BAYESIAN NON-LINEAR MODELING FOR THE PREDICTION COMPETITION

David J.C. MacKay Cavendish Laboratory, Cambridge, CB3 0HE. United Kingdom mackay@mrao.cam.ac.uk

ABSTRACT. The 1993 energy prediction competition involved the prediction of a series of building energy loads from a series of environmental input variables. Non-linear regression using 'neural networks' is a popular technique for such modeling tasks. Since it is not obvious how large a time-window of inputs is appropriate, or what preprocessing of inputs is best, this can be viewed as a regression problem in which there are many possible input variables, some of which may actually be irrelevant to the prediction of the output variable. Because a finite data set will show random correlations between the irrelevant inputs and the output, any conventional neural network (even with regularisation or 'weight decay') will not set the coefficients for these junk inputs to zero. Thus the irrelevant variables will hurt the model's performance.

The Automatic Relevance Determination (ARD) model puts a prior over the regression parameters which embodies the concept of relevance. This is done in a simple and 'soft' way by introducing multiple regularisation constants, one associated with each input. Using Bayesian methods, the regularisation constants for junk inputs are automatically inferred to be large, preventing those inputs from causing significant overfitting.

An entry using the ARD model won the competition by a significant margin.

1 Overview of Bayesian modeling methods

A practical Bayesian framework for adaptive data modeling has been described in (MacKay 1992). In this framework, the overall aim is to develop probabilistic models that are well matched to the data, and make optimal predictions with those models. Neural network learning, for example, is interpreted as an **inference** of the most probable parameters for a model, given the training data. The search in model space (*i.e.*, the space of architectures, noise models, preprocessings, regularizers and regularisation constants) can then also be treated as an inference problem, where we infer the relative probability of alternative models, given the data. Bayesian model comparison naturally embodies **Occam's razor**, the principle that states a preference for simple models.

Bayesian optimization of model control parameters has four important advantages. (1) No validation set is needed; so all the training data can be devoted to both model fitting and model comparison. (2) Regularisation constants can be optimized on-line, *i.e.* simultaneously with the optimization of ordinary model parameters. (3) The Bayesian objective function is not noisy, as a cross-validation measure is. (4) Because the gradient of the evidence with respect to the control parameters can be evaluated, it is possible to optimism a large number of control parameters simultaneously.

Bayesian inference for neural nets can be implemented numerically by a deterministic method involving Gaussian approximations, the 'evidence' framework (MacKay 1992), or

by Monte Carlo methods (Neal 1993). The former framework is used here.

NEURAL NETWORKS FOR REGRESSION

A supervised neural network is a non-linear parameterized mapping from an input \mathbf{x} to an output $\mathbf{y} = \mathbf{y}(\mathbf{x}; \mathbf{w})$. Here, the parameters of the net are denoted by \mathbf{w} . Such networks can be 'trained' to perform regression, binary classification, or multi-class classification tasks.

In the case of a regression problem, the mapping for a 'two-layer network' may have the form:

$$h_j = f^{(1)} \left(\sum_k w_{jk}^{(1)} x_k + \theta_j^{(1)} \right); \quad y_i = f^{(2)} \left(\sum_j w_{ij}^{(2)} h_j + \theta_i^{(2)} \right)$$
 (1)

where, for example, $f^{(1)}(a) = \tanh(a)$, and $f^{(2)}(a) = a$. The 'weights' w and 'biases' θ together make up the parameter vector \mathbf{w} . The non-linearity of $f^{(1)}$ at the 'hidden layer' gives the neural network greater computational flexibility than a standard linear regression. Such a network is trained to fit a data set $D = {\mathbf{x}^{(m)}, \mathbf{t}^{(m)}}$ by minimizing an error function, e.g.,

$$E_D(\mathbf{w}) = \frac{1}{2} \sum_{m} \sum_{i} \left(t_i^{(m)} - y_i(\mathbf{x}^{(m)}; \mathbf{w}) \right)^2.$$
 (2)

This function is minimized using some optimization method that makes use of the gradient of E_D , which can be evaluated using 'backpropagation' (the chain rule) (Rumelhart et al. 1986). Often, regularisation or 'weight decay' is included, modifying the objective function to:

$$M(\mathbf{w}) = \beta E_D + \alpha E_W \tag{3}$$

where $E_W = \frac{1}{2} \sum_i w_i^2$. The additional term decreases the tendency of a model to 'overfit' the details of the training data.

NEURAL NETWORK LEARNING AS INFERENCE

The above neural network learning process can be given the following probabilistic interpretation. The error function is interpreted as the log likelihood for a noise model, and the regularizer is interpreted as a prior probability distribution over the parameters:

$$P(D|\mathbf{w}, \beta, \mathcal{H}) = \frac{1}{Z_D(\beta)} \exp(-\beta E_D); \ P(\mathbf{w}|\alpha, \mathcal{H}) = \frac{1}{Z_W(\alpha)} \exp(-\alpha E_W).$$
 (4)

The minimization of $M(\mathbf{w})$ then corresponds to the **inference** of the parameters \mathbf{w} , given the data:

$$P(\mathbf{w}|D,\alpha,\beta,\mathcal{H}) = \frac{P(D|\mathbf{w},\beta,\mathcal{H})P(\mathbf{w}|\alpha,\mathcal{H})}{P(D|\alpha,\beta,\mathcal{H})} = \frac{1}{Z_M} \exp(-M(\mathbf{w})).$$
 (5)

This interpretation adds little new at this stage. But new ideas emerge when we proceed to higher levels of inference.

Setting regularisation constants lpha and eta

The control parameters α and β determine the flexibility of the model. Bayesian probability theory can tell us how to set these parameters. All we need to do is write down the inference we wish to make, namely the probability of α and β given the data, and then use Bayes' theorem:

$$P(\alpha, \beta | D, \mathcal{H}) = \frac{P(D | \alpha, \beta, \mathcal{H}) P(\alpha, \beta | \mathcal{H})}{P(D | \mathcal{H})}$$
(6)

The data-dependent term, $P(D|\alpha, \beta, \mathcal{H})$, is the normalizing constant from our previous inference (5); we call this term the 'evidence' for α and β . This pattern of inference continues if we wish to compare our model \mathcal{H} with other models, using different architectures, regularizers or noise models. Alternative models are ranked by evaluating $P(D|\mathcal{H})$, the normalizing constant of inference (6).

Assuming we have only weak prior knowledge about the noise level and the smoothness of the interpolant, the evidence framework optimizes the constants α and β by finding the maximum of the evidence. If we can approximate the posterior probability distribution by a Gaussian,

$$P(\mathbf{w}|D, \alpha, \beta, \mathcal{H}) \simeq \frac{1}{Z_M'} \exp\left(-M(\mathbf{w}_{\text{MP}}) + \frac{1}{2}(\mathbf{w} - \mathbf{w}_{\text{MP}})^{\text{T}}\mathbf{A}(\mathbf{w} - \mathbf{w}_{\text{MP}})\right),$$
 (7)

then the maximum of the evidence has elegant properties which allow it to be located online. I summarize here the method for the case of a single regularisation constant α . As shown in (MacKay 1992), the maximum evidence α satisfies the following self-consistent equation:

$$1/\alpha = \sum_{i} w_i^{\text{MP } 2}/\gamma \tag{8}$$

where \mathbf{w}^{MP} is the parameter vector which minimizes the objective function $M = \beta E_D + \alpha E_W$ and γ is the 'number of well-determined parameters', given by $\gamma = k - \alpha \mathrm{Trace}(\mathbf{A}^{-1})$, where k is the total number of parameters, and $\mathbf{A} = -\nabla\nabla\log P(\mathbf{w}|D,\mathcal{H})$. The matrix \mathbf{A}^{-1} measures the size of the error bars on the parameters \mathbf{w} . Thus $\gamma \to k$ when the parameters are all well-determined; otherwise, $0 < \gamma < k$. Noting that $1/\alpha$ corresponds to the variance σ_w^2 of the assumed distribution for $\{w_i\}$, equation (8) specifies an intuitive condition for matching the prior to the data, $\sigma_w^2 = \langle w^2 \rangle$, where the average is over the γ effective parameters; the other $k - \gamma$ effective parameters having been set to zero by the prior.

Equation (8) can be used as a re-estimation formula for α . The computational overhead for these Bayesian calculations is not severe: one only needs evaluate properties of the error bar matrix, \mathbf{A}^{-1} . In my work I have evaluated this matrix explicitly; this does not take a significant time if the number of parameters is small (a few hundred). For large problems these calculations can be performed more efficiently (Skilling 1993).

AUTOMATIC RELEVANCE DETERMINATION

The automatic relevance determination (ARD) model (MacKay and Neal 1994) is a Bayesian model which can be implemented with the methods described in (MacKay 1992).

Consider a regression problem in which there are many input variables, some of which are actually irrelevant to the prediction of the output variable. Because a finite data set will

show random correlations between the irrelevant inputs and the output, any conventional neural network (even with regularisation) will not set the coefficients for these junk inputs to zero. Thus the irrelevant variables will hurt the model's performance, particularly when the variables are many and the data are few.

What is needed is a model whose prior over the regression parameters embodies the concept of relevance, so that the model is effectively able to infer which variables are relevant and switch the others off. A simple and 'soft' way of doing this is to introduce multiple regularisation constants, one ' α ' associated with each input, controlling the weights from that input to the hidden units. Two additional regularisation constants are used to control the biases of the hidden units, and the weights going to the outputs. Thus in the ARD model, the parameters are divided into classes c, with independent scales α_c . Assuming a Gaussian prior for each class, we can define $E_{W(c)} = \sum_{i \in c} w_i^2/2$, so the prior is:

$$P(\lbrace w_i \rbrace | \lbrace \alpha_c \rbrace, \mathcal{H}_{ARD}) = \frac{1}{\prod Z_{W(c)}} \exp(-\sum_c \alpha_c E_{W(c)}), \tag{9}$$

The evidence framework can be used to optimism all the regularisation constants simultaneously by finding their most probable value, *i.e.*, the maximum over $\{\alpha_c\}$ of the evidence, $P(D|\{\alpha_c\}, \mathcal{H}_{ARD})$. We expect the regularisation constants for junk inputs to be inferred to be large, preventing those inputs from causing significant overfitting.

In general, caution should be exercised when simultaneously maximizing the evidence over a large number of hyperparameters; probability maximization in many dimensions can give results that are unrepresentative of the whole probability distribution. In this application, the relevances of the input variables are expected to be approximately independent, so that the joint maximum over $\{\alpha_c\}$ is expected to be representative.

2 Prediction competition: part A

The American Society of Heating, Refrigeration and Air Conditioning Engineers organized a prediction competition which was active from December 1992 to April 1993. Both parts of the competition involved creating an empirical model based on training data (as distinct from a physical model), and making predictions for a test set. Part A involved three target variables, and the test set came from a different time period from the training set, so that extrapolation was involved. Part B had one target variable, and was an interpolation problem.

The task

The training set consisted of hourly measurements from September 1 1989 to December 31 1989 of four input variables (temperature, humidity, solar flux and wind), and three target variables (electricity, cooling water and heating water) — 2926 data points for each target. The testing set consisted of the input variables for the next 54 days — 1282 data points. The organizers requested predictions for the test set; no error bars on these predictions were requested. The performance measures for predictions were the Coefficient of Variation ('CV', a sum squared error measure normalized by the data mean), and the mean bias error ('MBE', the average residual normalized by the data mean).

¹The quantity equivalent to γ is $\gamma_c = k_c - \text{Trace}_c(\mathbf{A}^{-1})$, where the trace is over the parameters in class c, and k_c is the number of parameters in class c.

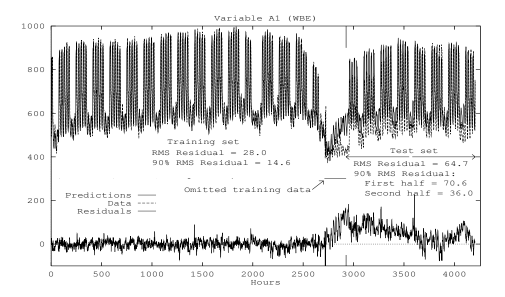


Figure 1: Target A1 — Electricity

The three target variables are displayed in their entirety, along with my models' final predictions and residuals, in figures 1–3.

Метнор

A large number of neural nets were trained using the ARD model, for each of the prediction problems. The data seemed to include some substantial glitches. Because I had not yet developed an automatic Bayesian noise model that anticipates outliers (though this certainly could be done (Box and Tiao 1973)), I omitted by hand those data points which gave large residuals relative to the first models that were trained. These omitted periods are indicated on some of the graphs in this paper. 25% of the data was selected at random as training data, the remainder being left out to speed the optimizations, and for use as a validation set. All the networks had a single hidden layer of tanh units, and a single linear output (figure 4). It was found that models with between 4 and 8 hidden units were appropriate for these problems.

A large number of inputs were included: different temporal preprocessings of the environmental inputs, and different representations of time and holidays. All these inputs were controlled by the ARD model. ARD proved a useful guide for decisions concerning preprocessing of the data, in particular, how much time history to include. Moving averages of the environmental variables were created using filters with a variety of exponential time constants. This was thought to be a more appropriate representation than time delays, because (a) filters suppress noise in the input variables, allowing one to use fewer filtered inputs with long time constant; (b) with exponentially filtered inputs it is easy to create (what I believe to be) a natural model, giving equal status to filters having timescales 1, 2, 4, 8, 16, etc..

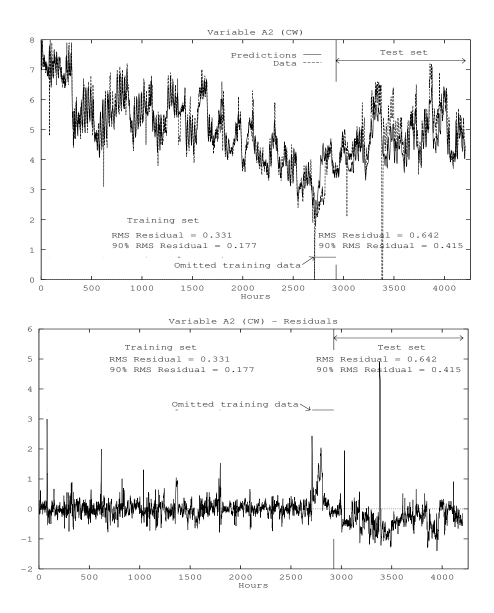


Figure 2: Target A2 — Cooling water

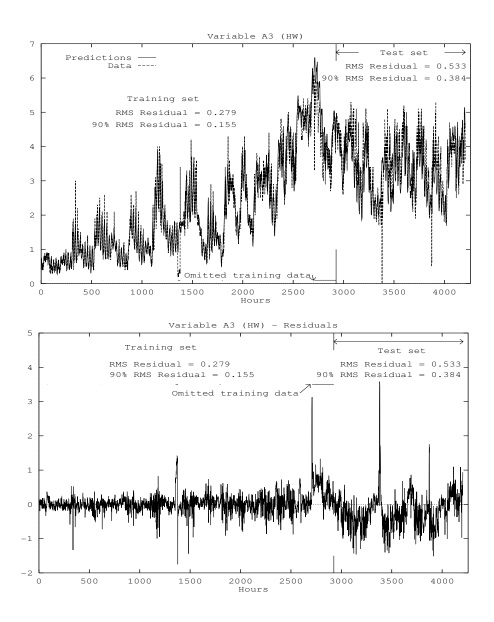


Figure 3: Target A3 — Heating water

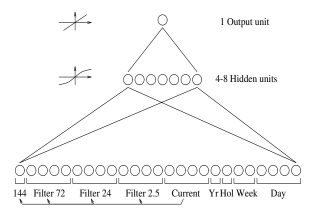


Figure 4: A typical network used for problem A

The filters produced moving averages of the four environmental inputs on three time-scales: 2.5, 24 and 72 hours. The temperature variable was also given a 144 hour filter. Time was represented using the cos of the year angle, a holiday indicator, and the cos and sin of: the week angle, the day angle, and twice the day angle. All hidden and output units also had a connection to a bias unit (not shown).

The on-line optimization of regularisation constants was successful. For problem A, 28 such control constants were simultaneously optimized in every model. The optimization of a single model and its control constants took about one day on a Sun 4 workstation, using code which could probably be made substantially more efficient. About twenty models were optimized for each problem, using different initial conditions and different numbers of hidden units. Most models did not show 'overtraining' as the optimization proceeded, so 'early stopping' was not generally used. The numerical evaluation of the 'evidence' for the models proved problematic, so validation errors were used to rank the models for prediction. For each task, a committee of models was assembled, and their predictions were averaged together (see figure 5); this procedure was intended to mimic the Bayesian predictions $P(\mathbf{t}|D) = \int P(\mathbf{t}|D,\mathcal{H})P(\mathcal{H}|D)d\mathcal{H}$. The size of the committee was chosen so as to minimize the validation error of the mean predictions. This method of selecting committee size has also been described under the name 'stacked generalization' (Breiman 1992). In all cases, a committee was found that performed significantly better on the validation set than any individual model.

The predictions and residuals are shown in figures 1–3. There are local trends in the testing data which the models were unable to predict. Such trends were presumably 'overfitted' in the training set. Clearly a model incorporating local correlations among residuals is called for. Such a model would not perform much better by the competition criteria, but its on-line predictive performance would be greatly enhanced.

In the competition rules, it was suggested that scatter plots of the model predictions versus temperature should be made. The scatter plot for problem A3 is particularly interesting. Target A3 showed a strong correlation with temperature in the training set (dots in figure 6b). When I examined my models' predictions for the testing set, I was surprised to find that, for target A3, a significantly offset correlation was predicted ('+'s in figure 6a).

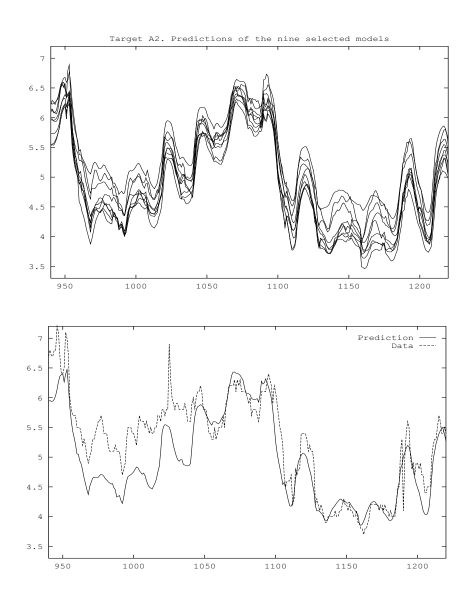


Figure 5: Target A2 — detail from test period

This figure shows detail from figure 2 and illustrates the use of a 'committee' of nine equally weighted models to make predictions. The diversity of the different models' predictions emphasizes the importance of elucidating the *uncertainty* in one's predictions. The *x*-axis is the time in hours from the start of the testing period. The prediction (lower graph) is the mean of the functions produced by the nine models (upper graph).

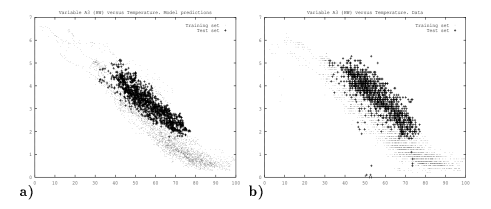


Figure 6: Predictions for target A3 (HW) versus temperature a) Model predictions. This graph shows that my model predicted a substantially different correlation between target A3 and temperature (+) from that shown in the training set (·). b) Data. This predicted offset was correct. Units: hot water /10⁶Btu versus temperature / F.

This change in correlation turned out to be correct ('+'s in figure 6b). This indicates that these non-linear models controlled with Bayesian methods discovered non-trivial underlying structure in the data. Most other entrants' predictions for target A3 showed a large bias; presumably none of their models extracted the same structure from the data.

In the models used for problem A3, I have examined the values of the parameters $\{\alpha_c, \gamma_c\}$, which give at least a qualitative indication of the inferred 'relevance' of the inputs. For prediction of the hot water consumption, the time of year and the current temperature were the most relevant variables. Also highly relevant were the holiday indicator, the time of day, the current solar and wind speed, and the moving average of the temperature over the last 144 hours. The current humidity was not relevant, but the moving average of the humidity over 72 hours was. The solar was relevant on a timescale of 24 hours. None of the 2.5 hour filtered inputs seemed especially relevant.

HOW MUCH DID ARD HELP?

An indication of the utility of the ARD prior was obtained by taking the final weights of the networks in the optimal committees as a starting point, and training them further using the standard model's regularizer (i.e., just three regularisation constants). The dotted lines in figure 7 show the validation error of these networks before and after adaptation. As a control, the solid lines show what happened to the validation error when the same networks were used as a starting point for continued optimization under the ARD model. The validation error is a noisy performance measure, but the trend is clear: the standard models suffer between 5% and 30% increase in error because of overfitting by the parameters of the less relevant inputs; the ARD models, on the other hand, do not overfit with continued training. The validation errors for the ARD model in some cases change with continued training, because my restarting procedure set the α_i to default values, which displaced the model parameters into a new optimum.

On the competition test data, the performance difference between these two sets of

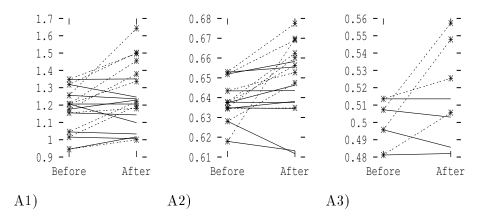


Figure 7: Change in validation error when the ARD prior is suspended The solid lines without stars show the performance of ARD models. The dotted lines with stars show the models with ARD suspended. In most cases, these standard ('ARD off') models get significantly worse.

models is not so pronounced, because the residuals are dominated by other effects. Maybe the greatest contribution of the ARD method to this problem was that it guided the choice of input variables to include large time-delays.

After the competition, it was revealed that the building in this study was a large university engineering center in Texas. Some of the glitches in the data were caused by the bursting of cold water pipes during a frost — a rare event apparently not anticipated by Texan architects!

The holiday period for staff ended on January 1st, but the student population did not return to the building for a couple of weeks. This may account for the significant bias error in the predictions of electricity usage (figure 1). Another factor which changed between the training period and the test period is that the Computer Science department moved to another building. This too will have caused a reduction in electricity usage. The reduction in electricity consumption may also account for some fraction of the biases in the cold and/or hot water supplies: one might expect less cooling water to be used, or more heating water, to make up the missing energy. The observed average electrical power deficit (according to my model) of 50 kW corresponds to an expected decrease in CW or increase in HW consumption of $0.17 \times 10^6 \text{Btu}$ (assuming that the CW and HW figures measure the actual energy delivered to the building). This is only about a fifth of the overall shift in correlation between HW and temperature shown in figure 6b. In fact, relative to my models, both CW and HW showed an increase of about $0.2 \times 10^6 \text{Btu}$.

3 Prediction competition: part B

The data for part B consisted of 3344 measurements of four input variables at hourly intervals during daylight hours over about 300 days. Quasi-random chunks of this data set had been extracted to serve as a test set of 900. The other 2444 examples were accompanied by a single target variable. The physical source of the data were measurements of solar

Problem A1	RMS	Mean	\mathbf{CV}	MBE	$RMS_{90\%}$	${ m Mean_{90\%}}$	RCV
ARD	64.7	50.3	10.3	8.1	54.1	42.2	11.1
ARD off	71.2	56.2	11.4	9.0	59.3	47.3	12.2
Entrant 6			11.8	10.5			
Median			16.9	-10.4			
Problem A2	RMS	Mean	$\overline{ ext{CV}}$	MBE	$RMS_{90\%}$	${ m Mean}_{90\%}$	RCV
ARD	.642	314	13.0	-6.4	.415	296	11.2
ARD off	.668	367	13.5	-7.4	.451	349	12.2
Entrant 6			13.0	-5.9			
Median			14.8	-7.6			
Problem A3	RMS	Mean	$\overline{ ext{CV}}$	MBE	$RMS_{90\%}$	Mean _{90%}	$\overline{\text{RCV}}$
ARD	.532	204	15.2	-5.8	.384	167	9.15
ARD off	.495	121	14.2	-3.5	.339	094	8.08
Entrant 6			30.6	-27.3			
Median			31.0	-27.0			
Problem B	RMS	Mean	$\mathbf{C}\mathbf{V}$	MRE	RMS	Meanage	\mathbf{RCV}

Problem B	RMS	Mean	\mathbf{CV}	MBE	$RMS_{90\%}$	${ m Mean_{90\%}}$	RCV
ARD	11.2	1.1	3.20	0.32	6.55	0.67	.710
Entrant 6			2.75	0.17			
Median			6.19	0.17			

Key:

My models:

ARD The predictions entered in the competition using the ARD

model.

ARD off Predictions obtained using derived models with the standard

regularizer.

Other entries:

Entrant 6 The entry which came 2nd by the competition's average CV

score.

Median (by magnitude) of scores of all entries in competition.

Raw Performance measures: Robust Performance measures: RMS Root $RMS_{90\%}$ Root mean mean squaresquare residual. of the smallest 90% of the MeanMean residual. residuals. \mathbf{CV} Mean of those residuals. Coefficient $Mean_{90\%}$ \mathbf{RCV} $RMS_{90\%}/(90\% \text{ data range}).$ of variation (percentage). The competition perfor-

mance measure.

MBE Mean Bias Error (percentage).

Normalizing constants:	$\operatorname{Problem}$	Mean of test data	90% data range
	A1	624.77	486.79
	A2	4.933	3.7
	$\mathbf{A3}$	3.495	4.2
	В	350.8	923

Table 1: Performances of different methods on test sets

flux from five outdoor devices. Four of the devices had a fixed attitude. The fifth, whose output was to be predicted, was driven by motors so that it pointed at the sun. The aim is to enable four cheap fixed devices to substitute for one expensive moving one. Clearly, information such as the day of the week and past history of the input variables was not expected to be relevant. However, I did not realize this, and I spent some time exploring different temporal preprocessings of the input. Satisfyingly, all time-delayed inputs, and the time of the week, were correctly found to be irrelevant by the ARD model, and I pruned these inputs from the final models used for making predictions — without physical comprehension of the problem.

The inputs used in the final models were the four sensor measurements, and a five dimensional continuous encoding of the time of day and the time of year. For training, one third of the training set was selected at random, and the remaining two thirds were reserved as a validation set. This random selection of the training set was later regretted, because it leaves randomly distributed holes where there are no training data. This caused my models' predictions to become unnecessarily poor on a small fraction of the testing data. As in part A, a committee of networks was formed. Each network had between 5 and 10 hidden units.

RESULTS

Problem B was a much easier prediction problem. This is partly due to the fact that it was an interpolation problem, with test data extracted in small chunks from the training set. Typical residuals were less than 1% of the data range, and contrasts between different methods were not great. Most of the sum-squared error of my models' predictions is due to a few outliers.

4 Discussion

The ARD prior was a success because it made it possible to include a large number of inputs without fear of overfitting.

Further work could be well spent on improving the noise model, which assumes the residuals are Gaussian and uncorrelated from frame to frame. A better predictive model for the residuals shown in figures 1–3 might represent the data as the sum of the neural net prediction and an unpredictable, but auto-correlated, additional disturbance. Also, a robust Bayesian noise model is needed which captures the concept of outliers.

In conclusion, the winning entry in this competition was created using the following data modeling philosophy: use huge flexible models, including all possibilities that you can imagine might be appropriate; control the flexibility of these models using sophisticated priors; and use Bayes as a helmsman to guide the search in this model space.

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