A Bayesian Neural Network for Hail Size Prediction

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Abstract

The National Severe Storms Laboratory has developed numerous algorithms that compute a number of Doppler radar and environmental attributes believed to be relevant for the detection/prediction of severe hail. Based on these attributes a neural network has been developed for the estimation of hail size. In this article, the development of two sets of neural networks is outlined: one for predicting the hail size, and another for assigning a probability of belonging to one of three hail size classes. Emphasis is placed on multidimensional (i.e., nonscalar) means of assessing performance. It is shown that the network designed to predict hail size outperforms the existing method for estimating hail size, and the network designed for classifying hail size produces highly reliable and discriminatory probabilities for two of the three hail-size classes - the smallest and the largest; forecasts of midsize hail, though highly reliable, are found to be mostly nondiscriminatory.
1 Introduction

The National Severe Storms Laboratory (NSSL) has developed numerous algorithms for the detection of various atmospheric phenomena. These algorithms reside collectively within an “umbrella” program called the Severe Storm Analysis Package (SSAP). Two such algorithms designed for the detection of tornados and/or damaging wind have recently been supplemented with Neural Networks (NNs) (Marzban and Stumpf 1995, 1998) and their performance has been compared with conventional statistical methods in Marzban, Paik, and Stumpf (1997). The NSSL has also developed a Hail Detection Algorithm (HDA) that computes a number of attributes believed to be relevant for the detection/prediction of severe hail (Witt et al. 1998a). As such, it is natural to develop a similar NN complementing the HDA.

The statistical theory of NNs and the art of employing them is now well developed (Bishop 1996, Ripley 1996). It has become common practice in many model-building tasks to examine NNs along side traditional methods. There is usually only one disadvantage, and that is NN’s tendency to overfit data. Overfitting occurs when a model captures the statistical noise in the data rather than the underlying signal. A number of popular methods for dealing with the problem of overfitting utilize several independent data sets; these methods fall under the general classes of split-sample and resampling methods. Examples are cross-validation, jack-knifing, and bootstrap (Bishop 1996). In these methods, one or more of data subsets are employed for estimating the parameters of the NN (i.e. training), and the remaining subsets are employed for determining the optimal complexity (i.e., nonlinearity) so as to prevent overfitting.

There exists an approach, referred to as the evidence framework (MacKay 1996) which in principle allows the use of the entire data for training, and without overfitting. Such a method has been illustrated in the context of tornado prediction (Marzban 1998). One feature of the evidence framework is its reliance on Bayesian inference. On the other hand, it invokes certain assumptions that may or may not be satisfied (Bishop 1996). However, the use of Bayesian inference to prevent overfitting is a valid and desirable feature. So, it is possible to utilize this feature in combination with some split-sample or resampling method. The use of split-sample or resampling and Bayesian inference in
order to preclude overfitting is particularly important when the sample size is small as compared to the number of predictors, as is the case here. Indeed, a non-Bayesian NN for hail size prediction developed recently (Marzban and Witt 2000) does display some symptoms of overfitting.

The development of a NN model for HDA can be divided into two sub-tasks: One of developing a model that predicts the occurrence of severe hail, and another that predicts the size of severe hail, given that severe hail has occurred. Only the latter will be considered in this article, as the former model is currently under construction.

Even a NN for the prediction of size alone can be developed in two independent ways: One can develop a NN that predicts the size of hail in some physical unit, or one can assess the probability of belonging to some size range. The data suggests that severe hail reports fall naturally into three different classes, corresponding to coin-size, golfball-size, and baseball-size, in an ordinal fashion. As such, it is possible to assess the probability of a hail report belonging to each of these three classes. The former approach falls in the domain of regression, while the latter is an example of a classification problem. Both NNs will be considered herein, and will be referred to as the regression NN and the classification NN, respectively.

In what follows the data and the methodology are further described, and multidimensional (e.g., distribution-based) measures of performance are set forth in terms of which the performance of the NNs is gauged. Finally, a discussion section offers some final thoughts on the matter of a NN for hail size prediction.

2 Data

The input variables provided to the NN include a mix of Doppler-radar derived parameters along with several parameters representing the near-storm environment. The radar parameters include two based on reflectivity data, cell-based vertically integrated liquid (Johnson et al. 1998) and the severe hail index (Witt et al. 1998a), as well as two based on velocity data, storm-top divergence (Witt and Nelson 1991) and midaltitude rotational velocity (Witt 1998). The near-storm environment parameters include three based on thermodynamic data and two based on kinematic data (Table 1). The vertically-integrated
wet-bulb temperature parameter is computed by integrating the wet-bulb temperature profile from the surface to the height of the wet-bulb zero. These near-storm environment parameters are either calculated within the SSAP using numerical model data, or they can be calculated from sounding data and manually entered into an adaptable parameter file. For this study, all the near-storm environment parameters were calculated from sounding data.

The verification data on hail size comes from Storm Data. Because Storm Data is a collection of severe weather reports, the minimum hail size in this study is 19 mm (0.75 inch). There are numerous problems associated with using Storm Data for verifying radar-based algorithm predictions (Witt et al. 1998a,b). For prediction of maximum hail size, the primary concern involves the possibility that any given hail report is not representative of the maximum size being produced by the storm (at the time of the report). To minimize the impact of this possibility, the analysis was restricted to the maximum size observed per hailstorm (Witt 1998). For each severe hailstorm, a 20 minute time window (Witt et al. 1998a) was used to relate the predictor variables to the maximum reported hail size. For the radar-based parameters, the maximum value within the time window was used, whereas for the near-storm environment parameters, an average value was used.

In situations where there are multiple reports of the same maximum hail size for a storm, the report corresponding to the time period when the storm appeared to be weaker was used, since these radar characteristics were indicative of the minimum strength necessary to produce the observed maximum hail size. For example, suppose one is using Vertically Integrated Liquid (VIL) to predict maximum hail size. Then, if there are two reports of golfball-size hail with a storm, and the VIL is 50 for one report and 60 for the other, it is reasonable to assume that a VIL of 50 is the representative value associated with golfball-size hail for this storm. However, this condition (using the weaker period) only applies to multiple reports where all the predictor variables have been measured within the time window, i.e., the storm-top divergence or midaltitude rotational velocity parameters are not “missing” due to range folding. In cases where the choice was between using a stronger period with a full set of data or a weaker period which includes missing data, then the stronger period was used.

Algorithm output was generated for 81 storm days from across the U.S. (Table 2), on
which 386 severe hailstorms were observed. The distribution of maximum reported hail sizes for these 386 reports is shown in Fig. 1. The common practice of reporting hail size using familiar circular or spherical objects (e.g., various coins or balls) is clearly evident, as reports tend to be clustered along discrete sizes. The highest frequency corresponds to dime, nickel and quarter (coin) size hail (19 - 25 mm), golf ball size hail (44 mm) and baseball size hail (70 mm). It would appear that few hail reports are actually measured to obtain a precise reading of their size, and that most hail sizes are estimated. Hence, one must assume that a certain amount of “rounding-off” error exists in the observations, and this error appears to increase as hail size increases. Because of all this, the prediction of maximum hail size was approached from two distinct directions - regression and classification.

3 Bayesian Neural Network

All nonlinear regression and classification models can overfit data; overfitting occurs when the flexibility/complexity of a model allows it to fit a data set or a decision boundary to such high accuracy that the fit is driven by the statistical fluctuations in the data. Consequently, such a model has no predictive capability. The true cause of this phenomenon is the finiteness of the sample size, and it occurs mostly when the model is allowed to be highly nonlinear.

In NNs, nonlinearity is determined by two quantities, the number and the magnitude of the independent parameters (weights). The former is determined by the number of hidden nodes (if the inputs are not collinear), and the latter is determined by the variance of the initial weights. If the number of independent weights greatly exceeds the sample size, and the magnitude of the weights greatly exceeds unity (if the inputs are normalized in some fashion), then an NN can overfit data. If, however, either of these conditions is not met, then overfitting is unlikely. For instance, if the magnitude of the weights is restricted to be much less than one, then most activation functions are linear, and so an NN is nothing but linear regression, regardless of the number of hidden nodes.

As such, it is possible to restrain overfitting, even with a large NN (i.e., with many hidden nodes), if the weights are prevented from becoming too large. The question is then
“How large is too large?” The magnitude of the weights can be constrained by introducing a weight-decay term into the error function that is to be minimized (Bishop 1996). The question then becomes one of determining the optimal value of the coefficient of this term. It is for this purpose that techniques of Bayesian inference can be employed. One of the advocates of this method is MacKay (1996), and the methodology followed in this article is also that of MacKay. However, the quantity defined as evidence will not be utilized here; instead, resampling will be employed to further restrict overfitting. The NN program employed in this research is a variation on that developed by MacKay and made available on the Web at http://wol.ra.phy.cam.ac.uk/mackay.

Both the split-sample and resampling methods rely on splitting the data into three sets - a training set, employed to estimate the parameters of the model, a validation set for optimizing the complexity of the model, and a test set for estimating the unbiased performance of the model. So, only two sets - a training and a validation set - are sufficient for the purpose of selecting a model. For small sample sizes, it is preferable to employ a resampling method instead of a split-sample method such as cross-validation (Bishop 1996). An example of a resampling method for model selection is Bootstrapping (Efron and Tibshirani 1993). In its simplest form, one repeatedly trains with subsamples of the data, and the optimal NN is selected to be the one with the lowest average error over the unused subsamples. From the variance (over the subsamples) of the validation errors one can construct a confidence interval for the performance of the NN.

In addition to the variance due to resampling, in NNs there exists a variance due to local minima. In other words, there is no assurance that all the NNs trained on the different subsamples rest in the same minimum of the error function. Of course, in practice, only the deepest, practically reachable local minimum is of interest. The question then arises as to what precisely is the “deepest” local minimum. The answer lies in the range over which the local minima vary. Specifically, the distribution (over the different subsamples) of the values of the error function at the local minima can aid in assessing the likelihood of obtaining a deeper local minimum than a given one. Thus, a local minimum that is highly unlikely to be won over by a deeper one is considered to be the deepest local minimum. This issue has been thoroughly discussed in Marzban (2000) and Marzban and Witt (2000); suffice it to say that the “global” minimum referred to in
the current article is the deepest local minimum as outlined above.

4 Method

As mentioned previously, the problem of predicting maximum hail size can be approached either from the point of view of regression or classification. This can be illustrated by examining the distribution of the hail size data set (Fig. 1). The number of distinct values (i.e., 14) is sufficiently large to warrant a regression analysis wherein the predictand is hail size. On the other hand, the existence of clusters/peaks in that distribution suggests three distinct classes for hail size, corresponding to “coin size”, “golfball size”, and “baseball size.” As such, a classification approach is also feasible. Both approaches are fruitful in that the former provides estimates for hail size in some physical unit, and the latter can assess the probability of belonging to one of the three classes of hail size.

In the regression approach, an appropriate measure of error is the Mean Square Error (MSE), defined as

$$MSE = \frac{1}{N} \sum [t - y(x, \omega)]^2,$$

(1)

where $x$ is the vector of inputs (attributes), $\omega$ is the vector of the weights, and $t$ is the target value that is to be estimated by the output $y(x, \omega)$. The summation is over the number of cases, $N$, in the relevant data set (training, validation, etc.). The activation function for the hidden nodes is taken to be the logistic function, $f(x) = 1/(1 + \exp(-x))$, while that of the single output node is the linear function. The former introduces the necessary nonlinearity into the NN, and the latter allows for the output node to take the full range of values taken by the target.

In the classification approach, the appropriate error function is cross-entropy, defined as:

$$S = -\frac{1}{N} \sum [t \log y(x, \omega) + (1 - t) \log (1 - y(x, \omega))].$$

(2)

In the c-class case, with c output nodes representing class membership, if the activation functions are the softmax function (a generalization of the logistic function to multiple output nodes), then the minimization of $S$ yields outputs that can be interpreted as the posterior probability of class-membership, given the inputs (Richard and Lippmann 1991). Such a probability is precisely what is required for forecasting purposes. An additional
weight-decay term of the form

\[ S_\omega = \alpha \frac{1}{2} \sum \omega^2, \]

restrains the size of the weights (and therefore, overfitting) without affecting the probabilistic interpretation of the output nodes. The total error function is, therefore, \( S + S_\omega \).

Another assumption of Richard and Lippmann (1991) is that 1-of-c coding be employed for coding the classes among the output nodes. This means that 3 output nodes are required to represent 3 classes, with the largest output node designating the corresponding class. Accordingly, the NNs described herein have 3 output nodes. This is in contrast to the NNs discussed in Marzban and Witt (2000) wherein the problem of predicting the 3 classes was decomposed into three 2-class problems. In other words, three NNs were developed - one for discriminating between coin-size hail and otherwise, a second for discriminating between golfball-size hail and otherwise, and a third NN for discriminating between baseball-size hail and otherwise. As such, each network required only one output node, and a simple application of Bayes’ theorem showed that the output nodes represented posterior probabilities for the corresponding classes. The reason for this was to reduce the chance of overfitting by reducing the number of parameters between the hidden and the output layer. In the current analysis, however, overfitting is restrained through the weight-decay term in the error function, and so it is safe to allow for 3 output nodes.

In both the regression and the classification schemes the number of input nodes is 9 (or 7, if there are missing data; see the last paragraph in this section.), namely the total number of hail attributes computed by HDA and SSAP. Experimentation with smaller number of input nodes suggests that nothing is gained by employing subsets of the nine attributes as inputs (see the Discussion section). Each attribute is centralized (i.e., converted to z-scores) by subtracting the mean and dividing by the respective standard deviation. This can stabilize and even expedite the training procedure.

As mentioned previously, the number of hidden nodes (on one hidden layer) was determined via bootstrapping, and it was found unnecessary to go beyond one hidden layer. As expected, what is gained by employing the Bayesian procedure for inferring the strength of the weight-decay term (\( \alpha \)) in conjunction with bootstrapping is that no loss of performance is found even for larger number of hidden nodes. In other words, the
combination of the two methods yields a range of number of hidden nodes within which the NN’s performance is insensitive to the precise number of hidden nodes.

There is also the matter of missing data. One common approach is to simply replace any missing data with the average of the nonmissing data for each predictor. For a skewed distribution, however, this approach can be unsatisfactory. In the present application, the fortunate situation arises in which the missing data always appear in only two of the predictors. As such, it is possible to develop two NNs, one based on all the data and all 9 inputs (when all 9 inputs are available), and another trained on only 7 inputs for which there is no missing data. This was done for both the regression and the classification NN.

5 Performance Measures

Performance is a multifaceted quantity. Although single, scalar (i.e., one-dimensional) measures such as MSE and Cross entropy are minimized during the training phase, it is important to assess the performance of the NN in terms of multidimensional quantities such as scatter plots, distribution plots, or attributes diagrams (Wilks 1995, Murphy and Winkler 1992).

The performance of the regression network can be assessed in terms of the scatterplot of the actual versus predicted hail size. Such a plot can display regions of forecasts that may be problematic as well as regions where the NN performs superbly. Of course, it is possible to distill this two-dimensional measure into a single, scalar measure such as $R^2$ which reflects the amount of variance explained by the model, but such a reduction is apt to lead to loss of information. Another multidimensional view of performance is offered by residual plots (Draper and Smith 1981); a plot of the residues (i.e., actual minus predicted) as a function of predicted size; this, too, can display a wide range of multidimensional features of the model.

For the classification network, there are many more ways of expressing performance. Classification performance can be assessed not only in terms of scalar and multidimensional measures, but also through categorical and probabilistic measures. This gives rise to four possibilities, examples of which are given in Table 2. For instance, an example of a scalar, categorical measure is the Heidke Skill Score (Wilks 1995), and an example of a
probabilistic, scalar measure is the Ranked Probability Score (a multi-class generalization of the Brier Score) (Wilks 1995). Multidimensional measures are typically diagrams of some sort, such as ROC or attributes diagrams (more, below), and are clearly desirable and preferred over scalar measures.

The starting point for computing categorical measures is the contingency table. The elements of a contingency table, $C_{ij}$, are the number of observations in the $i^{th}$ class that are forecast as belonging to the $j^{th}$ class. One can construct numerous categorical measures each of which captures a different facet of performance. A (nonexhaustive) examination of such measures based on $2 \times 2$ contingency tables is given in Marzban (1998). While many such measures are easily generalized to the $3 \times 3$ case, others are constructed from more basic quantities which are defined only for a 2-class problem. That problem is exacerbated when forecasts are probabilistic. For instance, reliability and attributes diagrams do not generalize to multiple classes. \footnote{One generalization of reliability diagrams has been introduced by T. Hamill (1987). However, it appears that the proposed diagrams assess not reliability but some other facet of performance quality. The proposed diagrams have been computed for the current application; however, they suggest that the facet being assessed is totally “perfect.” In other words, deviations from perfect quality are all statistically insignificant. As such, the proposed diagrams do not offer a useful assessment of performance, and are not presented herein.}

Typically, one reduces the 3-class problem to three 2-class problems. In other words, the problem of classifying three objects - labeled “1”, “2”, and “3” - is broken down to three 2-class problems: classification of class “1” or otherwise, class “2” or otherwise, and class “3” or otherwise. Although this treatment of a 3-class problem can lead to some loss of information (Wilks 1995), the loss is partially compensated by the multidimensionality brought about via reliability or attributes diagrams. The 3-class problem in the present application is reduced in this fashion.

In a 2-class problem (i.e., with yes/no observations and forecasts), with “1” (“2”) representing “no” (“yes”), two basic quantities are the probability of detection and the false alarm ratio, defined as

$$
POD = \frac{C_{22}}{C_{21} + C_{22}}, \quad FAR = \frac{C_{12}}{C_{12} + C_{22}}.
$$

Another basic quantity is the false alarm rate, defined as

$$
FAT = \frac{C_{12}}{C_{11} + C_{12}}.
$$
One multidimensional measure of performance is the ROC diagram. A ROC diagram (Masters 1993) is simply a parametric plot of POD versus FAT, as a probability threshold is varied from 0 to 1. It is easy to show that a classifier with no ability to discriminate between two classes yields a diagonal line of slope one and intercept zero; otherwise, the ROC curve lies in the region above the diagonal line. The area under the curve is often taken as a scalar measure of the classifier’s performance, and so, a perfect classifier would have an area of 1 under its ROC curve, while a random classifier would have an area of 0.5 (i.e., the area under the diagonal line). In addition to its multidimensionality, another virtue of a ROC diagram is its ability to express performance without a specific reference to a unique probability threshold. A specific choice of the threshold calls for knowledge of the costs of misclassification which are user dependent. In this way, a ROC diagram offers a user-independent assessment of performance. Because the ROC diagram is based on inherently 2-dimensional quantities (POD and FAT), it has no natural extension to the 3-class case. Therefore, three ROC diagrams must be considered, one for each of the three classes.

Since the network produces probabilistic forecasts, it is possible to assess the quality of the forecasts without categorizing the forecasts. One commonly employed probabilistic measure is the Brier Skill Score which is based on the MSE in equation (1), and is defined as (Wilks 1995)

\[ BSS = 1 - \frac{\text{MSE}}{p(1 - p)}, \]

where \( p \) is the climatological probability of hail of a given size. Due to its scalar nature, BSS fails to capture all the facets of probabilistic forecasts. Consequently, the quality of the probabilities will be assessed within a probabilistic scheme (Murphy and Winkler, 1987, 1992; Wilks, 1995). According to Murphy and Winkler (1992), the various aspects of the quality of probabilistic forecasts can be captured by three diagrams - reliability (or Calibration), refinement, and discrimination diagrams. In terms of the conditional distribution of forecasts, \( f \), and observations, \( x \), a reliability diagram is a plot of \( p(x = 2\lvert f) \) as a function of \( f \).\(^2\) A refinement plot is the plot of the marginal distribution \( p(f) \) as a function of \( f \), and a discrimination plot is a plot of \( p(f\lvert x = 1), p(f\lvert x = 2), \ldots, p(f\lvert x = c) \) as a function of \( f \), where \( c \) is the total number of classes in the problem (here 3). Reliability

\(^2\)\( x = 1, \) and \( x = 2 \) refer to the nonexistence and existence of an event, respectively.
diagrams can be supplemented with the contribution of the forecasts to BSS and their resolution, with the resulting diagram referred to as an attributes diagram (Murphy and Winkler 1992). Resolution is the variance of the difference between the unconditional observations and conditional (on forecast) observations (Murphy and Daan 1985).

The reliability portion of an attributes diagram displays the extent to which the frequency of an event at a given forecast probability matches the actual forecast probability. A refinement diagram displays the sharpness of the forecasts, i.e., the extent to which very high or very low probabilities are issued, and a discrimination plot exhibits the extent to which the forecasts discriminate between the classes. All of these together present a (somewhat) complete representation of performance quality.

6 Results

As mentioned previously, two NNs have been developed - a regression NN designed to estimate hail size, and a classification NN for modeling the probability of the three hail-size classes. The optimal number of hidden nodes for both networks was found via bootstrapping, as described in the Method section. However, as expected within the Bayesian approach, the final results are quite insensitive to the precise number of hidden nodes. In particular, for the regression NN it was found that any number of hidden nodes between 0 and 4 yields results that are statistically equivalent. For the classification NN the range of the number of hidden nodes leading to comparable performance is 2 to 6. As a result, the number of hidden nodes was set at 2 for the regression NN, and at 4 for the classification NN.

According to the bootstrap approach, in order to assure that the results are not jeopardized due to sampling effects, one must repeat the entire training/validation process with different training and validation data. For the current analysis, 4 such trials are made and the results are presented either separately, or averaged over the trials.

Having identified the optimal NNs, one can turn to their performance. Again, only multi-dimensional measures will be employed in order to avoid any loss of information brought about by the use of scalar measures. Consequently, the following assessment is heavily diagram oriented.
We begin with the regression NN. In this case, it is possible to compare the performance of the NN with an existing model which is currently in operation in the WSR-88D HDA. The existing model is an empirically derived equation for the Maximum Expected Hail Size. The model, hereafter referred to as “MEHS”, is a nonlinear model based only on the severe hail index (Witt at al. 1998a). Figure 2 shows the training and the validation errors for the four bootstrap trials of the NN and the MEHS models. It can be seen that not only does the NN have training and validation mean square errors that are lower than those of MEHS, they are also more clustered. In other words, the NN consistently outperforms MEHS in terms of the mean square error of the forecasts.

Figure 3 shows the scatter plot of the NN and MEHS for one of the bootstrap trials. The dark dots are the training data and the circles are the validation data. From the general pattern of these figures, it is evident that the NN outperforms MEHS: (reluctantly) regressing to a scalar measure like $r^2$ of the fits provides a more quantitative measure of the goodness of fit. Values of $r^2$ approaching 1 imply a better fit. The $r^2$ of the training and validation data for the NN are 0.51 and 0.40, respectively. By comparison, the same quantities for MEHS are 0.34 and 0.29. Clearly the NN provides a far better fit to the data than MEHS does. Also shown, are the regression fits to the corresponding plots. It can be seen that whereas the NN’s fit to both the training and validation data sets produces diagonal lines of slope 1, the MEHS slopes fall short of that ideal value. This means that on the average there is perfect agreement between NN-predicted size and the actual size; by contrast, the MEHS-predicted size is typically higher than the actual size.

An examination of the residual-plots (Figure 4) is also informative. First, the performance of both the NN and MEHS deteriorates with increasing hail size. That, of course, is partially a consequence of the skewed nature of the distribution of hail size (Figure 1). Furthermore, the NN displays far less scatter about the horizontal line than MEHS does. As such, the NN’s predictions are more accurate than those of MEHS. Again, a scalar but quantitative reflection of these observations is offered by the $r^2$ of the residuals. Here a small $r^2$ is desirable, for that implies more accurate predictions. For the NN, the training and the validation $r^2$ are 0.0002, and 0.0017, respectively, while the same two quantities for MEHS are 0.21, and 0.13. Also, the regression fits to the NN data yield horizontal lines of slope zero for both the training and the validation sets. This is desirable because
it implies that the NN has correctly captured the underlying relation between the predictors and hail size. By contrast, the same fit for the MEHS data yields lines that have significant negative slopes. In other words, MEHS’s predictions not only deteriorate with the magnitude of the prediction (as do also the NN’s), they are also heavily biased toward larger estimates.

Continuing with performance issues, that of the classification networks is assessed in terms of ROC, discrimination, refinement, and attributes diagrams. The former requires the introduction of a probability threshold, and so, will be treated last. The discrimination diagrams for the three classes are displayed in Figure 5. It can be seen that the class 1 forecasts clearly discriminate between the three classes; the distribution of class 1 observations is peaked to the right, while those of the other two classes are either flat or peaked to the left. This is a desirable result, although ideally one would want one curve peaked to the right and two curves peaked to the left. By contrast, class 2 forecasts display little or no discriminatory capability. Although, the class 1 curve is peaked to the left (as desired), the curve that is peaked to the right is the class 3 curve, and not the class 2 curve. Finally, class 3 forecasts are quite discriminatory, but with an interesting twist; they derive their discriminatory capability not only from the identification of class 3 observations, but also (in fact, mostly) from the identification of observations that do not belong to class 3.

Several facets of the quality of the forecasts can be assessed through attributes diagrams. Figure 6 shows these diagrams for forecasts belonging to each of the three classes. It can be seen that the reliability of nearly all the forecasts is within statistical limits of perfect forecasts (i.e., the diagonal line). The error bars are 95% confidence intervals due to sampling. The horizontal line corresponds to forecasts that have no resolution, and the bisector of the angle formed by it and the diagonal marks the locus of forecasts with no skill (i.e., BSS=0). The shaded area defines forecasts that contribute positively to skill. Therefore, it can be seen that in addition to being highly reliable, all the forecasts also contribute positively to skill. This is true even for the class 2 forecasts for which almost no discriminatory capability exists (Figure 5, middle). The lack of discriminatory class 2 forecasts and their positive contribution to BSS may seem paradoxical; however, it must be noted that the vicinity of the reliability curve (dark circles) in Figure 5b to the region
of no skill, in conjunction with the size of the error bars, suggests that the contribution of the forecasts to skill may not be statistically significant.

Although the probabilities are all highly reliable, the range of the forecasts is quite varied. Class 1 forecasts span a relatively wide range of 10% to 90%, whereas class 2 forecasts are restricted to the range 20%-50%. This reflects the difficulty in predicting class 2 hail with a high degree of confidence. It is interesting that class 3 forecasts can reach probabilities nearing 100%, albeit rarely.

It is convenient to superimpose the refinement diagram upon the attributes diagram, the former labeled with squares in Figure 6. Evidently, the 3 forecast classes have distinct levels of refinement. The class 1 forecasts display a mild degree of the desired U-shaped pattern. Class 2 forecasts have an uncommon and undesirable bell-shaped pattern, indicating that most of the forecasts are in the vicinity of 20%. The highly left-peaked forecasts for class 3 suggest that the most common forecasts are at 0%. This is partially a consequence of the rarity of class 3 observations in the data.

Finally, the introduction of a probability threshold can dichotomize the forecasts and allow for the computation of ROC diagrams (Figure 7). These diagrams support the previous findings that class 3 forecasts appear to have the highest performance, followed by class 1, and class 2 forecasts, respectively. In fact, from the validation data (Figure 7b), one might suspect that class 2 forecasts are nearly random because the diagonal line goes through the corresponding error bars.

7 Summary and Discussion

Two neural networks have been developed for maximum hail-size prediction, one for estimating the size in some physical dimension, and a second for assessing the probability of belonging to one of three hail size classes. The parameters of the networks have been inferred via Bayesian methodology in order to alleviate the problem of overfitting. Four Doppler radar, and five environmental variables have been employed as inputs to the networks. Emphasis is placed on multidimensional measures of performance. It is shown that the network designed to predict maximum hail size outperforms the existing method in WSR88-D Hail Detection Algorithm. Since the existing method does not produce proba-
ibilities, it is not possible to compare the performance of the second network; however, it is found that the network-produced probabilities constitute high quality forecasts, with quality gauged in terms of reliability, refinement, and discrimination diagrams. Attributes diagrams suggest that the forecasts contribute positively to skill. The one possible exception is the forecasts for mid-size hail (∼ 40mm) which display no statistically significant skill. This is easy to understand, for discriminating between extreme events is a relatively simple task. In contrast, disambiguating intermediate events from the extreme ones is more difficult. It is expected that all three classes can be reasonably discriminated as more data becomes available.

To better understand the relation between the individual attributes (predictors) and hail size, it is useful to compute the corresponding linear correlation coefficients, r. The r between hail size and the four “radar attributes” (attributes 1-4 in Table 1) all have approximately equal correlation with hail size, namely 0.5 (standard error=0.04). The “environmental attributes” (attributes 5-9 in Table 1) have linear correlations that vary from 0.01 to 0.1, but are practically zero within the standard errors (0.05). Of course, as evidenced by the nonzero number of hidden nodes in the NNs, the true underlying relationship is nonlinear. In fact, the exclusion from the NN of the variables with a small correlation with hail size leads to inferior performance. This implies that in spite of their weak linear correlation with hail size, the environmental attributes do play a significant role in a nonlinear and interactive model (e.g., NN) for hail size prediction.

Furthermore, the r between the various attributes themselves is important in ascertaining the amount of collinearity among the inputs. Identifying collinear inputs and their exclusion as inputs to the NN can reduce the likelihood of overfitting the data. The most collinear pair of attributes is (5,6) with r = 0.83, followed by the pair (6,7) with r = 0.70. Neither of these r’s is sufficiently large to justify the exclusion of either member of either pair. As a result, no attributes were excluded as input nodes.

What remains to complete the augmentation of the Hail Detection Algorithm with neural networks is the development of the latter for assessing the probability of severe hail, regardless of size. Such a network would issue a probability for the existence of hail (within some time window) after which the networks outlined in this article would enter to provide information regarding the size of the forecast hail. The former network
is currently in development and will be discussed in a separate article.

8 Acknowledgements

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9 References


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Figure Captions

Figure 1. The distribution of reported hail size.

Figure 2. The training and validation mean square errors of the NN (square) and MEHS (circle) for four bootstrap trials.

Figure 3. Scatterplot of the actual vs. predicted hail size of the NN (left) and MEHS (right) for the training (dark dots) and validation (circles) data. Also shown are the corresponding regression fits.

Figure 4. Residual plot of the NN (left) and MEHS (right) for the training (dark dots) and validation (circles) data. Also shown are the corresponding regression fits.

Figure 5. Discrimination diagrams for class 1 (top), class 2 (middle), and class 3 (bottom) forecasts.

Figure 6. The attributes and refinement diagrams for the class 1 (top), class 2 (middle), and the class 3 (bottom) forecasts. Also shown are the 95% confidence intervals due to sampling. The curve consisting of the squares is the refinement diagram, the hashed region corresponds to forecasts that contribute to BSS, and the horizontal line defines forecasts with no resolution.

Figure 7. ROC diagrams for class 1, 2, and 3 forecasts, based on the training set (top) and the validation set (bottom). The error bars in the x and y directions are the one-standard deviation intervals based on bootstrapping.
Table 1: The nine predictors computed by HDA and SSAP.

<table>
<thead>
<tr>
<th>No.</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Cell-based vertically integrated liquid</td>
</tr>
<tr>
<td>2</td>
<td>Severe hail index</td>
</tr>
<tr>
<td>3</td>
<td>Storm-top divergence (delta-V in m/s)</td>
</tr>
<tr>
<td>4</td>
<td>Midaltitude rotational velocity (m/s)</td>
</tr>
<tr>
<td>5</td>
<td>Height of the wet-bulb zero (km MSL)</td>
</tr>
<tr>
<td>6</td>
<td>Height of the melting level (km MSL)</td>
</tr>
<tr>
<td>7</td>
<td>Vertically-integrated wet-bulb temp</td>
</tr>
<tr>
<td>8</td>
<td>Wind speed at the equilibrium level (m/s)</td>
</tr>
<tr>
<td>9</td>
<td>Storm-relative flow at $-20^\circ C$ level (m/s)</td>
</tr>
</tbody>
</table>
Table 2: Summary of the 81 storm cases analyzed.

<table>
<thead>
<tr>
<th>Region</th>
<th>No. days</th>
<th>No. hailstorms</th>
</tr>
</thead>
<tbody>
<tr>
<td>Western U.S.</td>
<td>13</td>
<td>21</td>
</tr>
<tr>
<td>High Plains</td>
<td>7</td>
<td>27</td>
</tr>
<tr>
<td>Southern Plains</td>
<td>22</td>
<td>115</td>
</tr>
<tr>
<td>Midwest</td>
<td>15</td>
<td>85</td>
</tr>
<tr>
<td>S.E. U.S.</td>
<td>19</td>
<td>94</td>
</tr>
<tr>
<td>N.E. U.S.</td>
<td>6</td>
<td>44</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td><strong>81</strong></td>
<td><strong>386</strong></td>
</tr>
<tr>
<td>Category</td>
<td>Scalar</td>
<td>Multi-dimensional</td>
</tr>
<tr>
<td>-------------------</td>
<td>---------------------------------------</td>
<td>--------------------------------------</td>
</tr>
<tr>
<td>Categorical</td>
<td>Heidke Skill Score</td>
<td>ROC Diagram</td>
</tr>
<tr>
<td>Probabilistic</td>
<td>Ranked Probability Score</td>
<td>Attributes Diagram</td>
</tr>
</tbody>
</table>

Table 3: Examples of four types of performance measures.