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A NEURAL ARCHITECTURE FOR
NONLINEAR ADAPTIVE FILTERING OF
TIME SERIES

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INTRODUCTION

The need for nonlinear adaptive filtering may arise in different types of filtering tasks such as prediction, system identification and inverse modeling [17]. The problem of predicting chaotic time series has been addressed by several authors [6],[11]. In the latter case a feed-forward neural network was used both for prediction and for identification of a simple, nonlinear transfer function (system identification). These filtering tasks can be solved using a filtering configuration shown in Fig. 1 [17].

\[ y(k) = F[y(k-1), \ldots, y(k-M), x(k), \ldots, x(k-L+1); \theta] \]  

(1)

where \( F[\cdot] \) is an unknown nonlinear function parameterized by \( \theta \), \( k \) is the discrete time index and \( L, M \) are filter orders.
The general structure of equation (1) enables one to model any nonlinear discrete system. $\theta$ is assumed to be slowly time varying and consequently $y(k)$ is quasi stationary. The use of a recursive, nonlinear filter does, however, pose serious difficulties regarding stability. The filter may display limit cycles, chaotic behaviour and unboundedness. The scope of this paper is to implement only the nonrecursive part of (1).

We propose a modularized architecture for the nonlinear filter in Fig. 1 including algorithms for adapting the filter. Further we develop simple guidelines for selecting a specific filter design within the proposed architecture given a priori knowledge of the distribution and origin (type of modeling problem) of the input signal $x(k)$ and the desired response $d(k)$. Finally we present simulations in order to further investigate the nature of the relations between filter design and the statistics of $x$ and $d$.

**NONLINEAR FILTER ARCHITECTURE**

The proposed filter architecture is shown in Fig. 2. The filter, which may be viewed as a generalization of the Wiener Model [15], is divided into three partially independent (depending on specific design) sections: A preprocessing unit containing the filter memory, a memoryless, multidimensional nonlinearity (MMNL) and a linear combiner. The structure is selected in order to modularize the modeling problem which ensures a proper and sparse parameterization, and facilitates incorporation of a priori knowledge in contrast to the limited possibilities when using an ordinary feed-forward neural network. The main objective of the preprocessor is to extract the essential information contained in $x_k = [x(k), x(k-1), \ldots, x(k-L+1)]'$ ensuring that $z$ has a dimension, $p \leq L$. The nonlinearity is memoryless and transforms the vector $z$ into the vector $v$, and finally the linear combiner forms a weighted sum $y(k)$ of the terms in $v$. This corresponds to rewriting (1) as:

$$y(k) = a'F_n[F_p(x(k))]$$ (2)
where \( F_p(\cdot) \) is the preprocessor, \( F_w(\cdot) \) is the MMNL and \( a \) the weights of the linear combiner. The filter could be viewed as a heterogeneous multi-layer neural network. All sections could be adapted, but this is not always necessary, as we shall see in the next section.

**FILTER DESIGN**

**Signal Dependence**

It seems reasonable that the specific design of the filter depends on the origin and distribution of the signals \( z(k) \) and \( d(k) \), and we will summarize some guidelines for choosing an appropriate design as follows:

<table>
<thead>
<tr>
<th>Case</th>
<th>( x )</th>
<th>( d )</th>
<th>Model</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Gaussian</td>
<td>Gaussian</td>
<td>( d(k) = a'x_k + \epsilon(k) )</td>
</tr>
<tr>
<td>2</td>
<td>Gaussian</td>
<td>Non-Gaussian</td>
<td>( d(k) = F[x_k] + \epsilon(k) )</td>
</tr>
<tr>
<td>3</td>
<td>Non-Gaussian</td>
<td>Gaussian</td>
<td>( d(k) = F[x_k] + \epsilon(k) )</td>
</tr>
<tr>
<td>4</td>
<td>Non-Gaussian</td>
<td>Non-Gaussian</td>
<td>( d(k) = F[x_k] + \epsilon(k) )</td>
</tr>
</tbody>
</table>

1. The linear filter is optimal in this case, so \( v = [1 \ x']' \), see e.g. [13, Theorem 14.3].
2. The Wiener model, i.e. a bank of orthogonal linear filters in the preprocessing unit followed by a fixed (non-adaptive) polynomial nonlinearity, provides a filter which can be adapted in a very simple and fast way [15]. The linear filters may be determined using principal component analysis (PCA) on \( x \). This case is associated with the problem of nonlinear system identification where \( d(k) \) is the output of an unknown system and \( z(k) \) the input.
3. In this case there is no obvious choice of filter design. The case arises e.g. in inverse modeling where \( d(k) \) is the driving signal of the unknown system and \( z(k) \) the resulting output.
4. This case relates to prediction of nonlinear time series where \( z(k) = d(k-\tau) \) is, in fact, a delayed version of \( d(k) \). Previous simulation studies [11] indicate that the nonlinearity should be constructed from bounded functions (e.g. the commonly used \( \tanh(\cdot) \)) rather than polynomials, which have the inconvenient property of growing fast towards infinity.

**Preprocessing Methods**

We present two possible methods for dimensionality determination. If the unknown system being modeled can be described in the form of a nonlinear differential equation it is possible, in some cases, to determine dimensionality by letting \( z = [x(k), Dx(k), \ldots, D^{p-1}x(k)]' \) where \( D \) is a discrete derivative operator. We denote this preprocessor: The derivative preprocessor (DPP).
The following example (the pendulum) illustrates that we often have \( p < L \) without losing information:

\[
\frac{d^2 x(t)}{dt^2} + \beta \frac{dx(t)}{dt} + \alpha \sin[x(t)] = d(t)
\]

(3)

\[
D^2 z(k) + \beta Dz(k) + \alpha \sin[z(k)] = F[D^2 z(k), Dz(k), z(k)] = d(k)
\]

(4)

Equation (4) which is a discrete approximation of (3) clearly shows, that \( d(k) \) may be expressed as a function of only \( p = 3 \) variables i.e. the derivatives of \( z(k) \). To ensure that the approximation of the derivative operator \( D \) is accurate (i.e. approximates the continuous time derivative operator) over a broad range of frequencies it must be implemented using a linear filter with high order. A tapped delay line used without a preprocessing element would necessarily need the same length \( L \) to hold the same information so \( L \) is obviously greater than \( p \) in this example. In practice there exist two major problems with this kind of preprocessing: 1. Derivatives amplify noise (SNR is often low at high frequencies which are amplified the most) and 2. The optimal differentiating filter is non-causal. The first problem may be dealt with by noise reducing lowpass-filtering (see [3] for an optimal approach). The second obstacle may be circumvented by delaying \( d(k) \) thus allowing for non-causal filtering i.e. estimating \( d(k - r) \) \((r \geq 0)\) using \( z(k), z(k - 1), \ldots \).

Another method is principal component analysis (PCA) which serves two purposes: 1. PCA makes the components in \( z \) mutually uncorrelated (convergence speed-up for certain weight estimation algorithms, e.g. Backpropagation [8]) and 2. It determines the dimensionality which is done by removing the last \( L - p \) principal components (PC’s) with eigenvalues close to zero. The amount of lost information can be estimated as the total variance of the removed PC’s divided by the total variance of \( x \), i.e. \( L - V\{z(k)\} \). The remaining PC’s constitute the optimal linear projection (in the mean square sense) of \( x \) on the space spanned by the first \( p \) eigenvectors of the covariance matrix of \( x \). A theoretical wellfounded procedure for on-line estimation of the PC’s has recently been described [18]. Other schemes are given in [8, Chap. 8.3].

Further support for the use of PCA can be found in an information theoretical interpretation: Maximize the mutual information \( I(x; z) \) between \( x \) and \( z \). It can be shown that if \( z \) is Gaussian (also valid for certain similar probability density functions): \( \max I(x; z) = \max H(z) \leftrightarrow \max \det V\{z\} \) where \( V\{\} \) is the variance and \( H(.) \) is the entropy. PCA is in fact done by maximizing \( \det V\{z\} \) [10, p. 682] which implicates that dimensionality determination by means of PCA is equivalent to maximizing \( I(x; z) \). When dealing with signals corrupted by noise PCA is not always preferable (especially if the signal to noise ratio is low) because the PC’s then will reflect the noise. Furthermore using PCA when the spectral overlap of the signals \( z(k) \) and \( d(k) \) is small is not reasonable. This is due to the fact that the spectrum of the PC’s corresponding to large eigenvalues mainly contains the
dominating frequencies in $x(k)$ thus neglecting the frequencies that dominate the spectrum of $d(k)$.

Memoryless Multidimensional Nonlinearities

When approximating the MMNL, $F_n[x], z \in I$ where $I$ is the input space, we distinguish between local and global approximation methods [6]. In a local approximation context $I$ is divided into smaller domains. $F$ is now approximated in each domain by separate nonlinearities. This results in a modularization of the MMNL which ensures a sparse parameterization. By global approximation is meant, that no dividing of $I$ is done at all. In general there is a trade off between the number of domains and the complexity of the subsequent nonlinearities.

Global Approximation Methods. In this case we deal with only one nonlinearity which must have the ability of approximating $F$ arbitrarily accurate. We will discriminate between fixed and adaptive nonlinearities.

A natural choice of a fixed nonlinearity (FNL) is to let $v$ contain all possible products of $z_i$, e.g. terms of the form $\prod_{i=1}^p z_i^{s_i}$ up to some order $s = \sum_{i=1}^p s_i$. When these terms are added by the linear combiner it all amounts to a multidimensional Taylor expansion which combined with the linear filters in the preprocessor defines a discrete Volterra filter. Fréchet showed [15] that any continuous $F(x(t))$ can be represented by a Volterra filter with uniform convergence when $s \to \infty$ for $x(t) \in J, J \subseteq I$. A convenient representation can be obtained by using a complete set of orthogonal polynomials $P_i$, where $i$ is the order of the polynomial. If $z_i \in N(0, 1), P_i$ are identical to the Hermite polynomials. With these polynomials convergence in mean is assured over a suitable interval $[a, b]$. The generalization to the multidimensional case is done by forming all products of polynomials in different variables e.g. $\prod_{i=1}^p P_i(z_i)$ (see [15] for details). In general the probability density $f_z(x)$ is unknown which makes it impossible to find the orthogonal polynomials. Instead we propose the use of Chebychev polynomials preceeded by squashing functions that limits the $z_i$ to the interval $[-1; 1]$ thereby limiting the $n_i$ to $[-1; 1]$.

An obvious choice for an adaptive nonlinearity (ANL) is a layered feed-forward neural network composed of sigmoidal neurons. It is well-known (see e.g. [5]) that a two layer feed-forward network with a linear output neuron (under rather mild conditions on the activation function, $g$) can uniformly approximate any function as the number of neurons in the hidden layer reaches infinity. We suggest that the nonlinearity is composed of two layers. The first layer consists of $p$ neurons which maps $z_i, 1 \leq i \leq p$ into $g(z_i w^1 + w^2_0)$ ($w^1_0$ ensures a proper scaling, see below). The second layer consists of $q$ neurons ($q \to \infty$ for an arbitrarily accurate approximation) and the outputs then form the nonlinear terms $[v_1, \ldots, v_{q-p}]^t$. It is further suggested to explicitly model the linear part by letting $[v_{q-p+1}, \ldots, v_q]^t = [z_1, \ldots, z_p]^t$.

Local Approximation Methods. A possible way to divide the input space is to use Localized Receptive Fields [14]. The output from each re-
ceptive field is then fed into separate nonlinearities. As above they could be either fixed or adaptive. Note that there is a trade-off between the number of domains in the input space and the complexity of the succeeding nonlinearities. Other local approximation schemes can be found in [6], [16], [15, Chap. 21].

Scaling of $z$. Scaling of $z$ serves two purposes. First, we have to restrict $z_i$ to an interval where the nonlinearity is slowly varying; i.e. neither growing towards infinity (as polynomials for large arguments) nor being constant (like $\tanh(\cdot)$ for large arguments). Secondly, we have to ensure that only the significant amplitude range of $z_i$ (i.e. the interval where $f_{zi}(z_i) > \varepsilon$, $0 < \varepsilon < 1$) is fed into the filter. Otherwise very unlikely values of $z_i$ will be weighted too much in the cost function thus resulting in a poor performance. Scaling with a suitable measure of $z_i$, e.g. 2-3 standard deviations, serves this purpose.

Weight Estimation Algorithms
The task is to estimate the weights $\theta$ so that the cost function $\sum_{k=1}^{N} \lambda^{k-1} e^2(i)$ is minimized [7] where $e$ is the difference between the desired and the actual response and $0 < \lambda \leq 1$ is the forgetting factor.

Fixed Nonlinearity. In designs with a FNL it is only necessary to adapt the linear combiner. This is especially simple if $x$ is Gaussian and PCA is used as preprocessing making $z$ white (independent) and Gaussian. Now if $z_i$ is scaled to unity variance and Hermite polynomials are used in the nonlinearity then the $v_j$ will be uncorrelated and the weights may thus be updated using the crosscorrelation method proposed in [15]: $a_j = C\{v_j d\} / V\{v_j\}$, $1 \leq j \leq q$ where $C\{v_j d\}$ is the covariance between $v_j$ and $d$ and $V\{v_j\}$ is the variance of $v_j$. In most cases, however, $v$ is non-white but owing to the fact that $y(k)$ is linear in the weights, adaptive algorithms known from linear adaptive filtering such as the recursive least squares (RLS) [7, p. 385] or the least mean squares (LMS) [17, p. 99] are usable. The latter is perhaps the best choice for large values of $q$ because it needs less computations and memory capacity while the major advantage of the RLS is the much faster convergence for highly correlated inputs ($v_j$).

Adaptive Nonlinearity. Designs with ANL implicate that estimation of the weights is a nonlinear optimization task which in general is hard to solve (global optimization) but local optimization schemes have been given, e.g. Backpropagation (BP). BP is known to have very slow convergence [4]. There is therefore a need for development of algorithms with faster convergence. Several second-order algorithms (SOA) have been proposed, see e.g. [4]. A SOA incorporates the information contained in the Hessian ($H$) of the cost function and the weights are updated according to the Newton-Raphson algorithm. In contrast to BP the SOA parameters are given a natural interpretation. $0 < \lambda \leq 1$ is the exponential forgetting factor, $0 < \mu \leq 1$ is the stepsize which normally is non-critical, and $\delta (H^{-1} = \delta H)$ is initially chosen large. We suggest a further development that takes the problems of nearly...
singular Hessian matrices into account. This problem arises in “flats” parts of the cost function. It is proposed to use the U-D factorization of $H^{-1}$ due to Bierman [2].

SIMULATIONS

Simulated Systems

In order to compare filter designs when filtering signals with different origin and distribution we study three systems covering the cases 2–4 on p. 3.

<table>
<thead>
<tr>
<th>Example</th>
<th>x</th>
<th>d</th>
<th>Model</th>
</tr>
</thead>
<tbody>
<tr>
<td>System Identification (SI)</td>
<td>Band-limited Gaussian noise</td>
<td>Non-Gaussian</td>
<td>Equation (5)</td>
</tr>
<tr>
<td>Inverse Modelling (IM)</td>
<td>Non-Gaussian</td>
<td>Gaussian lowpass filtered noise</td>
<td>Equation (6)</td>
</tr>
<tr>
<td>Prediction (P)</td>
<td>Non-gaussian</td>
<td>Non-gaussian</td>
<td>Equation (7)</td>
</tr>
</tbody>
</table>

**System Identification.** A simple system describing a wave force problem is given by Morison’s equation [1, p. 234]. $x(t)$ is the wave velocity and $d(t)$ the wave force. The desired signal $d(k)$ is a discrete version of $d(t)$ sampled with the sampling period $\Delta T = 1$ and the same applies to $z(k)$.

$$d(t) = 0.2 \frac{dx(t)}{dt} + 0.8 z(t) x(t)$$

(5)

**Inverse Modelling.** We consider the pendulum where $x(t)$ is the angle deflection and $d(t)$ the force. The desired signal $d(k)$ is a discrete version of $d(t - \tau)$ (sampled with $\Delta T = 0.05$) where $\tau$ is a delay aiming to cancel both the delay between $d(t)$ and $x(t)$ and the delay in the preprocessing unit of the nonlinear filter. $z(k)$ corresponds to the angle $z(t)$.

$$\frac{d^2z(t)}{dt^2} + 0.2 \frac{dz(t)}{dt} + 4\pi^2 \sin[z(t)] = d(t)$$

(6)

**Prediction.** The signal $z(t)$ is generated by the chaotic Mackey-Glass equation which often is used in a benchmark test for nonlinear predictors [6]. $d(k)$ is a discrete version of $x(t)$ and $z(k)$ a delayed, discrete version of $x(t - \tau)$ where $\tau$ signifies how far ahead we predict. Sampling the signal with $\Delta T = 1$ $\tau$ equals 100 timesteps like in [6],[11].

$$\frac{dx(t)}{dt} = -0.1x(t) + \frac{0.2x(t - 17)}{1 + x(t - 17)^10}$$

(7)

The systems mentioned above have all been simulated using discrete approximations of the derivatives. These discrete filters are all non-recursive which means that a non-recursive nonlinear adaptive filter is adequate. The actual training and cross validation signals have been obtained by decimating the input and output signals in order to avoid oversampling.
Numerical Results

In the table below are listed the main results. A measure of the filter performance is given by the error index: \( E = \sigma_e/\sigma_d \), where \( \sigma_e \), \( \sigma_d \) denote the standard deviation of the error and the desired signals respectively (cross validation). The number of parameters \( W \) gives an indication of the complexity.

<table>
<thead>
<tr>
<th>Ex.</th>
<th>Prep.</th>
<th>( L )</th>
<th>( p )</th>
<th>Fixed</th>
<th>Adaptive</th>
<th>None</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>( E )</td>
<td>( W )</td>
<td>( E )</td>
</tr>
<tr>
<td>SI</td>
<td>PCA</td>
<td>14</td>
<td>4</td>
<td>0.263</td>
<td>94</td>
<td>0.215</td>
</tr>
<tr>
<td>SI</td>
<td>DPP</td>
<td>19</td>
<td>2</td>
<td>0.200</td>
<td>46</td>
<td>0.107</td>
</tr>
<tr>
<td>SI</td>
<td>None</td>
<td>14</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>IM</td>
<td>DPP</td>
<td>19</td>
<td>3</td>
<td>0.075</td>
<td>152</td>
<td>0.116</td>
</tr>
<tr>
<td>IM</td>
<td>None</td>
<td>19</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>P</td>
<td>PCA</td>
<td>20</td>
<td>4</td>
<td>0.152</td>
<td>170</td>
<td>-</td>
</tr>
<tr>
<td>P</td>
<td>None</td>
<td>20</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

In all simulations we have used 9000 samples for training and 8000 for cross validation. During training an algorithm based on a statistical test [12], [9] was used to eliminate non-significant weights which accounts for the variations in \( W \). The FNL consisted of bounded Chebychev polynomials and the ANL was implemented using a multilayer neural net. In both cases we used 2. order algorithms for adapting the weights. In general the simulations indicate that the nonlinear filters are clearly superior to the linear with respect to \( E \). This improvement is, however, gained at the expense of an increased complexity. The FNL and ANL's seems to show roughly equal performance on the selected examples, with the ANL having a better parameterization in the SI example (more significant weights and lower \( E \)) and vice versa in the IM example. The two preprocessing methods seem to complement each other. In the examples SI, IM, where the equations can be closely approximated with discrete derivative operators of low order, the use of discrete differentiating filters in the preprocessing unit yields a better performance with lower complexity than the use of PCA. This shows that the discrete derivatives are more informative than the PC's in the chosen examples. Using PCA in the example with the pendulum is in fact extremely bad because the PC's mainly reflect the low frequency components in \( z(k) \) while the high frequencies carry most of the information about \( d(k) \). In contrast, PCA works very well for the example of prediction whereas it is not easy to approximate the Mackey-Glass equation using only low-order derivatives (i.e. the use of a DPP is a bad choice). Finding an appropriate preprocessing method seems thus to require knowledge of an approximate mathematical model for the unknown system. Alternatively a rule of thumb saying that the preprocessor should make the spectra of the \( z(k) \) "close" to the spectrum of \( d(k) \) could be used. In the prediction example we have used a PCA with \( L = 20 \) and \( p = 4 \) which allows us to compare our results with the ones obtained in [11] where
a performance of $E = 0.054$ was found using an ANL and a total of 171 weights and $E = 0.28$ using a 6th order fixed polynomial nonlinearity and 210 weights. This indicates, that although the ANL still performs better on this example an increase in performance of the FNL has been gained by using bounded polynomials.

CONCLUSION

In this paper a neural architecture for adaptive filtering which incorporates a modularization principle is proposed. It facilitates a sparse parameterization, i.e. fewer parameters have to be estimated in a supervised training procedure. The main idea is to use a preprocessor which determine the dimension of the input space and further can be designed independent of the subsequent nonlinearity. Two suggestions for the preprocessor are presented: The derivative preprocessor and the principal component analysis. A novel implementation of fixed Volterra nonlinearities is given. It forces the boundedness of the polynomials by scaling and limiting the inputs signals. The nonlinearity is constructed from Chebychev polynomials. We apply a second-order algorithm for updating the weights for adaptive nonlinearities based on previous work of Chen et al. [4] and the U-D factorization of Bierman [2]. Finally the simulations indicate that the two kinds of preprocessing tend to complement each other while there is no obvious difference between the performance of the ANL and FNL.

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