A Machine Learning Artificial Neural Network Calibration of the Strong-Line Oxygen Abundance

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ABSTRACT

The H\textsc{ii} region oxygen abundance is a key observable for studying chemical properties of galaxies. Deriving oxygen abundances using optical spectra often relies on empirical strong-line calibrations calibrated to the direct method. Existing calibrations usually adopt linear or polynomial functions to describe the non-linear relationships between strong line ratios and $T_e$ oxygen abundances. Here, I explore the possibility of using an artificial neural network model to construct a non-linear strong-line calibration. Using about 950 literature H\textsc{ii} region spectra with auroral line detections, I build multi-layer perceptron models under the machine learning framework of training and testing. I show that complex models, like the neural network, are preferred at the current sample size and can better predict oxygen abundance than simple linear models. I demonstrate that the new calibration can reproduce metallicity gradients in nearby galaxies and the mass-metallicity relationship. Finally, I discuss the prospects of developing new neural network calibrations using forthcoming large samples of H\textsc{ii} region and also the challenges faced.

Key words: ISM: abundances – H\textsc{ii} regions – galaxies: ISM – methods: data analysis

1 INTRODUCTION

The H\textsc{ii} region oxygen abundance, a proxy for the metallicity of the interstellar medium (ISM), is a key quantity for studying chemical evolution of galaxies. Oxygen is the most common heavy element in the ISM (after helium and lithium). Together with other relatively abundant elements (e.g. nitrogen, sulfur), they produce a wealth of emission lines in the optical wavelength which are useful for constraining the ISM chemical abundances. Observing these optical emission lines has allowed us to significantly advance our understanding of fundamental chemical characteristics of star-forming galaxies, such as the oxygen abundance gradient (e.g. Searle 1971; Zaritsky et al. 1994; Sánchez et al. 2014; Ho et al. 2015) and the mass-metallicity relationship (e.g. Lequeux et al. 1979; Tremonti et al. 2004; Zahid et al. 2013).

Although observing emission lines from H\textsc{ii} regions is relatively straightforward, deriving accurate oxygen abundances is non-trivial. A variety of methods is available in the literature (see reviews by Stasińska 2004 and Peimbert et al. 2017); however, large systematic differences between different methods exist and the disagreement remains unresolved (Kewley & Ellison 2008). One of the more accurate ways is through measuring the electron temperature ($T_e$) and density ($n_e$) of the line-emitting gas (see Pérez-Montero 2017 for a tutorial). This method, known as the direct method, is built on the physical basis that the emissivity of collisionally excited lines (CELs) depends strongly on $T_e$. When $T_e$ and $n_e$ are known, ionic abundances can be inferred from CEL to hydrogen recombination line ratios. The total elemental abundance can subsequently be derived by accounting for unseen stages of ionisation. In this method, $n_e$ can be easily determined using the density-sensitive doublets [O\textsc{iii}]λ4363, 3729 and [S\textsc{ii}]λ6717, 6731. However, constraining $T_e$ requires detecting auroral lines originating from transitions between upper energy levels, e.g. [O\textsc{iii}]λ4363, [N\textsc{ii}]λ4575. A key limitation of the direct method comes from the faintness of the auroral lines, typically 2 orders of magnitude dimmer than the H\textsc{$\beta$} line.

It is worth noting that the direct method is not perfect and has an important caveat. Oxygen abundances estimated base on recombination lines (RLs) are almost always higher than those from the CELs, by about 0.2 to 0.3 dex (e.g. Peimbert 2003; Esteban et al. 2002, 2009; García-Rojas & Esteban 2007). One possible origin of this so-called “abundance discrepancy factor” problem is the presence of significant temperature, density, and possibly chemical com-
position fluctuation inside nebulae, which affect the direct method more than the RL method. The CELs are expected to originate from relatively low metallicity, high $T_e$, and low density regions inside a nebula, leading to an underestimate of the overall metallicity (see Peimbert et al. 2017 for a review). Recent spectroscopic studies of blue and red supergiant stars in nearby galaxies, however, report stellar oxygen abundances more consistent with gas-phase abundances of giant stars in nearby galaxies, however, report stellar oxygen abundances more consistent with gas-phase abundances measured from the direct method than the RLs (Bresolin et al. 2016 and references therein).

The faintness of the auroral lines severely limits its practical use when observations do not achieve excellent signal-to-noise ratios. Empirical strong-line calibrations are developed and have gained popularity among those using shallow (spectroscopic survey) data to study chemical properties of galaxies. Strong line indexes sensitive to metallicity, such as $R_{23}$ (Pilyugin 2001; Pilyugin & Thuan 2005), O3N2$^2$, N2$^3$ (Pettini & Pagel 2004; Marino et al. 2013), O32$^4$ (Maiolino et al. 2008; Curti et al. 2017), have been calibrated to the $T_e$ metallicity. In general, $T_e$ metallicities derived from high quality H\textsc{ii} spectra (or in some cases integrated galaxy spectra) are plotted against a selected strong line index. A simple linear or polynomial function is fit to the data and then reported as an empirical calibration.

The simplest cases are the linear calibrations of the O3N2 and N2 indexes. For the $R_{23}$ diagnostic, more complex functions are needed because the $R_{23}$ index is known to be double valued in metallicity (e.g. Pagel et al. 1979, 1980; Kobulnicky & Kewley 2004). In this situation, either two functions are fit separately to the upper and lower branches or a high-order polynomial is adopted. Photoionisation models, however, suggest that all these indexes have secondary dependency with the ionisation parameter (e.g. Kewley & Dopita 2002; Dopita et al. 2013). It is likely that, by including more line ratios, the calibration dispersion can be reduced.

It is non-trivial to include additional line ratios into the calibrations and find suitable analytical formulae. Steps moving from 1- or 2-dimensional calibrations (where one or two line ratios are used) to high-dimensional calibrations (where multiple line ratios are fit simultaneously) have been taken in various work (e.g. Maiolino et al. 2008; Curti et al. 2017). A particularly efficient approach was taken by Pilyugin & Grebel (2016). Pilyugin & Grebel recognised that the metallicity-dependent index N2 exhibits a secondary correlation with the excitation parameter $P$, particularly at high $T_e$ (low metallicities). By carefully designing the fitting formulae, Pilyugin & Grebel (2016) were able to develop two 3-dimensional calibrations: an $R$ calibration using [O\textsc{ii}], [O\textsc{iii}] and [N\textsc{ii}], and an $S$ calibration using [O\textsc{iii}], [N\textsc{ii}] and [S\textsc{ii}].

Another shortcoming of fitting one index at a time is that different indexes are sensitive to metallicity in different ranges, and the scatter could also be different. When multiple diagnostic indexes are available in a spectrum, different metallicities can be reported by using different indexes (and calibrations). A practical approach is to combine the different metallicities by taking the mean or minimising the $\chi^2$ in the space defined by the selected diagnostics (Curti et al. 2017; Maiolino et al. 2008).

Calibrating strong-line to $T_e$ metallicities is fundamentally searching for a function that can map the measured line ratios to $T_e$ metallicities using a set of well-characterised sample. This function is likely to have a high dimensionality (e.g. Vogt et al. 2014). The linear or polynomial fits in the literature describe only projections of this complex function onto certain index space. Here, I present an exploratory work of using a neural network model to recover the underlying function. The main goal of this work is to examine the possibility of fitting an arbitrarily complex, non-linear model to existing data (approximately 950 H\textsc{ii} regions) and improving the calibration accuracy by directly “learning” from the data. By performing supervised learning under a machine learning framework, I will “train” a strong line calibration that is data-driven and has no built-in physics. I will show that my first-try model can deliver reasonably good performance, even exceeding most popular literature strong-line calibrations. I will demonstrate that the neural network model can capture line ratios that are known to be sensitive to metallicity in photoionisation models, and successfully reproduce the well-characterised metallicity gradient and mass-metallicity relationship. I will end the article by discussing the prospect of this new approach and some challenges faced ahead.

Through out the paper, the following convention is followed: \[[\text{O}\text{ii}]=\text{[O\text{ii}]}\lambda\lambda6583,3729, \text{[O}\text{iii}]=\text{[O\text{iii}]}\lambda\lambda4959,5007, \text{[N}\text{ii}]=\text{[N\text{ii}]}\lambda\lambda6583, \text{[S}\text{ii}]=\text{[S\text{ii}]}\lambda\lambda6716,6731\]. The best-fit neural network model and corresponding PYTHON wrapper script is publicly available at https://github.com/

\footnote{P \equiv [\text{O}\text{iii}]\lambda4959,5007/([\text{O}\text{ii}]+[\text{O}\text{iii}])}

Figure 1. $T_e$ oxygen abundance distributions of different samples. Literature H\textsc{ii} region data compiled by Pilyugin & Grebel (2016, P16) are shown in yellow. Data from the CHAOS Project are shown in blue (Berg et al. 2015, CHAOS I), green (Croxall et al. 2015, CHAOS II), and red (Croxall et al. 2016, CHAOS III). The black solid line indicates the working sample adopted in this work.

\begin{itemize}
  \item No. of regions
  \item $T_e$ + log(O/H)
\end{itemize}
2 METHODS: COMBINING LITERATURE AURORAL LINE DATA AND A SIMPLE ARTIFICIAL NEURAL NETWORK

2.1 Sample

The H II region sample is drawn from two sources, containing originally 1009 oxygen abundances estimated based on auroral line(s). A sub-sample of 956 regions forms the final sample for this study.

The first source is 858 literature H II region data compiled first by Pilyugin et al. (2012) and Pilyugin & Grebel (2016). An updated version of this literature compilation is adopted here. The sample contains 70 regions where the [O III] λ 4363 line is detected and 151 regions the [N II] λ 6585 is detected. A small fraction of the sample has both auroral lines 90. The Te oxygen abundances are derived following the prescription described in Pilyugin et al. (2012) and assuming the commonly adopted t2 – t3 scaling relationship (Campbell et al. 1986; Garnett 1992). When both auroral lines are detected which yields two estimates of the oxygen abundance, the average is adopted. The second source comes from the on-going CHEmical Abundances Of Spirals (CHAOS) survey conducted with the Multi-Object Double Spectrograph (MODS) on the Large Binocular Telescope (LBT). In total, 151 H II regions are drawn from the three CHAOS publications, 45 from Berg et al. (2015, CHAOS I), 29 from Croxall et al. (2015, CHAOS II) and 77 from Croxall et al. (2016, CHAOS III). The oxygen abundances tabulated in their Tables 5, 5, and 4 are adopted, respectively.

Figure 1 presents the sample distributions. The P16 sample contains mostly low-metallicity regions that produce bright [O III] λ 4363. The CHAOS survey fills in the high-metallicity regime by virtue of the sensitivity of MODS and its capability to probe multiple auroral lines simultaneously ([O III], [N II], [S II], [O II] and [S I]).

To arrive at the final, working sample (956 regions), an abundance cutoff of 7.4 is imposed to remove the low low-metallicity tail. This is to avoid the calibration being affected by the sparse sampling in a relatively large metallicity space.

Calibrate strong-line abundance using ANN

2.1.1 Bootstrapping

The working sample does not uniformly cover the metallicity range of [7.4, 8.9], despite removing the low metallicity tail and including the high-metallicity CHAOS regions. To avoid the fitting being biased by the non-uniform distribution, I bootstrap the working sample to generate uniformly distributed samples.

The (non-uniform) working sample is first sorted and grouped in bins of approximately 30 regions. In each bin, new samples are drawn randomly (with replacement) and added to the bin such that the new sample number is proportional to the width of the bin. This re-sampling is done for every bin, except for the reference bin where the bin width is the narrowest. The reference bin defines the targeted bin-width to sample number ratio. The bootstrapped samples, each containing 2516 regions, are adopted to fit neural network models and quantify their performance. Performing analysis on the bootstrapped samples is equivalent to imposing higher weights on regions having high or low metallicities.

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2.2 Multi-layer perceptron

I build a simple, feed-forward artificial neural network using the multi-layer perceptron (MLP) framework. A schematic of the network architecture is shown in Figure 3. The MLP is constructed using scikit-learn\(^6\) (Pedregosa et al. 2011) version 0.19.2 in PYTHON3. The fully-connected MLP has three components: input layer, hidden layer(s), and output layer.

The input layer ingests “features”, i.e. logarithmic line ratios, from the training data. The 16 features (based on strong forbidden lines) are listed in Table 1. Here, all the line fluxes are normalised to H β and the logarithmic line ratios are adopted as the input features. To optimise the MLP performance, the 16 inputs are first passed through standard scaler functions such that the means are zero and the standard deviations are unity.

The 16 features are selected by taking possible arithmetic combinations of line ratios, i.e. single line, ratio of line pairs and sum of line pairs. Some of the features are line ratios well-known to be sensitive to metallicity. For example, the [N II], [N II]/[O II], and [N II]/[O III] are popular diagnostics (e.g. Kewley & Dopita 2002; Pettini & Pagel 2004). The combination of [S II]/[N II] and [N II] is also shown to be a robust metallicity indicator (Dopita et al. 2016). [O III]/[O II] is known to be sensitive to both ionisation parameter and metallicity (e.g. Kewley & Dopita 2002). [O I] + [O II], or equivalently R23, is yet another widely adopted metallicity indicator (Pagel et al. 1979). The rest of the features are not traditionally adopted in metallicity diagnostics. These line ratios could be sensitive to other physical properties of the ISM which may help improve the prediction of metallicity. The intention is to test if by non-linearly combining

Figure 2 shows where the working sample lies in the BPT line ratio space (Baldwin et al. 1981).
In the hidden layers, each neuron is a simple perceptron with the output being passed through an activator function. Each perceptron performs a simple calculation

\[ p(x) = \sum_i w_i x_i + b \]

where \( w_i \) and \( b \) (i.e. weight and bias) are the model variables and \( x_i \) are input from the previous layer. The rectified linear unit is adopted as the activator function that adds nonlinearity to the model. That is:

\[ A(x) = \max(0, x) \]  

In other words, each neuron in the hidden layers ingests outputs of the previous layer, \( x_i \), and performs the calculation

\[ A(p(x)) = \max(0, \sum_i w_i x_i + b) \]  

Two models are explored below: one hidden layer with \( n_1 \) neurons and two hidden layers with \( (n_1, n_2) \) neurons. The final, best model has only one hidden layer.

The output layer comprises of one single perceptron with no activator function attached. The perceptron in the output layer is used to predict the targeted unknown quantity, i.e. oxygen abundance.

The MLP framework described above is controlled by two hyper-parameters. The first parameter is the number of neurons in the hidden layers, \( n_1 \) and \( n_2 \). A larger number of neurons in the hidden layers allows the model a higher degree of complexity. Since complex models are prone to over-fitting the data, a second parameter \( \alpha \), the L2 penalty parameter (for regularisation), is introduced. At a fixed \( (n_1, n_2) \), \( \alpha \) of zero imposes no prior on all the possible models under the MLP configuration. A non-zero, positive \( \alpha \) penalises the chi-square by the sum of the weight squares (multiplied by \( \alpha \)), effectively guiding the fitting towards models with neuron weights closer to zero.

2.3 Train and test the neural network

Training the MLP network was done using the standard solver implemented in scikit-learn. The weights of the perceptrons are initialised randomly, and then a limited-memory version of the Broyden-Fletcher-Goldfarb-Shanno algorithm was adopted to minimise the loss function. The minimisation terminated when the the loss function did not improve by 10\(^{-4}\) for two consecutive iterations. Since the training sample is moderate in size, all the training sample was used simultaneously without performing any stochastic minimisation using mini-batches.

2.3.1 Hyper-parameter tuning

In general, hyper-parameter tuning is required to avoid overtraining a neural network, i.e. using a model too complex for the available data, resulting in the model to “remember the answers by-heart”. This problem is usually avoided by withholding a part of the data to test the model performance, and tuning the complexity of the model until the model performs equally well on data that it has and has not seen (i.e. training and testing samples). Below, I systematically investigate how the two hyper-parameters affect the network performance. This investigation will be done at different sample sizes to further understand whether more complex models can be adopted with increasing sample sizes.

For a given combination of hyper-parameters and train-sample size, a training sample is randomly drawn from the working sample and the remaining data are used for testing. Both the training and testing samples are bootstrapped (separately) to avoid the non-uniformity discussed in Section 2.1.1. After the model is trained, the standard deviation of the residuals, i.e. predicted versus \( T_e \) metallicities, is adopted to quantify the model performance. Both the residuals of the training and testing sets are calculated to quantify the in- and out-of-sample performance, respectively.

To avoid my analysis being biased by the random components described above, i.e. the random weight initiali-
poor performance. At each sample size, there is a family of models that can perform equally well. With increasing sample size, the achievable best-performance improves. That is, with more H\textsc{ii} regions complex models can be developed, which achieve better out-of-sample performance.

To avoid over-fitting the data, the hyper-parameters need to be chosen to deliver comparable in-and out-of-sample performance with the latter being as good as possible. This can be done by comparing Figures 4 and 5. At a given sample size, the appropriate model complexities can be achieved by different combinations of (log $\alpha$, $n_1$), networks with fewer neurons and less regularisation (smaller $\alpha$ and $n_1$) or with more neurons and more regularisation (larger $\alpha$ and $n_1$). The parameter log $\alpha$ of approximately 1.0 appears to avoid over-training over a wide range of $n_1$.

Finally, it is worth noting that the $n_1 = 1$ models are not optimal in all five sample sizes. Even with little regularisation, the single neuron model is still too simple to fully extract information from the data. The performance difference between $n_1 = 1$ and $n_1 > 1$ models is increasingly more obvious with increasing sample size. The single neuron model describes essentially linear combinations of the 16 logarithmic line ratios. Some existing strong line calibrations only fit linear combinations of logarithmic line ratios to the data, e.g. using the N2 or O3N2 indexes. These calibrations are similar to the single neuron models. This suggests that properly trained MLP models with more than one neuron will out-perform these literature calibrations. In other words, given the available literature samples complex models utilising multiple line ratios could better calibrate strong line metallicity than those using only one or two line ratio pairs.

2.3.3 Two hidden layers

The performance of a single hidden-layer network saturates at large $n_1$. It is worth exploring whether including a second hidden-layer could further increase the network performance. Here, I fix $n_1$ to 16 and systematically explore the network performance at different $n_2$ and $\alpha$. This exploration is done only for the 80%/20% train/test split. As before, 100 models with different random states are trained for each grid.
Using the best MLP model, I perturb one feature at a time while keeping the other features unchanged and compute the changes in the output metallicity. The small perturbation is set to 3% of the standard deviation of the input feature of the working sample. The sensitivity analysis can be achieved by computing the partial derivative of the output with respect to the input features.

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will report the median value of the predictions from the 100 model realizations. The data points are the working sample described in Section 2.1. The error bars are the standard deviations of the 100 predictions from the 100 model realizations. The dashed line is the one-to-one line.

In Figure 7, I compare input $T_e$ metallicities with those predicted by the MLP model from strong lines. The error bars show the 1σ standard deviations of the 100 predictions. In general, the spread in metallicity due to the random components in the fitting process is much smaller than the dispersion of the calibration. Figure 7 suggests that the best MLP model performs reasonably well over a wide range of metallicity.

I have demonstrated how the hyper-parameters and sample size affect the choice of model complexity, and showed that one hidden layer is sufficient for the current sample size. Here, I train the final MLP model using all the HII region sample, without withholding any data for testing. I adopt the hyper-parameters $(\log \alpha, n_1, n_2) = (1.2, 10, 0)$. This parameter combination delivers in- and out-of-sample performance of approximately 0.18 and 0.18, respectively, for the 80%/20% train/test split (right panels in Figures 4 and 5). Given more training data for the final model, these hyper-parameters should deliver an appropriate model complexity that does not over-fit the data. As before, I train 100 models with different random states to avoid being biased by the random components in the fitting process. Henceforth, I would report the median value of the predictions from the 100 model realizations. The data points are the working sample described in Section 2.1. The error bars are the standard deviations of the 100 predictions from the 100 model realizations. The dashed line is the one-to-one line.

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the perturbations is calculated for the entire working sample. Figure 8 presents histograms of the resulting changes in the output metallicities in the form of a “violin” plot. The top three features that the network output is most sensitive to are [S ii]/[N ii], [N ii]/[O ii], and [N ii] (deviate most from zero). Interestingly, photoionisation models have demonstrated that the [N ii]/[O ii] ratio is a good metallicity diagnostic (at above 0.2 solar) due to its insensitivity to ionisation parameter (Kewley & Dopita 2002; Dopita et al. 2013). The combination of [N ii]/[S ii] and [N ii]/Hα is also suggested by Dopita et al. (2016) to be an excellent diagnostic that is insensitive to ISM pressure and ionisation parameter. The neural network appears to rely more heavily on features that are also important in diagnostics developed from photoionisation models, despite that this information is not given a priori. This demonstrates the ability of the neural network to directly “learn” from the data.

3 RESULTS

3.1 Compare with other calibrations

It is worth comparing the MLP performance with other literature strong line calibrations to examine the putative improvement. Below, I apply different strong line diagnostics to the working H ii region sample and quantify their performance using the metallicity residuals, i.e. strong-line versus $T_e$ metallicity.

In Figure 9, I compare the strong line diagnostics using the O3N2 and N2 indexes calibrated by Pettini & Pagel (2004)\textsuperscript{7} and Marino et al. (2013). The H ii regions with indexes beyond the diagnostic limits are shown in grey. The calibrations provide 1D linear functions describing simple relationships between the input indexes and metallicity. Although the O3N2 and N2 indexes are not explicitly included in my feature list (Table 1), equivalent line ratios, [N ii]/[O iii] and [N ii]/Hβ, are a part of the input features. In Figure 9, I also compare the R and S calibrations developed by P16. These two calibrations each ingest three forbidden Balmer line ratios. The R calibration uses [O i], [O iii], and [N ii], and the S calibration [O i], [S ii], and [N ii]. These calibrations describe more complex 3D mappings from line ratios to metallicity.

In Tables 2 and 3, I quantify the residual distributions of the different calibrations. Table 2 tabulates the median, mean, and standard deviation derived from the entire sample. Since a large fraction of the sample was adopted to construct the R- and S- calibrations, a second comparison in Table 3 is done using only the CHAOS sample that has on average a higher metallicity than the P16 sample (see Figure 1). Even with the very simple architecture (Figure 3) and virtually picking the input features blindly, the MLP out-performs almost all the calibrations with its smallest median, mean, and standard deviation (Table 2). In Table 3, the MLP standard deviation is slightly larger than the R. calibration but its mean and median are smaller. The improvement of MLP over other literature calibrations is somehow marginal (a few to 30 percent in dispersion), how-

\textsuperscript{7} For the N2 calibration, the linear fit is adopted.
ever, the prediction accuracy of the ANN method increases with increasing sample size, as demonstrated in Figure 5. It is likely that when a larger training sample is available, the prediction accuracy can be improved further (see Section 4).

Below, I will apply the MLP calibration to H\textsc{ii} region data and its well-behaved radial gradient that covers a wide range of metallicity. The MLP calibration is applied to only regions that the model has not seen before. Theses include 25, 9, 18, and 24 regions from Croxall et al. (2016), van Zee et al. (1998), Li et al. (2013) and Kennicutt & Garnett (1996), respectively. The auroral lines were undetected in these regions. An additional 15 regions from the recent work by Hu et al. (2018, their Table 1) is also included.

In Figure 10, I show the metallicity gradient from the MLP calibration in the left panel. For reference, I show the metallicities from Croxall et al. (2016), Li et al. (2013) and Kennicutt & Garnett (1996) before. Thees include 25, 9, 18, and 24 regions from Croxall et al. (2016), van Zee et al. (1998), Li et al. (2013) and Kennicutt & Garnett (1996), respectively. The auroral lines were undetected in these regions. An additional 15 regions from the recent work by Hu et al. (2018, their Table 1) is also included.

### 3.2 Abundance gradients in nearby galaxies

I apply the MLP calibration to a sample of H\textsc{ii} regions in M101. The M101 galaxy is chosen because of its rich literature H\textsc{ii} region data and its well-behaved radial gradient that covers a wide range of metallicity. The MLP calibration is applied to only regions that the model has not seen before. Thees include 25, 9, 18, and 24 regions from Croxall et al. (2016), van Zee et al. (1998), Li et al. (2013) and Kennicutt & Garnett (1996), respectively. The auroral lines were undetected in these regions. An additional 15 regions from the recent work by Hu et al. (2018, their Table 1) is also included.

In Figure 10, I show the metallicity gradient from the MLP calibration in the left panel. For reference, I show the T_e metallicities from Croxall et al. (2016), Li et al. (2013) and Kennicutt & Garnett (1996) in the right panel. These H\textsc{ii} regions are in the training sample. In both panels, the grey dashed lines indicate the best-fit gradient from Croxall et al. (2016). Figure 10 demonstrates that the MLP calibration produces a reasonable metallicity gradient and comparable scatter, compared with the T_e method.
In the left panel of Figure 12, I present the mass-metallicity relationship derived from SDSS and SAMI spectra using the MLP calibration. In the right panel of Figure 12, I show the mass-metallicity relationships from applying the PP04 O3N2 and N2 calibrations to SDSS spectra (Kewley & Ellison 2008). I also show the relationship derived from oxygen auroral lines measured in SDSS stacked spectra (Andrews & Martini 2013). In both panels, the blue and red star symbols indicate the mass-metallicity relationship derived from blue and red supergiant stars (BSGs and RSGs) in nearby galaxies (Davies et al. 2017 and references therein). The MLP calibration yields reasonable slope and the zero point of the mass-metallicity relationship and follows closely the BSG/RSG relationship over 1 order of magnitude in oxygen abundance.

4 PROSPECT AND CHALLENGE

I have demonstrated that by fitting literature H\textsc{ii} region data with one of the simplest form of neural network, i.e. the multi-layer perceptron, it is possible to calibrate a strong-line diagnostic that performs reasonably well over a wide range of oxygen abundance. The MLP calibration performs equally well as the R and S calibrations by Pilyugin & Grebel (2016) and outperforms some popular literature calibrations base on N2 and O3N2.

The very first, simple attempt of this methodology demonstrates the prospect of applying this approach to large datasets coming in the near future. In particular, Figures 4 and 5 suggest that with larger training samples complex models can be developed and more accurately predict oxygen abundance. Although in this work I only develop a calibration requiring five strong forbidden lines, customised models can be developed and more accurately predict oxygen abundance. Although in this work I only develop a calibration requiring five strong forbidden lines, customised calibrations can be easily developed and their performance quantified by following the same work-flow.

The non-linear nature of the neural network, to an arbitrarily complex degree, delivers several advantages for developing strong-line diagnostics. It is no longer critical to
navigate through high-dimensional space via a series of 2-dimensional projections. For example, it is common practice to first select a primary feature sensitive to metallicity and search in the residuals for potential correlations with other line ratios. Non-linear correlation can also be more easily captured. For example, existing calibrations using the R_{23} index sometimes provide two different fits for the upper and lower branches due to the turn-over of the R_{23} index with metallicity (e.g. Pilyugin 2001; Pilyugin & Thuan 2005). Such upper/lower branch separation is integrated automatically in the MLP model. Furthermore, different line ratios are often sensitive to metallicity in different metallicity intervals. The MLP model adapts to this internally without needing to fit different indexes over different intervals and report the fit parameters and the applicable intervals (e.g. Curti et al. 2017). This makes both the development and application of a calibration more straightforward.

In this work, no attempt was made to optimise the choice of input features. The input features are picked virtually blindly base on possible arithmetic operations of the strong lines. While some features are known to be sensitive to metallicity, others might carry only limited information. It is thus expected that the network can perform even better with an optimised input feature set. The optimisation can be guided by Figure 8 but will require some trial-and-error by, for example, dropping one feature at a time and re-training the network and re-tuning the hyper-parameters. This is because that the network may response to feature trimming non-linearly and many existing features are degenerate.

Some additional features might also be important and are currently missing. For example, the electron density of the nebula, traced by either the [S ii]λ6716 to [S ii]λ6731 ratio or the [O ii]λ3726 to [O ii]λ3729 ratio, is a fundamental, yet missing physical quantity. The ionisation parameter is another important quantity that could improve metallicity calibration. Although the ionisation parameter in principle is encoded in the [O iii]λ5007 to [O ii]λλ3726, 3729 ratio already seen by the model, the ratio is also sensitive to metallicity (e.g. Kewley & Dopita 2002). The [S ii]λλ6766, 6732 to [S ii]λ6716, 6731 ratio is another key feature for passing the ionisation parameter information to the model given its insensitivity to metallicity (e.g. Kewley & Dopita 2002). These line ratios are currently missing in my attempt due to their non-uniform availability in the literature. Exploring the importance of these features using coming data from, for example, the CHAOS Project may be fruitful.

Despite the prospect and advantages of the neural network over traditional calibration methods, there are limitations and challenges. The predictive power of a neural network depends heavily on the quality and quantity of the training sample. The training sample in this work has only 43 H II regions (4.5%) more metal rich than 12+log(O/H) = 8.69 (solar), only 3 regions (0.3%) above 8.87 (1.5 solar), and no regions above 8.99 (2 solar). The lack of high metallicity training data is directly reflected in the mass-metallicity relationship (Figure 12). The SDSS and SAMI galaxies have no MLP oxygen abundances larger than approximately 12+log(O/H) = 8.8 or 1.3 solar. The mass-metallicity relation from the blue and red supergiants, however, reaches up to 12+log(O/H) = 8.9. The systematic underestimate of MLP metallicities for the high-metallicity and high-mass galaxies results in the SDSS median mass-metallicity relation to saturate at approximately 12+log(O/H) = 8.7 at the high-mass end. Furthermore, a low-metallicity floor of 7.4 was also imposed due to the sparse sampling in the low-metallicity end (see Section 2.1). The lack of high and low metallicity training data puts a fundamental constraint on the interval where the calibration is reliable. For most literature calibrations, extrapolating the calibrations outside the applicable metallicity interval can be easily visualised, and in some cases justified by comparing with photionisation models (e.g. Maiolino et al. 2008). Nonetheless, the non-linear nature of the neural network puts a fundamental constraint on making predictions outside the parameter space populated by the training sample.

The uniformity of the training data is also key to the quality of the predicted metallicity. Non-uniformly analysed or low quality input data will introduce noise and bias to the network. Artefacts arise from non-uniform analysis can already been noticed in my MLP model. In Figure 13, I present the residuals of H II regions from different sources. The P16 residual distribution is very similar to the overall residual distribution, because the P16 sample comprises the majority (84%) of the working sample. The CHAOS distributions, however, are noticeably different from the overall distribution, with CHAOS I significantly skewed to negative residuals and CHAOS II and III more consistent with the overall distribution.

The differences come from how the oxygen abundances are derived from auroral line measurement(s). In the P16 sample, the temperature scaling relationship of Garnett (1992) is always used to infer the high- or low-state oxygen temperatures. In fact, 92% and 19% of the P16 sample have T([O iii]) and T([N ii]) measured from the [O iii]λ4363 and [N ii]λ4575 lines, respectively. As in the two-zone model, T([O ii]) is approximated by T([N ii]). A small fraction of the sample (11%) has both auroral lines detected and the mean of the two metallicities is adopted. In the CHAOS Project, multiple auroral lines are typically detected in each regions. Different auroral lines and scaling relationships are adopted.
to determine the low and high state temperatures. For the low state, T([O\text{II}]) is usually approximated by T([N\text{II}]) even when T([O\text{II}]) is measured. For the high state, CHAOS I prioritizes T([S\text{III}]) over T([O\text{III}]) while CHAOS III adopts the opposite. CHAOS II has no [O\text{III}],4436 detections and adopts T([S\text{III}]). When only one of the two states is measured, scaling relationships are adopted. CHAOS I and II adopt the G92 relationships, and CHAOS III adopts updated relationships fit to their sample. These various factors determine the final T([O\text{II}]) and T([O\text{III}]) used for calculating the oxygen abundances. Figure 14 shows the T([O\text{II}]) and T([O\text{III}]) “used” (right panel) and those “measured” (left panel). For the CHAOS II and III sample, the temperatures used in determining oxygen abundances lie close to the G92 scaling relationship, consistent with the P16 sample. This consistency is directly reflected in the residual distributions in Figure 13. A significant fraction of the CHAOS I sample, however, falls systematically above the G92 scaling relationship, resulting in the skewed residual distribution in Figure 13.

In general, there is an unavoidable trade-off between sample uniformity and sample size. In this exploratory work, sample size is prioritized over uniformity. Thus, all the CHAOS samples are included even though a significant fraction (56%) has no [O\text{III}],4436 detections. Nonetheless, it is expected that in the next few years, uniform analyses on large samples will be feasible by virtue of two emerging high quality H\text{II} region catalogs. The CHAOS Project has detected multiple auroral lines in a few tens H\text{II} regions in each of the three published galaxies. With a sample size of a dozen nearby galaxies, close to 1,000 H\text{II} regions are expected. Large IFU surveys of nearby galaxies with 10-m class telescope (e.g. the MUSE/VLT large program by the PHANGS collaboration\(^9\); Kreckel et al. 2018; Ho et al. in preparation) will also report auroral line metallicities. The Local Volume Mapper of the planned SDSS V survey will detect auroral lines in thousands of H\text{II} regions in M31, M33, LMC, SMC, and the Galaxy (Kollmeier et al. 2017). These coming data will significantly improve the quality and quantity of the training data (even at super solar abundances), allowing both an increase of neural network complexity and its predictive accuracy.

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APPENDIX A: VARIATION OF MLP PREDICTION DUE TO LINE FLUX UNCERTAINTY

Non-linear models often respond non-linearly with input perturbations. In the presence of measurement noise, the non-linearity could be an undesirable characteristic for calibration. To understand how the MLP model reacts to measurement uncertainties of the input line fluxes, I perform Monte Carlo simulations. Gaussian noises are added to the four line fluxes ([O\text{II}], [O\text{III}], [N\text{II}], and [S\text{II}]) and the resulting features are adopted to derive MLP abundances. Five hundred realisations are created for each of the 962 regions. The five curves in each panel describe the distribution function of the MLP abundance fluctuation (2.3%, 15.9%, 50.0%, 84.1%, and 97.7%). The MLP abundance fluctuation is defined as the abundance difference between predictions using noise-free and noisy line fluxes.
is, adding noise to the line fluxes does not seem to skew the resulting MLP abundances in a statistically significant way.

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