Forecasting Air Quality Time Series Using Deep Learning

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**Introduction**

Tropospheric ozone (O3) is a secondary pollutant formed by complex photo-chemical processes that impacts human health, plants, and structural materials. Over 21,000 premature deaths in Europe are attributed annually to O3 exposure (Amann et al., 2008) with over 1.1 million deaths worldwide - over 20% of all deaths attributed to respiratory diseases (Malley et al., 2017). In cases where O3 levels will exceed standards for long periods of time, air managers may issue air quality warnings and even limit industrial and vehicular activities (Kuhlbusch et al., 2014; Welch et al., 2005). Improving forecast accuracy provides planning and decision options that can impact receptor health and local economies.

In this paper, we train a deep learning model, consisting of a recurrent neural network (RNN) with long short term memory (LSTM), to accurately predict local 8 hr averaged (ave) O3 concentrations based on hourly air monitoring station measurements as a tool to forecast air pollution. RNNs are particularly well suited for air concentration prediction because they incorporate sequential history into the training and processing of input data. Air quality measurements are time series datasets in which the order of data is important. Troposhperic O3 was chosen as a parameter to forecast because of its complex formation processes and sources.

**Background**

Ozone as an Ambient Air Pollutant

The majority of tropospheric O3 is generated through anthropogenic sources (Lelieveld and Dentener, 2000; Cooper et al., 2006) attributed to the photo-disassociation of NO2 as shown below in the simplified reaction (Finlayson-Pitts and J.N. Pitts, 1993):

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|  | (1) |

where *M* is a stabilization molecule used during the intermediate formation between O and O2. In addition to nitrogen oxides (NOx's), VOCs, chlorine, solar radiation (SR), relative humidity (RH) and ambient temperature also impact tropospheric O3 formation (Song et al., 2011). Local concentrations of O3 are further influenced by weather patterns and terrain that disperse the pollutants, precursors, and byproducts (Beck et al., 1998). At night, O3 reacts with NO2 to form NO3 (nitrate radical) (Finlayson-Pitts and J.N. Pitts, 1993):

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|  | (2) |

The NO3 radicals react with NO2 to form dinitrogen pentoxide (N2O5) which in turn forms nitric acid (HNO3) through hydrolysis with water or aqueous particles (Song et al., 2011, Thornton et al., 2010). The acid is finally neutralized by ammonia (NH3) to complete the reaction chain (Brown and Stutz, 2012).

Additional contributions to tropospheric O3 concentrations come from the stratosphere-troposphere exchange (STE) of stratospheric ozone (Tarasick and Slater, 2008). The percentage of O3 provided by STE at surface levels range from 13% (Cooper et al., 2006) to over 42% (Lelieveld and Dentener, 2000) depending on area and conditions. With so many chemical and transport phenomena taking place throughout the day and night, modeling O3 becomes a very complex task even before local terrain, sources and weather patterns are incorporated (Bei et al., 2014). Nonetheless, predicting ambient O3 concentrations, and in particular, concentrations that may exceed air quality standards, is important for air managers and at-risk populations.

Forecasting Ozone with Machine Learning

Due to the formation process of O3, the actual concentration a local population is exposed to may have been generated from precursors emitted hundreds or even thousands of kilometers away (Glavas and Sazakli, 2011). Populations living in coastal regions may be exposed to pollutants generated locally but transformed and mixed with other precursors in circulating land-sea breezes (Freeman et al., 2017). Tropospheric O3 is therefore a more complex pollutant to estimate than primary pollutants such as sulfur dioxide (SO2) or carbon monoxide (CO).

Many studies have used supervised machine learning techniques, such as artificial neural networks (ANNs) to predict O3 time series concentrations (Wirtz et al., 2005; Ettouney et al., 2009; Kurt et al., 2008; Biancofiore et al., 2017). ANNs have been shown to provide better predictive results than linear models such as Multiple Linear Regression (MLR) and time series modeling such as Autoregressive integrated moving averages (ARIMA) (Prybutok et al., 2000; Zickus et al., 2002; Liu, 2007; Vlachogianni et al., 2011). Support vector machines (SVMs) have also been applied to O3 prediction scenarios with results that often outperform ANNs (Luna et al., 2014; Papaleonidas and Iliadis, 2013; Singh et al., 2013).

The benefits of using ANNs include not requiring a priori assumptions of the data used for training and not requiring weighting of initial inputs (Gardner and Dorling, 1998). In practice, dimensionality reduction is often used to remove inputs to the model that are not independent and identically distributed (IID) or offer little influence on the overall training. Principal Component Analysis (PCA) is often used to reduce the overall inputs to the model by removing transformed components, but provides little variability to the actual number of features required to train (Singh et al., 2013; Wang et al., 2015).

Most of the studies that use ANNs apply a single hidden-layer feed forward neural network (FFNN) architecture trained with meteorological and concentration data and generally have limited success for forecasting air quality. The canonical FFNN model consists of an input layer, a hidden layer and an output layer. Each layer is constructed from interlinked nodes that generates a value (usually between -1 and 1 or 0 and 1). The individual node model is shown in Figure 1.

Figure 1 here

The node sums the weighted inputs of the previous layer, sometimes with a bias, and transforms the combined sum with a non-linear activation function, *σ*. The node activation equation is given by

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where *w* is an array of weights for the connections between the previous layer and the current layer, *x* is a vector of input values from the previous layer, and *b* is a bias value. Common activation functions include the *sigmoid*, *tanh*, and *relu* functions. A general property for activation functions is that they normalize the output and have a continuous first order derivative that can be used during the back propagation training process (Goodfellow et al., 2016). The common activation functions mentioned earlier are shown in Table 1 along with their first order derivative and output range.

Table 1 here

More recent studies, however, looked at the limitations of FFNNs, namely the difficulty in choosing a suitable architecture and the tendency to over-fit the training data, leading to poor generalization, particularly in situations where limited labeled data is available (Lu and Wang, 2005; Papaleonidas and Iliadis, 2013). The predicted outputs in previous air quality forecast studies (Arhami et al., 2013) were based on continuous concentration values measured in parts per million (ppm) or μg/m3 from single stations. Achar et al. (2011) investigated the intervals between O3 exceedances and maxima of daily concentration levels instead of estimating real time values. Their study of interoccurrence between peaks was used to determine overall improvements to air quality trends over time as compared to predicting future air quality concentrations (Achcar et al., 2011).

For our validation case study area in Kuwait, several studies were completed that focused on ambient air quality and modeling using ANNs. Abdul-Wahab (2001) used 5 minute measurements of precursors CH4, Non-Methane Hydrocarbons (NMHCs), CO, CO2, Dust, NO, NO2, and NOx) and meteorological inputs (WS, WD, TEMP, RH, and Solar Radiation) from a mobile site in the Khaldiya residential area to estimate ozone and smog produced (SP) using a single hidden layer FFNN (Abdul-Wahab, 2001). Al-Alawi and Abdul-Wahab later enhanced their model by applying PCA to reduce the dimensionality of the input data (Al-Alawi et al., 2008). Ettouney et al. (2009) used the same inputs as Abdul-Wahab (replacing dust with Methanated Hydrocarbons) and two FFNNs to predict monthly O3 concentrations from the Jahra and Um Al Hayman stations. They suggested that O3 in Kuwait often comes from outside the local area via long range transport (Ettouney et al., 2009).

Deep Learning and Time Series

Studies in atmospheric sciences and O3 concentration predictions using Deep Learning (DL) have not been as common as single hidden layer ANNs. DL refers to the families of ANNs that have more than one hidden layer or use advanced architectures such as recurrent neural networks (RNNs) and convolutional neural networks (CNNs) (Goodfellow et al., 2016). Partially recurrent network models, such as the Elman Network (EN), were used with air station inputs to predict ground level concentrations of O3 (Biancofiore et al., 2015) and PM2.5 (Biancofiore et al., 2017). The feedback provides memory to the system when a single input set is fed into the system. Different ANN architectures are shown in Figure 2. The simple node in Figure 1 is redrawn as a circle for comparison. The arrows between layers represent synaptic weights that interconnect each node.

Figure 2 here

Implementations of procedures such as LSTM for RNNs allow network training to take place without having long term parameters “explode” or “vanish” as a result of multiple learning updates (Pascanu et al., 2013). LSTM was first introduced by Hochreiter and Schmidhuber in 1997 to overcome these training issues (Hochreiter and Schmidhuber, 1997). Gomez (2003) was one of the first researchers to use a single layer RNN to forecast maximum ozone concentrations in Austria (Gomez et al., 2003). His model utilized LSTM to outperform other architectures such as ENs. Noting the gap of years from Gomez et al. in 2003 and the work by Biancofirore et al. as recently as 2017 using ENs show that the complexity of preparing and training RNNs that use LSTM has kept researchers from using DL methods. DL has recently become popular for many applications due to improvements in training procedures and software libraries such as Theano (Theano Development Team, 2016) and Keras (Chollet et al., 2015) and Tensorflow (Abadi et al., 2015). These libraries have made implementing DL models easier, and therefore more accessible for researchers outside of the Machine Learning fields. A discussion of the theory of RNNs is presented in the next section.

**Theoretical Background**

Time series data

Air quality data are continuous, multi-variate time series where each reading constitutes a set measurement of time and the current reading is in some way related to the previous reading, and therefore dependent (Gheyas and Smith, 2011). Measured pollutants may be related through photochemical or pre-cursor dependencies, while meteorological conditions are limited by physical properties. Time series are often impacted by collinearity and non-stationarity that also violate independence assumptions and can make forecast modeling difficult (Gheyas and Smith, 2011). Autocorrelation of individual pollutants show different degrees of dependence to past values. Correlation coefficients were calculated using the equation

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|  | (4) |

where X is the input vector of a time step and *τ* is the lag (in hours). The correlogram was plotted based on lags up to 72 hr, as shown in Figure 3.

Figure 3 here

The parameters of Figure 3 show clear diurnal cycles, with O3 having very strong relational dependence every 24 hr, regardless of the time delay. In contrast, the dependency of NOx falls rapidly over time, despite peaking every 24 hr. Non-stationarity, collinearity, correlations, and other linear dependencies within data are easily handled by ANNs if enough training data and hidden nodes are provided (Goodfellow et al., 2016). More important to time series are the near term history associated with the previous time step. RNNs incorporate near term time steps by unfolding the inputs over the time sequence and sharing network weights throughout the time sequence. Additionally, the sequence fed to the RNN has fixed order, ensuring that for that individual observation, the sequence follows the order it appeared in, rather than randomly sampled as is the case for FFNN training (Elangasinghe et al., 2014). Previous models using ANNs could assume that some historical essence of the data was incorporated into the weights during updating as long as the training data was fed in temporal order and not shuffled as most categorical applications are (Bengio, 2012). Another way of handling sequential data is to use a time-delay neural network (TDNN). This type of architecture takes multiple time steps of data and feeds into the network at the input, using extensions of the input to represent previous states and become the system memory. TDNNs, in modern terminology, are called 1-dimensional CNNs (Goodfellow et al., 2016). These models were not considered in this study.

Recurrent Neural Networks

As mentioned previously, RNNs are well suited for multivariate time series data, with the ability to capture temporal dependencies over variable periods (Che et al., 2016). RNNs have been used in many time series applications including speech recognition (Graves et al., 2013), electricity load forecasting (Walid and Alamsyah, 2017) and air pollution (Gomez et al., 2003; Fan et al., 2017). RNNs use the same basic building blocks as FFNNs with the addition of the output fed back into the input. This time delay feedback provides a memory feature when sequential data is fed to the RNN. The RNN share the layer's weights as the input cycles through. In Figure 4, X is the input values, Y is the network output, and H is the hidden layers. An FFNN is included to compare the data ow over time. By maintaining sequential integrity, the RNN can identify long-term dependencies associated with the data, even if removed by several time steps. An RNN with one time step, or delay, is called an Elman Network (EN) and has been used successfully to predict air quality in previous studies (Biancofiore et al., 2015, 2017). The structure of the EN was shown in Figure 4b.

Fig 4 here

RNNs are trained using a modified version of the back propagation algorithm called back propagation through time (BPTT). While allowing the RNN to be trained over many different combinations of time, BPTT is vulnerable to vanishing gradients due to a large number of derivative passes, making an update very small and nearly impossible to learn correlations between remote events (Pascanu et al., 2013; Graves, 2013). Different approaches were tried to resolve these training challenges including the use of gated blocks to control network weight updates such as LSTM. LSTMs will be discussed in another section. While RNNs and LSTMs have been around for many years Hochreiter and Schmidhuber (1997), their use was limited until recently, in what Goodfellow et al. calls a “third wave of neural network research”. This period began in 2006 and continues to this day (Goodfellow et al., 2016). Like FFNN's, RNN's are trained on loss functions using optimizers to minimize the error. The loss function, or cost function, is the function that measures the error between the predicted output and the desired output (Goodfellow et al., 2016). In optimization theory, there are many loss functions that can be used including the Mean Square Error (MSE) and cross entropy (CE) functions being the most popular for machine learning applications. In our study, the MSE loss function was used instead of the CE function. Optimizers provide the method to minimize the loss function and include terms and parameters that determine the amount of incremental changes to the network weights during training. In addition to a learning rate, optimizer terms often include momentum and regularization. Momentum is used to speed up convergence and avoid local minima (Sutskever et al., 2013), while regularization describes terms that reduce generalization errors (Goodfellow et al., 2016). The most common optimizer used for the back propagation training algorithm used on most neural networks is stochastic gradient descent (SGD). The basic first order SGD equations with a classic momentum (CM) term is given as

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|  | (5) |

where *g* is the gradient update term, *μ* is the momentum factor (), *α* is the learning rate, and is the gradient of the loss function for a specific parameter, *θt*. The parameter is updated by

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|  | (6) |

Solving for the vanishing gradient

A major limitation of SGD for training very deep learning networks is the vanishing gradient problem, where the gradient update term becomes so small that no update takes place and the network parameters do not converge. Hinton et al. (2006) introduced greedy layer-wise pre-training in which a network was trained layer by layer and then integrated with SGD when compiled together (Hinton and Osindero, 2006). Since then, other first-order optimizers have been introduced that modify the SGD's basic algorithm by updating the learning rate and momentum terms during the training process (Sutskever et al., 2013). One such method was to apply a Nesterov accelerated gradient (NAG) (Nesterov, 1983) term to the SGD gradient update. The NAG update closely resembles the SGD update in eq 5 except for the addition of another momentum term in the parameter gradient.

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|  | (7) |

Other algorithms include the adaptive subgradient descent (AdaGrad) optimizer (Duchi et al., 2011), the root mean square propagation (RMSProp) optimizer (Tieleman and Hinton, 2012), the adaptive momentum (Adam) optimizer (Kingma and Ba, 2014), and the Nesterov adaptive momentum (Nadam) optimizer (Dozat, 2016). A summary of how these optimizers differ is shown in Table 2.

Table 2 here

After experimenting with all four of the optimizers in Table 2 and SGD, the Nadam optimizer proved to work the best with our study as described in the next section.

Long Short Term Memory

In order to preserve the memory of the data in the current state of the model, the RNN feeds parameters of its current state to the next state. This transfer can continue for multiple time steps and presented significant training challenges as mentioned earlier. The issue of vanishing gradients that took place during the BPTT updates was largely solved with the implementation of gating systems such as LSTM that allow nodes to forget or pass memory if it is not being used, thus preserving enough error to allow updates (Hochreiter and Schmidhuber, 1997). The LSTM uses a series of gates and feedback loops that are themselves trained on the input data as shown in Figure 5.

Figure 5 here

Each individual node in the LSTM acts like a standard FFNN node, similar to the one in Figure 1. The choice of activation function is another parameter to consider in the LSTM design. Common functions include the *sigmoid*, *tanh*, and *relu* functions as shown in Table 1. The difference to model at the input layer is that in addition to the observation data, X, additional input from the recurrent output, YR, representing a time delayed element of the network, is included for a composite input of

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|  | (8) |

The processed recurrent input, XR, feeds into several gates that allow the data to pass, represented by Φ in Figure 5. The weights that pass XR to the gate summations are trained as well. The use of LSTM in RNN architecture allows long term dependencies in data to be remembered within the model (Graves, 2013), a feature required when working with sequential series.

**Methodology**

Study area and data

For our validation case study area, we used datasets collected in the State of Kuwait. Kuwait is a small country of approximately 17,818 square km located on the northwest corner of the Persian Gulf, between longitudes 46:56o -48:37o East and latitudes 28:51o - 30:16o North with over 499 km of coastline (CIA 2015). The country is classified as a desert zone with the highest altitude reaching only 300 meters above sea level. In 2016, approximately 4.1 million people lived in Kuwait (CSB, 2016) with over 64% of its annual Gross Domestic Product (GDP) coming from the production of hydrocarbons (KAMCO, 2013). Over 98% of the population lives within 10 km of the coast and are subject to coastal effect winds, caused by the diurnal differential heating/cooling of the sea and land (Crosman and Horel, 2010; Cuxart et al., 2014). The land-sea breezes continuously shift direction and speed over the course of the day, recirculating pollution back and forth from land sources creating different zones of concentration mixing (Freeman et al., 2017). Fixed air monitoring stations are operated throughout the country near residential and industrial areas, but predominantly in the coastal zone areas (Freeman et al., 2017). For this study, data was collected from a station using OPSIS differential optical absorption spectroscopy (DOAS) analyzers (www.opsis.se) located near a local college as shown in Figure 6.

Figure 6 here

The location is centered between two major highways (5th and 6th Ring Highways) in a concentrated mixed commercial/residential area and north of the Kuwait International Airport. While not near the heavy refineries and industries in southern Kuwait, the site is impacted by land-sea breezes that recirculate emitted pollutants from throughout the Persian Gulf (Freeman et al., 2017). A data set from 1 Dec 2012 to 30 Sep 2014 was used for this study. Parameters available are shown in Table 3.

Table 3 here

The local O3 air quality standard is 75 ppb measured against an 8 hr moving average (KEPA, 2017). The DOAS station recorded O3 concentrations hourly, along with the other measured parameters. Measured units in μg/m3 were converted to ppb using the conversion formula

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|  | (9) |

where *C(ppb)* is the gas concentration in ppb, *C(μg/m3)* is the concentration in μg/m3, *R* is the ideal gas constant given as 8.3144 m3kPaK-1mol-1, MW is the molecular weight of the gas in g/mole (O3 = 48.01 g/mole), T is the ambient temperature in degrees Kelvin, and P is the atmospheric pressure at ground level in kPa. The station receives a prevailing wind from the northwest throughout the year as shown in Figure 7.

Figure 7 here

Seasonal effects are shown in bivariate polar plot in Figure 8. High O3 concentrations occur in the summer months from June to August, but come from the northeast, indicating the transport of pollutants from the coast. Not surprisingly, most of the compliance exceedances take place during this period.

Figure 8 here

Average hourly O3 and NOx concentrations are shown in Figure 9. The two variables are highly inverse correlated (R2 = -0.963) with common maxima/minima at 3:00 am, 6:00 am, 2:00 pm, and 10:00 pm. The raw hourly data is less correlated (R2 = -.576), but still higher than other variables. The NOx peaks correspond to rush hour traffic periods with winds blowing from the 6th and 7th Ring highways in the southwest. O3 levels peak in the afternoon, corresponding with solar radiation levels, but there is also a local maximum, or “morning bump” due to photolyzed chlorine ions reacting with N2O5 to form NO3 as part of the O3 formation cycle in eq 2 (Calvert et al., 2015). The formation of NO3 radicals was observed to be inversely related to O3 concentrations (Song et al., 2011).

Figure 9 here

Building the RNN

The RNN used in this study used the Keras (version 2.0.9) machine learning application programming interface (Chollet et al., 2015) with Theano back-end. Theano is a C++ library that allows mathematical expressions to be calculated symbolically and evaluated using datasets in matrices and arrays (Al-Rfou et al., 2016). The architecture used a single RNN layer with LSTM and a single output feed forward node. The output activation functions for both layers was the sigmoid function while the activation function for the recurrent nodes was the tanh function. The learning rate, α, was left at the default value of 0.002 (Chollet et al., 2015). Other Keras defaults included weight initialization (using a uniform distribution randomizer). Regulation was not used, although a dropout layer was included between the LSTM and the output layer.

Input Data preparation

Each available feature was compared to the maximum possible data range of 16,056 hourly measurements over the observation period. Gaps in data were assumed to be Missing Completely at Random (MCAR) and attributed to maintenance down-time, power failures, and storm related contamination (Le et al., 2007). Additionally, some data were clearly out of range or had negative readings (Junger and Ponce de Leon, 2015). Data recorded as a 0 was assumed to be censored and converted to the smallest recorded value within the data set of the individual parameter (Raa et al., 2015). Negative and missing data were converted to 0 and identified using a filter mask. Two different single imputation (SI) techniques were used based on the number of consecutive gaps in data. For gaps (g)< 8, the first and last measurement within the gap were used as a Bayesian estimator based on the previous observation to create a linear estimate of the missing data given by

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|  | (10) |

where *n* is the missing data point in sequence (), and

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|  | (11) |

For consecutive gaps > 8, the corresponding hourly measurement from the previous and preceding day was averaged.

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|  | (12) |

The value of 8 consecutive gaps was determined by comparing the root mean square error (RMSE) of the original data with data generated from the different statistical (imputed data) methods on artificially generated gaps (Junninen et al., 2004). The first method's error varied with gap size, while the second method had a higher, static error. The RMSE for O3 and NOx using the first method is shown in Figure 10 along with the intersection of the 2nd method.

Figure 10 here

Features with more than 50% missing data, like RH and Chlorine (Cl2), were discarded. The remaining variables from Table 3 had few missing data points, ranging from 41 missing points out of 16,053 (0.3%) for WS, WD and Temperature, to 137 missing points (0.9%) for NH3. The available data used for this study is larger than data sets used in other studies that had up to 16% missing data (Taspnar, 2015). Missing data adds noise to the training set and is often used to improve generalization and prevent overfitting the network to meet the training dataset. Adding dropouts, or intentional data removal, is often used during network training for this reason (Srivastava et al., 2014).

Cyclic and Continuous Data

WD and time of day were converted into representations that preserved their cyclic nature. Wind direction was converted into sine and cosine components (Arhami et al., 2013). Other parameters were transformed and scaled between values of 0 and 1 (Chatterjee et al., 2017) using the *MinMaxScaler* function in the Python Sci-Kit preprocessor library (Pedregosa et al., 2011). Overall, 25 features were prepared for initial training. These included the parameters measured in Table 3 with the addition of sine and cosine components for wind direction.

Feature selection using Decision Trees

Prior to training the RNN, features were reviewed to reduce input dimensionality by training multiple decision trees on the data sets and prioritizing features using the feature importance metric calculated during classifier training with the *DecisionTreeClassifier,* also in the Sci-kit library (Pedregosa et al., 2011). Decision trees and random forest classifiers have been used to reduce input dimensions for sensors (Cho and Kurup, 2011) and data classification competitions, outperforming other methods such as PCA and correlation filters (Silipo et al., 2014; Al-Alawi et al., 2008; Voukantsis et al., 2011). Decision trees are a supervised learning algorithm that recursively partition inputs into non-overlapping regions based on simple prediction models (Singh et al., 2013; Loh, 2011). Decision trees do not require intensive resources to train and evaluate, and keep their features, unlike PCA that transforms input variables into linear combinations based on the singular value decomposition (SVD)of the total data set's covariance matrix (Wang et al., 2016). While PCA is a form of unsupervised learning that allows dimensionality reduction by removing the number of transformed components fed to the input, decision trees identify which raw variables offer less impact so that they can be removed from the data collection stream. This can reduce future efforts required to collect, clean and prepare data sets.

Using PCA for model input also limits the inclusion of new data that may become available on real time systems. If a model is trained on transformed principal components only, any new data must also be transformed, thus changing the historical data set. Using a linear transformation method that only changes the individual observation and not the entire data set is more practical for time series applications where new data is expected to be added over time. Other feature reduction techniques, include evaluating contribution factors of weights to nodes to remove, or “prune” low value weights (de Oña and Garrida, 2013). This technique is not recommended for RNNs, especially with LSTMs, because of the way the weights are updated through different time steps. Alternative pruning techniques can be applied to RNNs to not only reduce inputs, but also reduce internal interconnections between the LSTM and look-back nodes (Narang, 2017). We did not explore internal pruning techniques in this paper. Instead, we evaluated the contribution of features externally and independent of the RNN using decision trees. Binary classification trees, a type of decision tree used for categorical separation, were trained to predict exceedances of 8 hr ave O3 over 1 hr to 12 hr horizons. Individual observations were first scaled using the *MinMaxScaler* function based on the equation

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where xmax and xmin are the maximum and minimum values in the data set respectively. It can be argued that this method also suffers from legacy biasing like PCA in that the system retains xmax and xmin in order to restore transformed data to original scale, similar to a key for encryption decoding. If new data is included that exceeds the xmax / xmin values, the data needs to be reprocessed using the new points. Since we are already working with an historical data set and not using real time data, there is no need to incorporate this issue. However, even if we were using real time data, we could safely assume that any value measured that exceeded xmax in our historical set was also an extreme point and could be accounted for in the model as a value > 1.

Eighty percent (80%) of the scaled data was divided into a training set with 20% reserved for testing. In larger data sets, a percentage of the total data is often reserved to allow parameter optimization prior to training in order to reduce time and system resources. Additionally, for FFNNs, training often take hundreds or thousands of epochs, where an epoch is a complete training cycle that uses all training sets. With our relatively small data set and fast training using the LSTM, reviewing system performance using the full training set did not take much time and therefore did not requiring reserving a subset.

Output exceedances were classified by the decision tree as either 0 (less than the exceedance standard of 75 ppb) or 1 (exceeds the standard). Overall accuracy of each horizon was measured using the accuracy score function in Sci-kit which calculated the standard error of exact matches of the observed output with the predicted (Raschka, 2016). Other classifiers in the Sci-Kit library were evaluated as well, including SVMs and Random Forest classifiers. The decision tree in classification mode using “gini” criteria to measure the data split at each decision node proved to be the most accurate. The importance of each feature was computed using the Sci-kit *feature\_importances\_* function that normalizes the total reduction of the criterion brought by individual features. Results of individual feature importance from classifying O3 exceedances at different horizons is shown in Figure 11. The computed values are unitless and displayed in relative importance to each other.

Figure 11 here

Output data preparation

The RNN output was trained to predict the 8 hr moving average of measured 1 hr O3. To predict future values, the calculated values were shifted in time based on the desired horizon so that input observations X(t = 0) was trained on Y (t = 12) if the prediction horizon was 12 hr. Output data was generated from 8 hr moving averaged O3 calculated from measured 1 hr O3 concentrations at each station. The first seven hours of both the input and output training data set was then discarded.

Tensor preparation for RNN input data

Data sets provided to the RNN were converted into 3 dimensional tensors based on the sample size of data. The sample size was based on the number of look-back elements within the RNN, as compared to an observation which represented one row of the original data set, *X*. The transformation of the original 2-dimensional data set *X* is illustrated in Figure 12 using Python notations. Assuming *X* is a data set of input data (for training or testing the RNN) with *n* observations and *p* variables, the total number of elements is the product of *n* and *p*, or 20 elements for the 5 x 4 data set in the figure. A tensor (*T*) is created with dimension (*s, l, p*) where *s* = # of samples given as *n* - *l*. The total number of elements within *T* is *s - l - p*. In the example of Figure 12, the dimensions of T are *s* = 5 - 2 = 3, *l* = 2, and *p* = 4.

Figure 12 here

**Analysis of Results**

Final parameter selection

Because of the Tensor formation for input, the actual number of samples provided for training and testing was based on the look ahead horizon and number of recurrent (look-back) units of the individual run. The farther out the prediction, the fewer samples were available because of the time shifting required. The total amount of samples available for training and testing could be calculated as total samples = (16,035 - *h*) where *h* is the prediction horizon (as an integer value > 1). The number of training epochs was limited after reviewing training error values up to 20 epochs for look-ahead horizons of 24 hr, 36 hr, and 48 hr as seen in Figure 13. An optimum number of 10 epochs was used for later model runs as it minimizes the training error without overfitting which begins to take place after 12 epochs, especially in Figure 13a.

Figure 13 here

Performance measures

Final parameter selection and performance were measured by Mean Absolute Error (MAE)

and Root Mean Square Error (RMSE). MAE and RMSE are widely used measures of continuous variables with RMSE criticized for over-biasing towards large errors (Chai and Draxler, 2014; Willmott and Matsuura, 2005). Both metrics were calculated for comparison, however MAE is used more often for descriptive analytics in this paper.

Impact of features and parameters on results

The network trained very well with all 25 input features from Table 3. An example of the predicted results compared with the observed measurements (8 hr ave O3) over a 24 hr horizon is shown in Figure 14.

Figure 14 here

The MAE for this scenario is 0.41 ppb during training, and 0.37 ppb during testing. The residuals of this scenario are shown in Figure 15 where they show a Normal distribution tendency (skewness = 0.411, kurtosis = 3.94, where Normal is 0 and 3, respectively) with a positive bias given by the distribution mean of 1.632 ppb. This is consistent with Figure 14 that shows the model slightly under-predicting.

Figure 15 here

The results of the decision tree analysis in Figure 11 showed that many features could be removed without impacting network performance. Features were removed based on the order of least importance in groups of 5 until the most prominent feature remained. The results in Figure 16 show that overall training error improves with fewer inputs.

Figure 16 here

Based on the training error curves in Figure 16, the 5 feature data set was used for evaluation because it provided stable error over the prediction horizons of interest. The features used were (in order of importance) 8 hr ave O3 in ppb, 1 hr O3 in ppb, SR, the cosine of WD, and CH4. Parameter sensitivity analysis was performed on a model with default values shown in Table 4.

Table 4 here

All other parameters were held constant as an individual parameter was varied. The prediction horizon value of 24 hr was held constant throughout all runs shown in Figure 17. In all cases, the error measurements, MAE and RMSE showed similar forms, despite the RMSE having a consistently higher value, as expected, and as seen in other research (Singh et al., 2012). The prediction horizon of the model using the 5 feature data set is shown in Figure 17.

Figure 17 here

As the prediction extends further into the future (> 80 hr), the training error climbs rapidly, while the test error appears to level off. The model began to overfit by this point and predictions past that range were considered to be unreliable. The results in Figure 18a show training error for different prediction horizons over several parameters. The parameter that influences the model performance the most is the number of look-back nodes in relation to the prediction horizon. A horizon value of +2 provides the lowest errors, while adding additional nodes increases the model complexity and makes training more challenging. Samples/batches in Figure 18b show relatively little error variance until many samples are included (>75). While more samples per batch are preferred to reduce training time, too many create bias in the loss function as the overall average of each sample reduces chances for updates. The number of recurrent, or look back nodes, in relation to the prediction horizon was considered in Figure 18b. Both training and test results are minimized at 26, the horizon value + 2. This was consistent with other horizon prediction values such as 36 and 48. As more look back nodes are added, the error increases. Finally, the use of dropout is recommended to improve generalization of the model and reduce overfitting (Gal and Ghahramani, 2016). For this model, dropout was applied only between the output of the LSTM layer and the FF output layer. The error shows reasonably good optimization at around 0.2. The errors level out at higher rates at around 0.35. The default values were the optimum parameters based on the results shown in Figure 18.

Figure 18 here

An LSTM model has many more variables that can be optimized compared to other models. The RNN used in this study with 25 input features had 16,485 update-able parameters, of which 16,432 were in the LSTM layer alone. As a comparison, an FFNN with 3 hidden layers (5 layers total), bias on all layers, and the same 25 feature inputs, had only 2,107 parameters. Nonlinearities are further introduced during the training phase in which the derivative of the activation function for each layer is used to influence the weight updates. It is therefore difficult to fully explain the mechanisms driving the output results of complex DL models. In order to insure the model is working, the output must be compared with known results and parameters adjusted to optimize performance.

Comparison to previous studies

Previous studies mentioned in Section 1 used RMSE and other error measurement methods than MAE. The 3 studies that used RMSE were compared to our results with an LSTM network. Luna et al. (2014) used SVMs and FFNNs (Luna et al., 2014). Feng et al. (2011) used a a multi-layered system that included an SVM and a genetic algorithm stabilized FFNN (Feng et al., 2011). Wang and Lu used an FFNN with a particle swarm optimization (Wang and Lu, 2006). All studies, except Gomez (2003), used PCA to pre-process the data. Comparing the results of the RNN to these previous studies gives an initial impression that the RNN has an order of magnitude improvement over the best FFNN or SVM models as shown in Table 5.

Table 5 here

The results cannot however be directly compared because they were based on different data sets. While the other studies used complex and hybrid architectures along with complicated pre-processing, the RNN model preprocessing was very simple after the features were prioritized using a decision tree. The RNN and LSTM are themselves complex algorithms with many internal parameters that undergo training and updates.

Comparison to different models

In order to evaluate how the RNN model compared to other forecasting models using the same data set, a comparison was made for a 24 hr prediction using a FFNN with three hidden layers, and an ARIMA model. The FFNN was built with the Keras library using relu activation functions in the hidden layers and sigmoid function for the output. The inputs included all the parameters from Table 3 and the model was allowed to train for 1,000 epochs. The number of nodes in the hidden layer was based on the estimate # of hidden nodes = (*SF* \* # of input nodes) + # of output nodes where *SF* is a scaling factor between 0.5 and 1 (Papaleonidas and Iliadis, 2013). For an *SF* of 0.75, the number of nodes is 20 nodes in each hidden layer. An MSE loss function and Nadam optimizer were also used to build the model. The ARIMA model was built using the *arima* function in R and fitted on the 8 hr ave O3 only. The formatting parameters, *p*, *d*, and *q* were automatically estimated using the *auto.arima* function as 3, 0, and 0, respectively (Hyndman and Athanasopolous, 2013). The results are shown in Figure 19.

Figure 19 here

**Conclusions and recommendations**

This is one of the first papers that employ advanced Deep Learning techniques in the prediction of air quality time series events. This new methodology produced very good results using our validation data set. A recurrent neural network with LSTM was trained on time series air pollutant and weather data from an air monitoring station in Kuwait to predict 8 hr ave O3 over different prediction horizons. Missing data and censored data were replaced using a first-order imputation technique that accounted for sequential influence of previous readings for small gaps (< 8 consecutive gaps) and seasonal effects for larger gaps. A decision tree was used to prioritize the most influential features for training by categorizing pollution exceedances using the input parameters.

Prioritizing and removing less important features allows for real-time observations to be fed into the model without transforming large blocks of data as is required when using principal components or wavelets. New observations need only be scaled by normalizing or standardizing with the scaling values calculated from historical data sets.

A sensitivity analysis of key parameters showed that the network could be tuned for optimal performance. Measurement of the performance, in terms of observed and predicted results, were consistent in form, but the RMSE was always biased higher than the MAE measurement. Either measurement would have produced the same conclusions based on observation of local minima and maxima regardless of the error value. Error increased with the complexity of the network, even with reduced features. This “Curse of Dimensionality” led to overfitting of the model, reducing the ability to generalize if new data sets were introduced. Slight overfitting is not a problem for time series data that follow predictable cyclic patterns and the main output product of interest goes higher than a set limit.

While the results cannot be directly compared to other studies because different data sets were used, the results should not be dismissed either. Comparing the same data set results to other common forecasting models such as ARIMA and a multi-layer FFNN shows that the RNN does perform significantly better. The complexity of RNN implementation has been dramatically reduced with the use of the Keras developmental library, allowing non-computer scientists the ability to use DL without the coding overhead. The LSTM model provided very good results for this case and can be applied to other environmental time series challenges such as forecasting wide area pollution exceedances from multiple stations and multiple pollutants. LSTMs could also be effective in predicting individual source emissions or modeling source apportionment under different criteria.

Reducing features and optimizing parameters assisted with lowering error of both training and test sets. Initial runs using local data showed excellent results compared to performance from FFNNs, even with the inclusion of complex pre-processing of input data and architecture of the model. The relative ease of model structure in the programming code is misleading though. The Keras and Theano libraries are some of the most advanced, validated, and complex libraries available in the research community. The underlying errors within the model implementation may not be resolved or even quantified. However, they are still useful tools for rapid prototyping and architecture validation. Using the data sets of the sources listed in Table 5 with our RNN would be a more direct way to prove which method works better. Further investigations will target multiple station influences on local concentration prediction, as well as how imputation techniques for missing data impacts overall prediction accuracy.

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Table 1: Common activation functions

|  |  |  |  |
| --- | --- | --- | --- |
| **Name** | **Equation** | **1st derivative** | **Output range** |
| *sigmoid* |  |  |  |
| *tanh* |  |  | -1, 1 |
| *relu* |  |  |  |

Table 2: Enhanced first order optimizers used for DL

|  |  |  |
| --- | --- | --- |
| **Optimizer** | **Description summary** | **Source** |
| AdaGrad | Divides the learning rate, α, by the L2 norm | Duchi (2011) |
| RMSProp | Divides gradient by a running average of its recent magnitude | Tieleman (2012) |
| Adam | Combines CM with RMSProp | Kingman (2014) |
| Nadam | Combines NAG with RMSProp | Dozat (2016) |

Table 3: Chemical and meteorological parameters captured at the AMS

|  |  |
| --- | --- |
| **Chemical Analytes** | **Meteorological** |
| Nitrous oxide (NO) | Wind direction (WD) |
| Ammonia (NH3) | Wind speed (WS) |
| Ozone (O3) | Temperature |
| Sulfur dioxide (SO2) | Atmospheric pressure (ATP) |
| Formaldehyde (CH2O) | Solar radiation (SR) |
| Nitrogen dioxide (NO2) | Relative humidity (RH) |
| Benzene (C6H6) |  |
| Toluene (C7H8) |  |
| p-Xylene (C8H10) |  |
| m Xylene (C8H10) |  |
| 1,2,3-trimethylbenzene (C6H3(CH3)3) |  |
| o-Xylene (C8H10) |  |
| Ethylene glycol tertiary butyl ether (ETB) (C6H14O2) | |
| Styrene (C8H8) |  |
| Chlorine (Cl2) |  |
| Carbon dioxide (CO2) |  |
| Methane (CH4) |  |
| Hydrogen sulfide (H2S) |  |
| Carbon monoxide (CO) |  |

Table 4: Default values for parameter sensitivity analysis.

|  |  |
| --- | --- |
| **Parameter** | **Default Value** |
| Input features | 26 |
| Prediction horizon (hr) | 24 |
| Look back nodes | 26 |
| Samples/batch | 72 |
| Dropout factor | 0.2 |

Table 5: Comparison of RNN test data results to previously published results.

|  |  |  |  |
| --- | --- | --- | --- |
| **Source** | **Prediction horizon** | **Results (RMSE)** | **RNN (RMSE)** |
| Luna et al. (2014) | 1 hr | 6.3-12.3 | 0.8 |
| Feng et al. (2011) | 12 hr | 5.5-86.9 | 1.5 |
| Wang and Lu (2006) | 24 hr | 7.9-11.2 | 2.5 |
| Gomez et al. (2003) | 24 hr | 6.9-9.9 |

Table 6: Comparison of different forecasting errors over a 24 hr period.

|  |  |  |  |
| --- | --- | --- | --- |
|  | **RNN** | **FFNN** | **ARIMA** |
| MAE (ppb) | 0.235 | 15.28 | 23.574 |