Wine Analyte Prediction using Neural Networks

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1 Abstract

It has been determined that certain analytes present in common wines cause negative variations in the perceived quality of the products. The main compounds that adversely affect wine quality are 4-ethylguaiacol (4EG) and 4-ethyphenol (4EP). Enoses have become a cost efficient mechanism of monitoring for these analytes at the production stage[1]. In this project, 38 time series readings of a particular eNose (MOS-Enose) sensor in the presence of 38 different wines are provided with corresponding 4EG and 4EP values. The task of the project is to learn a model that can predict 4EG/4EP values from MOS-Enose readings in the presence of an unseen wine. 8 such wine measurements are provided and the task of the project is to provide predictions as to their analyte values. This report details the approach used to provide such predictions.

2 Initial Analysis of Datasets

The first step in this project was to get a feel of the data to best plan an approach for how to build the relevant prediction $\operatorname{model^1}$ Looking statically at the dataset, it is clear that the dimensionality of the dataset is huge. For each of the 12 sensors there is a reading twice every second for 600 seconds. Consequently, using the raw dataset as is a model that represents 1200×12 dimensions would have to be considered. Such massive dimensionality would make a model incredibly susceptible to noise and generalisation would be very hard especially in the presence

of a small training set like the 38 training instances in this dataset. By performing pre-analysis on the

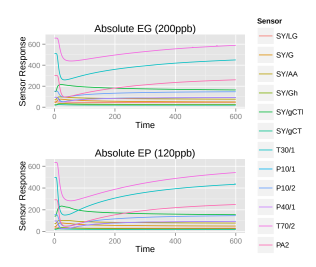


Figure 1: Absolute EG and EP readings of all sensors

dataset insights into whether certain sensors are redundant or correlated can be determined. If either of these properties can be found to be true then it is good evidence to suggest some dimensionality reduction that can be performed on all data prior to being fed into the predictive model.

Provided as part of the dataset is a calibration set. Here, readings from various artificially constructed wine concentrations of 4EG and 4EP are measured to observe the effects on the sensors. Along with these readings, benchmarks are also provided of wine without any contaminants present.

 $^{^1{\}rm The~source}$ for the analysis is provided in the R file calibration_analysis.r

2.1 Graphical analysis

The first step taken was to simply get a graphical feel of how each sensor responds, over time, to the presence of each analyte (see Figure 1^{2})

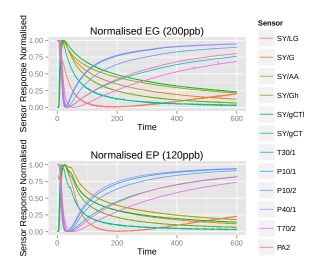


Figure 2: Normalised EG and EP readings of all sensors

Here it can be seen that the sensors clearly take time to react with the wine (probably due to diffusion of the aroma particular from the wine to the sensors). More over, at first glance it appears that certain sensors react more then others. This observation is based on the assumption that the scale that each sensor operates on is the same. If these sensors do not operate at the same scale then such a statement could be a very dangerous falsehood. Considering the normalised representations of these readings we can see that certain sensors contain a higher area under the graph which may be representative of that sensor reacting more than others. This shall be explored later. Also observable in these charts is that the data is quite noisy. To combat this, kernal smoothing (using NadarayaWatson kernel regression estimates[4]) has been applied to all further datasets in this report to smooth the noise out.

The next step is to consider how the sensors behave as a difference from the baseline. The idea how is to try and isolate the sensor reactions that respond to the analytes themselves. Background odours from the wine and its environment are going to contributing to the sensor reactions. By removing the baseline readings (where no analytes were present) an approximation can be derived for the relative target compound reactions. This requires a look at the relative readings when the baseline has been subtracted (see Figure 3)

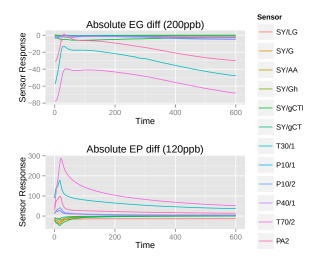


Figure 3: Absolute EG and EP readings of all sensors with baseline removed

Observing the effect of the difference over the relative baselines gives an indication that certain sensors are simply not reacting to the target compounds. From this analysis it appears that sensors PA2, T30/1 and T70/2 are stimulated the most. What is also apparent from this representation is that there is an indication that these sensors are correlated (co-insides with the conclusions of [1] and [2]). As one increases, so do the others. It can also be seen here how the sensors differ in the presence of 4EG and 4EP. 4EP appears to cause the sensors to increase their relative measurements whereas 4EG dampens the sensor's responses. This implies that the presence of the 2 substances could be at odds with each other.

 $^{^2{\}rm Note}$ that all data presented in this report is from averaged data of multiple readings of identical experiments where available

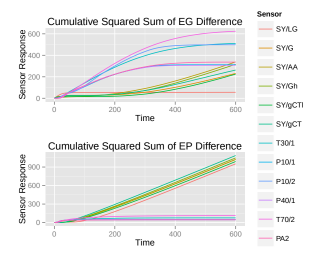


Figure 4: Monatomic Cumulative Squared plot of normalised EG and EP readings of all sensors

Also, although these mentioned sensors do appear to react heavily to the presence of the target analytes, it again is on the assumption that the operating scales of the sensors are consistant. As none of the other sensors are absolutely zero (except for maybe SY/gCTI) at this stage it is still not possible to conclude that they are not reacting to these components. As mentioned previously, looking at the integral of the normalised datasets might be useful to gauge how much 'power' each sensor has over the others when all normalised between 0-1. To visual demonstrate this analysis a cumulative squared summations³ of the baseline removed datasets were considered (see Figure 4. The results show that there are indeed some sensors that dominate over others for the presence of the various analytes. Sensors such as SY/LG clearly provide little information whereas T70/2 dominates for EG reading. At first glance it feels that a threshold could be set and all sensors that exhibit a total reading below that should be excluded but this doesn't help remove the correlations that are clearly visible in this dataset. It is clear that graphical analysis of the data is not enough to conclude if any sensors are redundant and more analytical approaches are required.

2.2 PCA

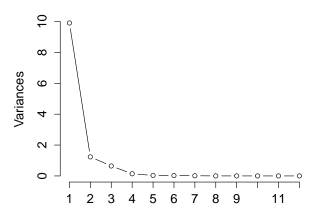
Principle Component Analysis is a process of projecting a dataset in a new space, represented as projections from new orthogonal basis vectors (eigenvectors)that separate the data dimensions so no correlations are present in the new space. Such a technique can be used to project a dataset onto this new completely othogonal space with zero mean and unit variance (whitening transform[3]). Doing so means that each dimension is in the same scale (normalised) and expresses the maximum amount of information for that dimension⁴ Furthermore, by looking at how the projections (or scores) of the original dataset on these new basis vectors vary, it can be easily observed which of the new dimensions contribute the most information of the dataset. Dimensions where the projections vary alot(high variance) imply that there is alot of information in this dimension. Dimensions where the projections do not vary much means that this dimension tells little information about how the data set evolves. Using this, components that hold little information can be removed from the dataset and thus the total dimensionality of the dataset can be reduced.

From studying the results of the PCA analysis on the EG difference results, it can be seen that PC1 contains 0.8258 of the total variance, whereas PC2 contains 0.1027. The rest of the components account for an accumulated 0.0715 suggesting that these can all be discarded as they dont contain much useful information. Using the kaiser criteria also confirms that only PC1 and PC2 need to be kept as they are the only components that have eigen values above 1. Looking at the EP difference dataset tells a similar story but this time PC1 contains the majority of the information of the dataset with a 0.9209 proportion of the variance. Again the scree plot and kaiser criteria suggest that PC1 need only be kept(see Figure 5). As another visual aid, the biplots show in Figure 6 show

 $^{^3}$ squared to remove negative values offseting the monotomic requirement of this analysis

 $^{^4}$ correlated dimensions in a sense share information and therefore are not as expressive as 2 dimensions that are uncorrelated

Scree Plot for EG difference



Scree Plot for EP difference

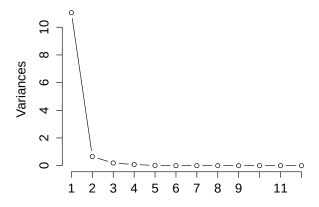
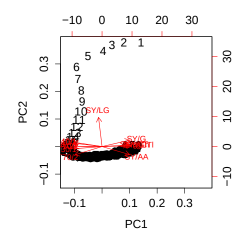


Figure 5: Scree Plot for EG/EP readings showing PC variances

how the time samples vary around the components⁵. Also on the plot are the relative sensor loadings that point in the direction of the PC they attribute to the most. To summarize, by performing PCA on the

data, the new feature space can represent the data in a consistant scale (removing the issues experienced using the graphical methods) and also provides the majority of the information in 3 principle dimensions (2 for EG and 1 for EP). Projecting a dataset onto these components greatly reduces the dimensionality of the dataset



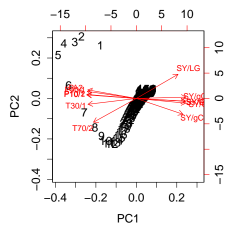


Figure 6: BiPlot for EG/EP Main Principle Components

 $^{^5{\}rm To}$ make the results more visible, the datasets have been desampled to 1 sample every 4 seconds, with values averaged between samples. See Section 2.3

2.3 Desampling

As already mentioned, using PCA, the new feature space has reduced the dimensionality to just 3 principle components. Given that there are still 1200 readings for each of these dimensions, this still leaves a total feature set containing 3600 dimensions. More work is needed to reduce it further. Currently the data is sampled at 2Hz meaning there are two readings every second. By desampling to a lower frequency and averaging the replaced samples, the dimensionality can be greatly reduced. To determine the correct degree to what degree of desampling should be performed, PCA can be considered again. The intuition here is that the dataset should be desampled as much as possible but only so much that the variance of the determined PCs does not drop too much. Too much in this case is so that the kaiser criteria is still up held to the point that EG (PC1,PC2) and EP(PC1) are still the dominent components (it might be possible to desample even further but then include more PC's but this will be left as possible future areas of study). Further to this, although values of < 0.025Hz (one reading every 40 seconds - see Figure 7) seem to maintain similar variance values of the PCs being considered, The performance of the model will be evaluated with different desampling values to tune this parameter appropriately. As a guidance, resampling at 0.025Hz provides a $3 \times 16 = 48$ dimensionality model. Something much more manageable then the original dataset.

3 Learning a model

Now that the data has been analysed and a proposed workable dataset with ≈ 48 dimensions, an appropriate prediction model needs to be chosen. Due to the computationally efficient learning and prediction phase, its intuitive geometric representation and simply because i have never implemented one before, the RBF neural network has been chosen for this prediction problem. Before discussing implementation details of the network itself, based on the findings from the previous section, a preprocessing layer is required for all data into the network to transform its high di-

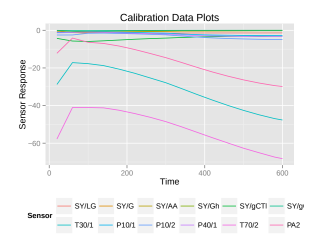


Figure 7: Example of desampling the EG difference plot to 0.025HZ. The main structure of the data is still present

mensional space into something more manageable.

3.1 PreProcessing

The next stage of this project was to perform a consistant series of steps to transform all data (both training and prediction sets) into a feature space more concise then the original dataset provided. The reduction transformation steps have already been discussed so the following serves as a formal reference to the pre processing steps used. It is worth pointing out however that the following steps are no longer performed on the calibration dataset (unless explicitly stated) but on the training and prediction sets as they enter the model. The order that the processing occurs is as presented. The preprocessing steps can be found in preprocessing.r and preprocessing-utils.r R source files.

- All Data is smoothed using NadarayaWatson kernel regression estimates
- 2. All Data is subtracted from the calibration set's baseline readings (average of both 4EG and 4EP readings) to reduce impact from other properties of the wine and its environment.

- 3. All Data is downsampled to x Hz where x<2. This value is a free parameter that will be evaluated as part of the network's performance.
- PCA basis vectors of desampled dataset calculated
- 5. All Data is projected onto the loadings from the PCA basis vectors ⁶.
- 6. All but PC1 and PC2 features are removed from the datasets.

3.2 Network

The network used in this project is an RBF network with k k-means positioned prototypes and it's implementation is provided in the network.r R source file. The general principle of the network is that a training set is used to cluster k centers or prototypes around the instances. Each prototype has a position in the feature space that radiates a field around it. When a instance is to be classified it is positioned in the feature space according to the values assigned to it. The position of this instance will sit in the radius fields of the prototypes defined with each field's strength being proportional to the distance from the instance to the field's origin prototype. The strength of the field will determine how much that instance is influenced by the prototype and it's influence is then multiplied by a learned weight attributed with the prototype. An instance can and most likely will be positioned over a number of fields so the overal output of the network is the summed value of all prototypes influences.

The centers are of the prototypes are positioned using the k-means algorithm whereby k randomly generated clusters (in the bounds of the min/max values of the instance set used) are iteratively moved towards clusters in the dataset. These clusters are defined by assigning membership of instances to the prototypes using the L2 norm distance:

$$||x|| = \sqrt{\sum_{i=1}^{N} (x_i - y_i)^2}$$

The prototype with the lowest distance to an instance (closest) is the prototype that gains this instance as membership. After each instance is assigned to a prototype, the mean distance of all prototype members is computed and the prototype is moved to this position. Membership is recomputed and the prototypes position moved until no more movement occurs.

An implementation consideration is what happens when a prototype does not have any members after the algorithm finished. A simple approach would be to simply remove the empty prototypes but this could, due to the starting initial conditions being random, mean that all instances could be assigned to a few prototypes, meaning that we have less positions to model our training data. As the dataset is very sparse (only 38 training instances), having less then k prototypes would have a large impact when measuring the performance of the network (especially when evaluating what k value to use Consequently a recursive implementation that retries the algorithm until each prototype is assigned a member. Although this scheme is a little brute force, in practice this doesn't add any noticeable overhead when $k \ll |instances|$. A better scheme using an optimization approach might be more appropriate if the instance numbers are much larger.

Once the prototypes have been defined and positioned, a radial basis function needs to be applied to them. For this project, a gaussian implementation of the form is used:

$$\phi(x) = \exp(-\frac{\|x\|^2}{2 \times \sigma^2})$$

The sigma used is calculate in either two ways. If there are enough instance examples (currently set to >50), then the covariance matrix of the actual instances are used. If there are <50 (which will always

⁶The loadings used are from both the EG and EP analysis. First the data is rotated in the EG space, then further rotated into a representative EP space

be the case with this training set) then the sigma of the gaussian is set at a constant width proportional to the maximum distance between any 2 prototypes in the network (to ensure every prototype can 'reach' any instance in the feature space). The sigma used is defined by

$$\sigma = d_{max}/\sqrt{2*k}$$

Its worth mentioning that because of the normalising whitening step used as part of the PCA preprocessing, the instances fed into the network will have 0 mean and unit variance meaning that the gaussian function will not need to use any skewed covariance matrix hence it should be acceptable to use the global sigma calculation.

Once the functions have been set up on the prototypes, the next step is to learn the weights of the network. This is accomplished arthimetically by minimizing the error terms (expected - actual) in the quadratic sense. Using an adaptive form of the psuedo inverse matrix transform (single value decomposition) multipled by the expected values (\underline{D}), the weights can be calculated with an error penalty λ term to help aid smoothing the error rates and thus help increase generalisation. The complete learning stage is performed using Poggio & Girosi's regularization applied form[5]:

$$\underline{W} = [\Phi^T \Phi + \lambda I]^{-1} \Phi^T \underline{D}$$

3.3 Determining Free Parameters

As already noted, this network contains a number of free parameters that need to be fixed prior to training and predicting phases. In summary, these parameters are:

- 1. n Hz desampling rate
- 2. k prototypes
- 3. λ error penalty term

To evaluate the best possible combination of these values prior to predicting the unseen dataset, a N fold cross validation scheme is used. This process involves

taking the 38 training instances and randomly splitting them into n equal bins. The process works by iterating over the bins, one by one, using each bin in turn as a test set and the rest of the bins as the training set. This continues until every bin has been the test set against the rest as the training set. The error rates of each iteration are then averaged using mean squared error and this value is deemed the overal error rate of that particular network (and its configuration). The idea here is that the performance of the network is evaluated on all combinations of instance training to test sets rather then just only, potentially lucky/unlucky configurations. Due to the fact that the training set is so small, having as many instances in the training fold is most important. For this reason, 18 fold X-validation was used. This equated to 18 tests of 36/2 train/test splits. The implementation is provided in evaluation.r.

4 Performance results

After letting the evaluation run to completion, it determined that that the best values for the free parameters are:

Parameter	$error_{min}^2$	$error_{max}^2$	$error_{avg}^2$
k = 25	130	231300	32970
n = 40 -	0.0	0.0	0.0
48D			
λ	0.0	0.0	0.0

The choice of the parameters was driven by the $error_{avg}$ whereby the parameter that generated the lowest average error was deemed the best⁷. Output of the evaluation scripts is presented in performance-results2.txt. It is worth noting, that using higher dimensionality feature spaces takes a lot of time to train (>20 mins). This may not be a problem with

⁷There are two evaluation scripts considered in this project. evaluate1() evaluates all combinations of all parameters to ensure that every parameter is considered. The idea being that the parameters are not independent and that there is an optimal configuration of textitall parameters. Due to exponential $O(k^{n^{\lambda}})$ nature of evaluate1 in evaluation.r, it simply took too long to execute. evaluate2 on the other hand evaluates each parameter one by one, holding the other parameters consistant. This implies that the parameters are independent but does complete in $O(k+n+\lambda)$

5 Wine Predictions

Based on the processes described in this report, the following values are the predictions for the 8 unseen wines. The code for this part of the project is presented in prediction.r

Wine Id	4EG	4EP
731	0.0	0.0
734	0.0	0.0
738	0.0	0.0
744	0.0	0.0
746	0.0	0.0
754	0.0	0.0
759	0.0	0.0
768	0.0	0.0

6 Conclusions

References

- [1] http://pubs.acs.org/doi/abs/10.1021/jf7037289
- [2] http://www.plosone.org/article/info% 3Adoi%2F10.1371%2Fjournal.pone.0006406
- [3] http://en.wikipedia.org/wiki/Whitening_transformation
- [4] http://en.wikipedia.org/wiki/Kernel_regression
- [5] http://cbcl.mit.edu/people/poggio/ journals/poggio-girosi-science-1990.pdf