Invertible Piecewise Linear Approximations for Color Reproduction

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Invertible Piecewise Linear Approximations for Color Reproduction

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Abstract

We consider the use of linear splines with variable knots for the approximation of unknown functions from data, motivated by control and estimation problems arising in color systems management. Unlike most popular nonlinear-in-parameters representations, piecewise linear (PL) functions can be simply inverted in closed form. For the one-dimensional case, we present a study comparing PL and neural network (NN) approximations for several function families. Preliminary results suggest that PL, in addition to their analytical benefits, are at least competitive with NN in terms of sum square error, computational effort, and training time.

1. Introduction

When a computer sends a color document to a laser printer, the color of each pixel is represented as a vector in a standard color space, such as Lab, a space based on the psychophysics of the eye. The printer transforms the vector into a device-dependent space, CMY, which specifies the quantity of each pigment the printer must lay down in order to reproduce the desired color. This transformation is nonlinear and, approximate first principles models notwithstanding, must be computed in practice from empirical data [1]. When collecting data, each experiment consists of specifying a CMY vector to the printer and measuring the output in Lab coordinates. Thus, the empirical data is gathered using the inverse of the desired transformation. Color science also indicates that the map will be injective, so it may be inverted, at least over its range. In current industry practice, the color transformation is performed by interpolation on a lookup table with approximately one to two thousand entries. Industry goals are to reduce the number of parameters required to perform the color transformation, thereby reducing the cost of calibration as well as hopefully yielding a functional representation which may be amenable to online updating methods as the function drifts and more calibration data is collected.

Beyond interpolation on a uniform grid, one of the most sophisticated techniques for approximating color space transformations currently in use in the color industry is sequential linear interpolation [2]. This approach applies asymptotic analysis from information theory to find the optimal (nonuniform) grid point placement.

Much attention has been given to various parameterizations of the space of approximations, especially nonlinear parameterizations such as Neural Networks (NN) and Radial Basis Function Networks (RBFN). In higher dimensions, the convergence per parameter rates for such nonlinear families can potentially be better than linear-in-parameter function families [3]. Unfortunately, popular nonlinear families like NN and RBFN generally do not admit the “leveraging” of additional domain knowledge about the function, such as invertibility. In our application setting, NNs and RBFNs require either i.) that a second network be trained in order to construct the function inverse or ii.) that some further numerical procedure be applied to generate the inverse. But in the color space transformation problem, the inverse map is just as important as the forward map. The printer physically realizes the inverse map and the application of control methodology seems most reasonable when working with the function most closely related to the physical system.

A novel approximation method suggested by Atkeson and Schaal [4] uses a population of local “experts.” Each “expert” is associated with an affine map and a Gaussian confidence. The “experts” vote on an output computed as the confidence weighted average of their affine components. Since the Gaussian bump has unbounded support, each expert has global influence.

Piecewise polynomial representations (splines) can also be applied to function approximation problems. Qualitatively, these may be thought of as local “experts” which have partitioned the domain. Analytical results are available for the one dimensional case for certain function families [5]. Algorithms exist for one dimension [6] as well as for higher dimensions [7], although analytical results hold only for the one-dimensional case. Stone et al. present a statistical theory for the rate of $L_2$ convergence [8].

2. Piecewise linear approximation

Piecewise linear approximations (PL), also known as linear splines with variable knots, comprise a function family in which the invertibility condition can be enforced, and the inverse can be calculated directly in closed form as well. In one dimension, the characterization of the piece-
wise linear approximation is straightforward. Consider a PL with \( n \) line segments on the domain \([0, 1]\). The PL is characterized by two vectors: \( d \in \mathbb{R}^{n+1} \) the vector of domain values, or knots, where \( 0 = d_0 < d_1 < \ldots < d_{n-1} < d_n = 1 \), and \( e \in \mathbb{R}^{n+1} \) the vector of codomain values. This gives \( 2n \) free parameters. Figure 1 shows a PL with four line segments.

Analytical results exist for the one-dimensional function approximation problem. In approximation, a completely known function is given and the objective is to find a PL which minimizes some norm, typically \( L_2 \), of the error. Barrow et al. [5] provide some generalized convexity conditions which imply the existence of a unique best \( L_2 \) fit from the class of piecewise linear approximations. Gayle and Wolfe [6] provide similar results for approximation using higher order splines. Their proof uses an algorithm to calculate the best approximation over the domain of all knot vectors, for which global, unique convergence is shown via application of the contraction mapping theorem.

In higher dimensions, the domain is partitioned into simplices: triangles in two dimensions, tetrahedra in three dimensions, and so on. Tourigny and Baines [7] present an algorithm for the two-dimensional function approximation problem, which can be generalized to higher dimensions. There are no corresponding analytical results. In order to produce an output for a given domain point in higher dimensions, the partition in which the domain point lies must first be identified. Since the partitions are nonuniform, this step rapidly increases in complexity with dimension. This points out a fundamental tradeoff between the complexity of the approximant (e.g., linear, quadratic, neural) and the complexity of the partition. (e.g., none, uniform, nonuniform)

### 3. Numerical studies

This section presents a numerical study designed to compare the relative approximation power of PL and NN approximations. The PL and NN were given the same number of free parameters in order to study the relative performance per parameter.

#### 3.1. Choice of function classes studied

Five different function families from the class of homeomorphisms on \([0, 1]\) were explored. That is, all functions mapped \([0, 1]\) to \([0, 1]\) and were continuous and invertible (i.e., monotonic). The five families fall into three groups: sigmoidal, piecewise linear and polynomial, chosen to "favor," respectively, NN, PL or neither. Examples of typical functions from these families are presented in Figure 2.

The sigmoidal group contains superpositions of hyperbolic tangents, of the form

\[
f(x) = k_1 \sum_{i=1}^{m} a_i \tanh(b_i (x - c_i)) + k_2
\]

where \( a_i \) and \( c_i \) are distributed uniformly on \([0, 1]\) and \( b_i \) is exponentially distributed with mean 30. Then \( k_1 \) and \( k_2 \) are chosen such that \( f(0) = 0 \) and \( f(1) = 1 \). The first family from the sigmoidal group has \( m = 5 \), so all functions in this family lie within the parameter space of the NN which was trained. Notice that this family is chosen to favor NN, since for \( m = 5 \) there exists a vector of NN parameter values which would give zero error. The second family has \( m = 15 \), so, while presumably favored, the neural network is underparameterized.

The piecewise linear group contains functions with \( m \) line segments, characterized by the points \( \{(x_i, y_i)\}_{i=0}^{m} \) with \( 0 = x_0 < x_1 < \ldots < x_{n-1} < x_n = 1 \) and \( 0 = y_0 < y_1 < \ldots < y_{n-1} < y_n = 1 \). The points \( x_i \) and \( y_i \) are chosen uniformly from \([0, 1]\) for \( i = 1, \ldots, n-1 \). The first family in the piecewise linear group has 10 line segments \((m = 10)\), so, again all functions in this family lie within the parameter space of the PL approximation. The second family has 30 line segments \((m = 30)\), so the PL is underparameterized.

The polynomial group consists of compositions of quadratic polynomials which satisfy \( f(0) = 0 \) and \( f(1) = 1 \) and are monotonically increasing. i.e. \( f'(x) \geq 0 \) for \( x \in [0, 1] \). Quadratic polynomials satisfying these constraints can be parameterized as

\[
f_{\alpha}(x) = (1 - \alpha)x^2 + \alpha x
\]

for \( \alpha \in [0, 2] \). Then the polynomial \( f = f_{\alpha_1} \circ f_{\alpha_2} \circ \ldots \circ f_{\alpha_m} \) is indeed a homeomorphism of \([0, 1]\), since it is the composition of homeomorphic functions, and it has degree \( 2^m \). The polynomial family presented here used \( m = 7 \) and the parameters \( \alpha_i \) were distributed uniformly over \([0, 1]\).

#### 3.2. Training methods

The PL algorithm employed here minimizes square error via gradient descent. Specifically, given a data set \( \{(x_i, y_i)\}_{i=1}^{N} \), the algorithm minimizes

\[
E = \frac{1}{2} \sum_{i=1}^{N} \left(y_i - f(z_i)\right)^2 = \frac{1}{2} \sum_{j=0}^{n-1} \sum_{x \in I_j} \left(y_i - f_j(z_i)\right)^2
\]
where \( I_j = [d_j, d_{j+1}] \) and \( f \) is the piecewise linear approximation, defined by

\[
f(x) = \begin{cases} \frac{c_{j+1} - c_j}{d_{j+1} - d_j} (x - d_j) & \text{for } x \in I_j \end{cases}
\]  

(4)

The partial derivatives are

\[
\frac{\partial E}{\partial c_i} = -R_{i-1} + R_i - S_i
\]  

(5)

\[
\frac{\partial E}{\partial d_i} = \Delta_i R_{i-1} - \Delta_i R_i + \Delta_i S_i
\]  

(6)

where

\[
R_i = \sum_{x_j \in I_i} \left( y_j - f_i(x_j) \right) \left( \frac{x_j - d_i}{d_{i+1} - d_i} \right)
\]  

(7)

\[
S_i = \sum_{x_j \in I_i} \left( y_j - f_i(x_j) \right)
\]  

(8)

\[
\Delta_i = \frac{c_{i+1} - c_i}{d_{i+1} - d_i}
\]  

(9)

The update law is then

\[
d_i^{k+1} = d_i^k - \mu \frac{\partial E}{\partial d_i}
\]  

(10)

\[
c_i^{k+1} = c_i^k - \mu \frac{\partial E}{\partial c_i}
\]  

(11)

where the superscript is the iteration number and \( \mu \) is the step size or "learning rate."

The PL algorithm takes advantage of the fact that it is approximating a function from the class of homeomorphisms on \([0, 1]\) by fixing \( c_0 = 0 \) and \( c_n = 1 \), in addition to \( d_0 = 0 \) and \( d_{n-1} = 1 \). This reduces the number of free parameters for a PL with \( n \) line segments from \( 2n \) to \( 2(n-1) \). Allowing a neural network to use this information would require significant modification of the backpropagation algorithm. The PL in the following experiments uses 10 line segments, giving a total of 18 free parameters. The PL gradient descent algorithm was implemented in Matlab.

The neural network was also implemented in Matlab using the Neural Network Toolbox. The network has six hyperbolic tangent neurons situated in a single hidden layer, providing a total of 18 free parameters for the NN. The standard backpropagation rule, which minimizes the square error via gradient descent, was used to train the network.

### 3.3. Design of study

One hundred functions were randomly chosen from each function family discussed above. The algorithms received two data sets generated from each function. The training set contains 136 input/output pairs evenly spaced over the domain, while the validation set has 68 pairs interspersed between the test data, following the heuristic that approximately 2/3 of the data should be used for training and the remaining third should be used for validation.

First the PL was trained on each function until a stopping condition based on the magnitude of the gradient was achieved or the maximum number of iterations (4000) was exceeded. The NN was trained on the same data, given the goal of attaining 1/4 the sum square error of the PL. The NN stopped when this goal was achieved or after the maximum number of iterations, or "epochs," was exceeded. The maximum number of epochs for the NN was set at 12000.

### 3.4. Results

The mean square error (MSE) on a data set \( \{(x_i, y_i)\}_{i=1}^N \) is defined as

\[
MSE = \frac{1}{N} \sum_{i=1}^N \left( y_i - f(x_i) \right)^2
\]  

(12)

where \( f(x) \) is the approximation given by NN or PL. Because MSE takes on a wide range of values, \( \log_{10}MSE \) is presented here. Table 1 shows the minimum, average, and maximum of \( \log_{10}MSE \) for PL and NN on each family, as well as the average number of iterations on that family. For simplicity, \( 10^{\log_{10}MSE} \) will be referred to hereafter as the mean MSE. Figure 3 shows the MSE for PL and NN on every function for each of the families. The solid line shows the ratio \( MSE_{PL}/MSE_{NN} \).

Table 1: Mean Square Error and Iteration Results

<table>
<thead>
<tr>
<th>Function Family</th>
<th>MSE (Validation Data)</th>
<th>Iter. ave</th>
</tr>
</thead>
<tbody>
<tr>
<td>tanh PL</td>
<td>-5.388 / -4.670 / -3.354</td>
<td>3026</td>
</tr>
<tr>
<td>m=5 NN</td>
<td>-4.538 / -3.236 / -2.423</td>
<td>12000</td>
</tr>
<tr>
<td>tanh PLH</td>
<td>-5.562 / -4.538 / -3.236</td>
<td>3216</td>
</tr>
<tr>
<td>m=15 NN</td>
<td>-4.427 / -3.566 / -2.869</td>
<td>12000</td>
</tr>
<tr>
<td>PL</td>
<td>-7.419 / -5.405 / -3.218</td>
<td>2783</td>
</tr>
<tr>
<td>m=10 NN</td>
<td>-4.749 / -4.257 / -3.721</td>
<td>3566</td>
</tr>
<tr>
<td>PL</td>
<td>-5.562 / -4.639 / -3.758</td>
<td>3216</td>
</tr>
<tr>
<td>m=30 NN</td>
<td>-4.213 / -3.430 / -2.685</td>
<td>12000</td>
</tr>
<tr>
<td>PL</td>
<td>-5.985 / -5.405 / -4.410</td>
<td>2243</td>
</tr>
<tr>
<td>m=7 NN</td>
<td>-5.177 / -4.276 / -3.526</td>
<td>12000</td>
</tr>
</tbody>
</table>

Notice that PL regularly achieves a smaller MSE than NN, with only a handful of exceptions, on all function families. This is true even for the two families which "favor" NN, superpositions of hyperbolic tangents. For the sigmoidal family with \( m = 5 \), it would be possible for NN to represent the functions exactly, but still the mean is \( 10^{-3.226} \), which is an order of magnitude worse than the PL. This is well illustrated by the ratios, \( MSE_{PL}/MSE_{NN} \). The results for the sigmoidal family...
The numerical results show that when the PL takes advantage of the fact that it is approximating a homeomorphism, it is able to achieve on average an order of magnitude lower mean square error on the tested function classes than a neural network with an equal number of parameters.

These results could in part be an artifact of the descent technique, since only simple gradient descent was used. Also, we did not investigate the change in the ratio of $MSE_{PL}/MSE_{NN}$ as the number of parameters given to PL and NN vary. Both the descent technique and the variation of MSE with parameters would be interesting followup work to this study.

Our future work will focus on higher dimensional algorithms for application in the color problem, and the investigation of new descent techniques, including non-gradient methods.

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References
Figure 2: Typical members of the function families

Figure 3: The mean square error attained by NN and PL on each function, as well as the ratio of their MSE, is plotted above. $MSE_{PL} \circ MSE_{NN} \times \frac{MSE_{PL}}{MSE_{NN}}$