mathcal{V}|$ is the cardinality of $\mathcal{V}$, and $f^{-\mathcal{V}}$ is the model estimated on the training set $\mathcal{T} = \mathcal{D}\backslash\mathcal{V}$. Note that $\mathcal{T}$ and $\mathcal{V}$ must be disjoint in order for $\hat{G}_{\text{SS}}$ to be an unbiased estimate of $G(f^{-\mathcal{V}})$.

The cross-validation estimator (CV) estimator [13], [14] resamples the validation and training sets from the available data in order

3Where $T = \mathcal{D}\backslash\mathcal{V}$ means $T = \{t \in \mathcal{D} \mid t \notin \mathcal{V}\}$.

4The split-sample method is sometimes referred to as “cross-validation” in the neural networks literature. This is inconsistent with the definition of [13], and we will here reserve the term exclusively for the averaging method (4).
to increase the reliability of the resulting estimator. In $L$-fold CV, $\mathcal{D}$ is split into $L$ disjoint subsets $(\mathcal{S}_j)_{j=1, \ldots, L}$ of roughly equivalent size $\bigcup_{j=1}^L \mathcal{S}_j = \mathcal{D}$. The split-sample estimators (3) calculated using each $\mathcal{S}_j$ in turn as a validation set are averaged over the subsets

\[
\hat{G}_{\text{CV}} = \frac{1}{N} \sum_{j=1}^L \sum_{t \in \mathcal{S}_j} (y(t) - f^{-\mathcal{S}_j}(\mathbf{x}(t)))^2
\]

\[
= \frac{1}{N} \sum_{t \in \mathcal{D}} (y(t) - f^{-t}(\mathbf{x}(t)))^2
\]

(4)

where for notational convenience, we introduce $f^{-t}$, which is the model estimated excluding the subset $\mathcal{S}_j$ containing $t$. This means that $\forall j, \forall (u,v) \in (\mathcal{S}_j)^2$, $f^{-u} = f^{-v}$. Note that (3) does not estimate the generalization error (2), but it does estimate its average over all possible datasets of the same size sampled from $p(\mathbf{x},y)$.

Finally, a number of analytical asymptotic estimators of the average generalization error have been proposed in the literature, e.g., final prediction error (FPE) [15], generalized prediction error (GPE) [16], final prediction error for regularized problems (FPER) [17], or network information criterion (NIC) [18]. Without loss of generality, we will settle for a GPE-like expression, i.e., an FPE with an effective number of parameters $\hat{P}$

\[
\hat{G}_{\text{FPE}} = \left( \frac{N + \hat{P}}{N - \hat{P}} \right) \hat{R}(f)
\]

(5)

where $\hat{R}(f)$ is the training error (or empirical risk) on dataset $\mathcal{D}$: $\hat{R}(f) = \sum_{t \in \mathcal{D}} (y(t) - f(\mathbf{x}(t)))^2$. For unregularized linear models, $\hat{P} = \hat{P}$, which is the number of parameters. For regularized and/or nonlinear models, we generally have $\hat{P} < \hat{P}$, and the exact expression for $\hat{P}$ depends on the estimator and the regularization method (see, e.g., [17] and [19]).

Estimators of generalization error, whether they are based on cross-validation or asymptotics, will now be used as evaluation criterion in order to derive an original delay extraction scheme.

III. EXTRACTION OF THE RELEVANT DELAYS

The above presentation of statistical feature selection applies to general regression problems. Time series prediction, and, to some extent, system identification, have a number of distinct characteristics. On the one hand, all potential features are available, but there is no upper bound on the maximum delay. This makes the extraction of relevant delays a rather bad candidate for backward elimination schemes. On the other hand, the chronological order yields a natural ordering of variables that our method uses as a natural selection criterion. The rationale for this scheme is that primary delays will always be tested for inclusion before secondary delays. As a consequence, secondary delays will never be included unless 1) the model is unable to represent the underlying mapping using primary delays alone, or 2) the secondary delay is also a primary delay.\(^6\)

\(^6\)Other estimators lead to similar expressions, and their subsequent use is straightforward.

The ERD method takes all delays in their natural order and adds a candidate variable if and only if it yields a significant decrease in generalization error. The algorithm can be described as follows.

1) Initialize: $d = 0$; no input selected; $G_{\min} = \sigma_y^2$ (time series variance).
2) Model: $d = d + 1$; add delay $t - d$ to selected inputs; Estimate generalization error $\hat{G}$ for resulting model.
3) Test: If $\hat{G}$ is significantly smaller than $G_{\min}$, then keep delay $t - d$ and cut $G_{\min} = \hat{G}$; else, discard delay $t - d$.
4) Iterate: Go to step 2 until stop condition is reached.

A. Optimality and Suboptimality

In the ideal case of a complete model (i.e., contains the target mapping) and sufficient amount of noise-free data (i.e., fairly larger than the number of parameters), the above method is optimal. To see this, consider the maximum primary delay $d_{\text{MAX}}$. Clearly, all models obtained for $d < d_{\text{MAX}}$ have strictly positive generalization errors $\hat{G} > 0$. Due to the completeness of the model class, there is a model using delays up to $d_{\text{MAX}}$, which yields $\hat{G} = 0$, and due to the noise-free assumption, this model can be estimated. As models using $d > d_{\text{MAX}}$ cannot yield any decrease in generalization error, no further delay will be selected.

For more general situations, we are not aware of any proof that the model is optimal. Indeed, the experiments presented below suggest that it is not, although results appear to be close to optimum. The correctness and near optimality of the extracted delays will depend on several factors, the approximation capabilities of the model and the noise level among them.

Note that the traditional caveat against sequential selection is its inability to handle variables that are combinations of other variables. This arises naturally in temporal modeling as each observation is a (possibly nonlinear) mapping of previous values. However, chronological selection ensures that each variable is tested for inclusion before the variables on which it depends, guaranteeing a parsimonious selection.

B. Significant Decrease in Error

Step 3 of the above algorithm requires that we assess the significance of an observed decrease in generalization error. This requirement avoids the inclusion of a delay leading to negligible decrease in estimated generalization error, which could happen by chance alone. We take advantage of the fact that the estimators presented in (3)–(5) are based on averaging over the data. The CV and asymptotic estimators can be put in the general expression

\[
\hat{G} = \frac{1}{N} \sum_{t \in \mathcal{D}} \epsilon(t)
\]

(6)

where $\epsilon(t) = (y(t) - f^{-t}(\mathbf{x}(t)))^2$ for CV and $\epsilon(t) = ((N + \hat{P})/(N - \hat{P})(y(t) - f(\mathbf{x}(t)))^2$ for GPE [16].\(^7\) For the split-sample estimator (3), $\epsilon(t) = (y(t) - f(\mathbf{x}(t)))^2$, and the average runs over $\mathcal{V}$ instead of $\mathcal{D}$.

\(^7\)Other estimators like FPER or NIC will lead to similar forms with different expressions for the penalty term of $\hat{P}$.
We wish to test whether $G_m$, which is the generalization error for the model with $m$ delays (current best in the algorithm), is significantly different from $G_{m+1}$, which is the generalization error for the model with $m+1$ delays (augmented with the candidate delay), using the residuals $e_m(t)$ and $e_{m+1}(t)$ and their average $\bar{G}_m$ and $\bar{G}_{m+1}$, which are unbiased estimators of $G_m$ and $G_{m+1}$, respectively. Assuming that the difference between the residuals $e_m(t)$ and $e_{m+1}(t)$ is approximately normally distributed and using the fact that all residuals are calculated on the same data, we assess the significance of an observed decrease in generalization error by using a one-tailed paired $t$ test [20] between the residuals. Note that by construction, the candidate model has an additional delay, meaning that fewer input–output pairs $(x(t), y(t))$ can be formed. In order to apply the paired $t$-test, we will discard the additional data points used by the smaller model. This waste of a couple of examples is counter-balanced by the superior power of the paired $t$-test, compared with the nonpaired version.

The choice of the $t$-test is typical for assessing the significance of a difference in means when we are willing to make a normal assumption about the individual differences $(e_m(t) - e_{m+1}(t))$. An efficient nonparametric alternative is the Wilcoxon matched-pairs signed-ranks test [21], which has an asymptotic relative efficiency (compared with the $t$-test) of 0.95 in the normal case. This means that the price for relaxing some of the parametric assumptions in the $t$-test is that it needs asymptotically only 5% more data to assess a given difference with the same significance.

As in standard iterative subset selection, the significance level $\alpha$ has an influence on the result. When $\alpha \to 0$, no delays are selected. However, contrary to the standard case, $\alpha \to 1$ does not necessarily select all delays. This is because the estimators of generalization take the increase in model complexity into account and do not always decrease for larger models (this is also observed with, e.g., Mallow’s $C_p$ [22]). Note that it has been reported that the traditional choice of $\alpha = 0.05$ (the 5% significance level) tends to be overly conservative. In agreement with standard practice in subset selection [23], we will choose $0.15 < \alpha < 0.25$, a range in which experimental results seem to be stable. Finally, an additional level of generalization estimation can be invoked to tune the value of $\alpha$ (e.g., CV in [24]), but the overall process becomes cumbersome and has not been pursued here.

C. Stop Condition

The stop condition is motivated by practical or by problem-specific considerations. As mentioned above, models with larger delays in the input will have less input–output pairs available; for a sequence of $T$ measurements and inputs with maximum delay $d_m$ (1), at most $(T - d_m)$ training examples can be formed. This means that with increasing delays, 1) the estimation becomes more difficult (less data for more parameters), and 2) the degrees of freedom in the statistical test decreases, such that the difference in estimated generalization error must be larger in order to become significant. The available data provides a natural upper bound for the maximum possible delay; the selection is stopped when the data becomes insufficient for proper estimation. Most parametric models, such as linear models or neural networks, require at least as many examples as parameters. Statistical rules-of-thumb suggest a ratio of at least 10 examples per parameter. For the Hénon map example below (Section IV-A) and a linear model, the maximum possible delay becomes 500 (data points) minus six (parameters in the candidate model), i.e., $d_{\text{max}} = 494$ (and $d_{\text{max}} = 440$ if one requires 10 data per parameter).

On some problems, and typically when modeling a physical system, it may make sense to introduce stronger constraints to reflect additional knowledge on the phenomenon or requirements on the model. In the fMRI experiments presented below (Section V), the resulting filter tentatively models the haemodynamic response to neuronal activation. As the experiment is performed as a series of consecutive runs and we are interested in the response to a typical activation pattern, it is sensible to limit the delay extraction to one run, i.e., 48 delays in that case.

Because the selection criterion is independent of the stop criterion, the influence of the latter is in how conservative the resulting model will be. Any criterion that stops the algorithm earlier will necessarily yield a model with fewer (or as many) delays selected, i.e., it will be more conservative. It is, thus, fair to say that the maximum delay gives the least conservative model (everything being otherwise equal). Adding stronger (e.g., physical) constraints on the maximum delay will potentially trade a more conservative model for an increase in interpretability. Note finally that due to the decreasing degrees of freedom when less data are available, the last included delay is usually (in the examples we have processed) much smaller than $d_{\text{max}}$.

IV. TIME SERIES PREDICTION

A. Hénon Map

Let us first consider a time series generated by the well-known Hénon map [25]

$$y(t) = 1 - 1.4(y(t - 1))^2 + 0.3y(t - 2), \quad (7)$$

We apply several methods on a dataset containing 500 points. An independent set of 10 000 elements is sampled from (7) to assess the resulting generalization abilities. The noisy map, with additive Gaussian noise ($\sigma^2 = 0.1$) is also investigated. We consider two models:

- linear model (obviously a bad choice to model the non-linear Hénon map) using FPE [15] as an estimator of generalization error;
- kernel smoother [26], [27] using the leave-one-out (i.e., $N$-fold) CV estimator;
and four delay selection methods

1) the $\delta$-test [5], estimating the embedding dimension;
2) the ERD method (Section III);
3) a “divine guidance” or large validation set (LVS) method;
4) the traditional $F$-inclusion scheme.

The “divine guidance” method selects delays on the basis of a large independent dataset providing a reliable estimator of generalization. This method is, of course, impractical on real data and is used in order to check the behavior of the estimators used in ERD.

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In $F$-inclusion, a candidate delay is included iff

$$F = \left( \frac{R_m(f) - R_{m+1}(f)}{NR_{m+1}(f)/(N-m-1)} \right) > F_{\alpha}$$

(8)

where $R_m(f)$ and $R_{m+1}(f)$ are the empirical risks, or training errors, for the models with $m$ delays and $m+1$ delays (respectively), calculated on the same data, and $F_\alpha$ is the $F$-distribution threshold for an $\alpha$ confidence level 1 and $(N-m-1)$ degrees of freedom. A candidate delay is therefore included when it yields a significant improvement in observed performance.

Figs. 1 and 2 show a plot proposed by Aleksic [28] to investigate the embedding dimension. All points correspond to a pair of data $(t_1, t_2)$ with $|\mathbf{x}(t_1) - \mathbf{x}(t_2)|$ small and give the output distance $|\mathbf{y}(t_1) - \mathbf{y}(t_2)|$ against $|\mathbf{y}(t_1) - \mathbf{y}(t_2)|$. Clearly, with only one delay in the input (Fig. 1), close inputs do not guarantee close outputs. Therefore, there is no continuous mapping from $\mathbf{x}(t) = \mathbf{y}(t-1)$ to $\mathbf{y}(t)$. On the other hand, with the first two delays (Fig. 2), all pairs with close inputs also have close outputs. This suggests that a continuous mapping between the first two delays and the time series value can be implemented. Accordingly, the $\delta$-test selects the first two delays from the dataset [5].

For noise-free data and the kernel smoother model, all other selection methods also select these two delays and are able to implement the Hénon map perfectly from the 500 available observations. In the remaining situations, they always select at least one additional delay, depending on the model-noise combination (see Table I).

All resulting models are tested on the large noise-free test set in order to check their generalization abilities (Fig. 3). Predictably, the kernel smoother provides much better performance than the linear model. Note that the $\delta$-test outperforms a statistical method ($F$-inclusion) only once: for the “kernel-noisy data” combination. This is due to its emphasis on extracting the primary delays; as efficient as the $\delta$-test may be at extracting the “true” delays, it suffers from not addressing the relevant goal in our context, namely, prediction of future time series values. The relative edge of the $\delta$-test for that particular model-noise combination is rather due to the inferior performance of $F$-inclusion.
This suggests that $F$-inclusion is ill-suited to flexible nonlinear models. This is because (8) does not penalize overfitting; on the contrary, when $R(f)$ becomes overly small, the $F$ statistic will be artificially increased, leading to an exceedingly optimistic selection (15 delays, instead of two or three). Notice the close agreement between L VS and ERD, suggesting that the estimators of generalization error (FPE and CV) perform well. ERD tends to be more parsimonious, yielding five and three delays for the linear model, instead of seven and seven, for similar performance. Finally, on noisy data, even with the flexible kernel smoother, ERD and L VS select an additional, secondary delay. This is because we try to learn the underlying relationship from a limited amount of data. This additional delay, although not theoretically necessary, yields a 30% improvement in performance.

B. Fraser River

Let us now turn our attention to a real dataset, which is analyzed in [29] and publicly available. The data records mean monthly measurements of the flow of the Fraser river in Hope, B.C., Canada, from March 1913 to December 1990. It contains 946 values with a rough periodicity and maxima every 11–13 months (see Fig. 4). The dataset is split in half: The first 473 observations are used for model estimation and delay extraction, and the last 473 are used for testing the generalization abilities of the resulting model. In the following experiments, the data have been log-transformed and centered. The selection method using the large validation set is, of course, not sensible in this context, as no extra data is available. Accordingly, we will consider only two selection schemes (the $\delta$-test and the proposed ERD) applied to three models:

- linear model;
- kernel smoother;
- nonlinear neural network model (multilayer perceptron with one hidden layer).

The parameters of the linear and neural network models are estimated by minimizing the mean squared error on the transformed data. The estimators of generalization error are the FPE [15] and GPE [16], respectively, for these models and the leave-one-out estimator, as above, for the kernel smoother. The nonparametric $\delta$-test selects six delays, whereas the ERD extracts six to 12 delays, depending on the model (see Table II). Not surprisingly, the more flexible models, i.e., kernel smoother and neural network, use fewer delays, whereas the linear model uses twice as many.

Notice that although the $\delta$-test and ERD extract the same number of delays for two out of three models, these delays do not coincide. Typically, ERD probes further into the past: across two periods for the neural network and even four for the linear model. This is an interesting artifact of the $\delta$-test method; as the size of the input space grows larger, fewer and fewer pairs of points have close inputs. As a result, the variance of the estimated probabilities increases, and the test is unable to reliably select additional delays.

The generalization abilities of the resulting models are assessed using the test set. On Fig. 5, points far above the dotted line indicate probable overfitting. Not surprisingly, the kernel smoother overfits the data quite severely. This model is known to suffer acutely from the curse of dimensionality and has difficulties handling high-dimensional inputs when only one smoothing parameter is used [30]. Surprisingly, the best overall performance is achieved by the combination of ERD and linear model. The flexible neural network model does marginally worse and seems to overfit slightly. Note that with a good training algorithm, the neural network should be able to outperform the linear model using the same inputs. However, the ERD scheme has

<table>
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<td>Fraser river</td>
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<tr>
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<tr>
<td>Kernel</td>
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Fig. 4. Mean monthly flow of the Fraser river recorded in Hope, B.C., Canada, between January 1925 and January 1945 (solid line). The one- and ten-year moving averages are shown as dotted and dashed lines, respectively.

Fig. 5. Result for three different models of both extraction schemes: the proposed extraction of the relevant delays (ERD, full marks) and the $\delta$-test (empty marks). The three models are linear, kernel smoother, and nonlinear neural network (NN).
predictably extracted different sets of delays for both models. In particular, the linear model uses three additional long-term delays (26, 35, and 48).

Again, these results illustrate the limitations of the physical approach when prediction performance is the target. The statistical approach based on model performance gathers more information in the past and yields an improvement in performance that is, here, most impressive for the linear model.

V. SYSTEM IDENTIFICATION

The ERD scheme can be extended to some simple system identification problems, such as the identification of linear or nonlinear finite input response (FIR) filters [31]. The data is acquired during a functional magnetic resonance imaging (fMRI) experiment. While the subject lies in the scanner, he is asked to either lie motionless (rest) or perform a simple motor task (activation), namely, left-handed finger-to-thumb opposition. Two 64 × 64 slices of whole brain echo-planar fMRI images are acquired every 2.5 s. The dimension of each voxel is 3.1 × 3.1 × 8 mm. Nine runs of 48 images are acquired in the following sequence: 12 images (30 s) of rest, 24 images (60 s) during activation, and 12 images of rest again. The complete time series consists of 432 measurements (18 min) in each voxel. In the following experiments, we focus on the 39 × 49 area containing the brain and only one slice.

The fMRI signal measures the haemodynamic response to focal neuronal activation. It is widely believed that the response in the brain can be characterized as a convolution of the reference stimulus signal representing the activation with a linear filter. In our experiment, the reference stimulus \( h(t) \) is a square wave with positive values for the scans acquired while the subject was performing the motor task and negative values during rest (Fig. 6). Previous approaches tried to characterize the haemodynamic response using several convolution filters: Poisson [32], Gamma [33], or FIR filter with a fixed length [34]. We will try to extend the latter to general FIR filters by extracting the delays that yield the best modeling performance. In other words, we are looking for the set of delays \( d_1, \ldots, d_m \) such that
\[
\hat{y}(t) = \sum_{i=1}^{m} \beta_i h(t - d_i) \tag{9}
\]

The performance of the resulting filters is compared with a FIR filter of fixed size. As the average haemodynamic delay has been observed between 5–10 s [35], i.e., two to four images, we will use an FIR filter with seven delays as reference. In Fig. 7, points above the dotted diagonal mean higher test error, i.e.,
ERD improves on traditional statistical selection methods by focusing on the relevant performance criterion, namely, the generalization error. An interesting side effect of the estimators of generalization error is that they naturally take into account the limited size of the available dataset. A similar effect is known from the AIC [36] and BIC [37] information criteria: BIC selects the asymptotically optimal model size, which AIC tends to overestimate, while effectively minimizing the average generalization error for finite datasets. Similarly, ERD selects additional delays that help prediction. In the Hénon map example, the selection of $d = 3$ yields better performance according to both the large validation set method and the independent test set. It should also be noted that it is the minimum generalization, rather than its value, that matters, such that the method could very well accommodate an estimator with a consistent bias.

The ERD method is essentially a forward selection method (see Section II-B). It is straightforward to extend it to backward elimination or stepwise regression using the same evaluation criterion. As we argued in Section III, backward elimination is ill suited to temporal modeling. However, stepwise regression can be applied by alternating forward and backward steps. With many models, this can be addressed during the model estimation, e.g., using an adaptive metric approach for kernel smoothing [30]. Gaussian processes [38], or pruning and regularization for linear models or neural networks [19], [39], [40]. These methods automatically discard irrelevant variables in the input vector. An additional interesting prospect for future research is the extension of the ERD scheme to more general system identification architecture, e.g., auto-regressive filter with exogenous input (ARX). Note that, in that case, the filter input consists of the system output $y(t)$ and the control signal $u(t)$, and the precise ordering of the candidate inputs has to be determined.

As a conclusion, we have presented a generalization-based method for the extraction of the relevant delays in temporal modeling with a focus on time series prediction and system identification. This method is related to traditional forward selection methods and is therefore well founded. It is easy to implement and requires little time in addition to model estimation. Furthermore, it accommodates efficiently the tradeoff between number of selected delays (model size) and model flexibility (approximation error). The method provides interesting results on a number of experiments performed in different contexts.

**ACKNOWLEDGMENT**

The Fraser river dataset (Section IV-B) is publicly available on statlib: http://lib.stat.cmu.edu/datasets/. The fMRI dataset (Section V) was acquired by R. Savoy at Massachusetts General Hospital, Boston. The author would like to thank P. Gallinari, J. Larsen, and the learning group at IMM, DTU, for valuable discussions on previous versions of this work.

**REFERENCES**

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