Novel Convolution-Based Signal Processing Techniques for an Artificial Olfactory Mucosa

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Abstract—As our understanding of the human olfactory system has grown, so has our ability to design artificial devices that mimic its functionality, so called electronic noses (e-noses). This has led to the development of a more sophisticated biomimetic system known as an artificial olfactory mucosa (e-mucosa) that comprises a large distributed sensor array and artificial mucous layer. In order to exploit fully this new architecture, new approaches are required to analyzing the rich data sets that it generates. In this paper, we propose a novel convolution based approach to processing signals from the e-mucosa. Computer simulations are performed to investigate the robustness of this approach when subjected to different real-world problems, such as sensor drift and noise. Our results demonstrate a promising ability to classify odors from poor sensor signals.

Index Terms—Convolution, electronic nose, signal processing.

I. INTRODUCTION

O
er the past 20 years, significant advances have been made in the understanding of the mechanism by which odors are detected by the human olfactory system. This has led to the concurrent development of instruments designed to detect odors and commonly known as artificial or electronic noses (e-noses) [1].

However, sensor-based electronic noses today generally suffer from significant weaknesses that limit their widespread application. Their sensing ability is heavily affected by environmental factors: general drift due to temperature, humidity and background noise, sensor variations and sensor poisoning. These problems, in addition to often wanting to detect very low concentrations (below PPM) of the odor in air [2], [3], make the design of an electronic nose difficult even with expensive autosamplers and the supply of clean air.

Faced with this challenge, novel devices are being designed to tackle these issues and improve detection thresholds and classification success rates; examples include combining an electronic nose array with a gas chromatography column or mass spectrometer. These analytical instruments are large and expensive and place significant limitations on the way in which their application and potential market.

More recently, it has been suggested that chromatography plays a role in odor discrimination within the biological olfactory system, known as nasal chromatography [4]. This report suggests that the aqueous layer covering the olfactory receptors in the olfactory epithelium (Fig. 1) acts as a retention layer as odors move along and through it, functioning similar to a stationary phase coated channel in a gas chromatographic system. Hence, different odors are partitioned and transported at different rates to olfactory receptor cells – leading to different temporal signatures. This may even be further amplified by the expressing of different binding proteins within the olfactory epithelium.

While e-noses are becoming more common, the processing methods used to analyze and classify sensor data are generally time-invariant models. Statistical methods extract time independent features from the sensor signals, such as baseline (zero gas) and peak (analyte present) values or some simple parameter from them (difference, ratio, etc.). These static features are then used to separate the groups of data and classify the results. Alternatively, nonlinear but still static methods such as artificial neural networks are used to classify these static features.

While these processing methods have proved to be sufficient for traditional e-nose devices, the new architecture of the e-mucosa provides novel data sets unlike those encountered in the
Fig. 2. Photograph of the embodiment of the artificial olfactory mucosa with 40 sensors distributed along a meandering coated channel [6].

Fig. 3. Sensor arrangement in the artificial olfactory mucosa [6].

The data produced by this device are now more information rich, containing information relating to both the spatially separated data from different sensors (classical sensor-based e-nose) and temporal differences between identical sensors at different locations (GC-like system). This novel spatio-temporal data set has been analyzed by treating the domains independently and treating them as individual features for the pattern recognition stage of processing. However, information relating to the relationship between these two domains is lost. A novel method of processing is required to fully utilize the information contained within these novel data.

This paper explores a simplified subsystem of the e-mucosa, employing one sensor array at the start of the channel, and a second set at the end of the channel. This simplified e-mucosa, or perhaps a kind of “tandem electronic nose,” considerably reduces the dimensionality of the problem to solve. The aim is then to apply novel convolution and signal combination techniques to this signal generated and thereby improve classification by low-cost instruments in the field of artificial olfaction.

II. PROCESSING METHODS

The statistical processing of data is carried out in several stages. First, a signal preprocessing stage is used to prepare the data for processing. This can involve normalizing and/or filtering the data. The functions carried out here should be considered carefully, as the quality of some aspects of the data will be improved while others are diminished. For example, normalization will make each set of data directly comparable; however weak signals dominated by noise will have their relative noise magnified greatly.

Features are then extracted from the data sets. The aim of this process is to determine a parameter that can be used to differentiate between the data from one input source and another. Multiple features can be extracted. This adds to the possibility of good discrimination, but also adds extra dimensionality to the data and the potential of redundancy in the data. This paper will only be considering single-feature processing methods.

The feature set is processed using a method that organizes data for pattern discrimination or classification. Various robust statistical methods for this process exist [7]. In the field of artificial olfaction, principle components analysis and discriminant function analysis are commonly used [8], [9]. These functions present high-dimensional data sets in a manner so that there is maximum variance in a minimum number of dimensions.

Classification is the final step in the processing procedure. This is done statistically, utilizing methods such as linear metrics and nearest neighbor linkage.

The methods presented hence focus on the feature extraction stage. These methods will be combined with different standard normalization techniques in the preprocessing stage. The extracted feature set will then be processed using principle components analysis, and benchmarked based on the quality of the separation in the principle component (PC) plots. Classification is not carried out explicitly, because we are seeking excellent linear separability of the samples.
A. Preprocessing (Normalization)

Since the magnitude of the sensor output can vary greatly, it is important to normalize each sensor’s raw data \((X^j)\), where \(j\) is the sensor index so that one sensor can be compared to another in an unbiased manner. However, any noise on small signals will be amplified greatly and must be carefully considered.

Two normalization schemes were used, signed auto-scaling and auto-ranging. Auto-scaling takes the data and scales the magnitude linearly so that all data points lie between 0 and 1, and is signed to preserve the direction of the magnitude

\[
S^j(t) = \frac{|X^j(t)|}{\max(|X^j|)},
\]

(1)

Auto-ranging scales the data in a linear manner so that the range of the data is 1. The advantage here is to preserve any swing that may occur in more advanced problems, as multiple odors may cause sensors to respond in both directions

\[
S^j(t) = \frac{|X^j(t)|}{\max(|X^j|) - \min(|X^j|)},
\]

(2)

These normalized data were passed to the feature extraction stage, along with unmodified raw data for comparison and benchmarking

\[
S^j(t) = X^j(t),
\]

(3)

B. Feature Extraction

In order to extract features from the two sensor arrays highlighted in Fig. 3, \(S_A\) and \(S_B\), the normalized signals from these arrays must be combined. Each sensor output from \(S_A\) is combined via a function with the equivalent sensor in \(S_B\). Different combinatorial functions will generate different characteristic signals. Convolution is often used in signal processing for combining and identifying similarities between signals [10]. An example of this kind of characteristic signal is shown in Fig. 4(b)

\[
y(t) = \int_{\tau=-\infty}^{\infty} S_B(\tau)S_A(t-\tau)\,d\tau.
\]

(4)

The product of the two data arrays can also form a characteristic signal. To preserve magnitude, a signed root product is used. Fig. 4(c) illustrated an example of this characteristic signal

\[
y(t) = \text{sgn}(S_A(t)S_B(t))\sqrt{|S_A(t)S_B(t)|},
\]

(5)

The third characteristic signal considered is a simple difference between two signals. The resulting characteristic signal is illustrated in Fig. 4(d)

\[
y(t) = S_A(t) - S_B(t).
\]

(6)

In order to be used in the main processing step, a feature needs to be chosen and extracted from these characteristic signals. The area under the curve was chosen, as this feature contains information on the shape of the signal. Combination signals obtain their shape from the shape of the constituent signals, with variations in the shape of the source signals altering the overall shape, and thus area, of the combined signal.

As the e-mucosa incorporates concepts of other traditional e-nose devices, these can also be extracted from the same data set and provide a point of comparison. For these devices, a suitable feature needed to be chosen for extraction and use in the main processing step. Peak magnitude was selected due to its prevalent use within standard e-nose devices [1]–[3]. Both an e-nose (\(S_A\)) and an advanced form of z-nose\(^\text{TM}\) (an e-nose preceded by a GC column) \((S_B)\) were extracted and processed, so both classes of device could be considered as comparison

\[
y(t) = S_A(t)
\]

(7)

\[
y(t) = S_B(t).
\]

(8)
The final feature extraction considered is referred to as cross-convolution. Whereas the convolution method already presented convolutes two identical sensors in different arrays, cross-convolution convolutes two different sensors in the same array. This process was applied to both $S_A$ and $S_B$, and the area under the curve was extracted as the feature for future processing:

$$y^{ij}(t) = \int_{-\infty}^{\infty} S_A^i(\tau) S_A^j(t - \tau) d\tau$$  \hspace{1cm} (9)$$

$$y^{ij}(t) = \int_{-\infty}^{\infty} S_B^i(\tau) S_B^j(t - \tau) d\tau.$$  \hspace{1cm} (10)

where $i$ and $j$ are different sensor indices. This process creates a large number of possible convolution pairs, some examples of which are illustrated in Fig. 5. The rapidly increasing computational cost as the number of sensors rises, and the addition of redundant noise make it impractical and undesirable to use all these possible convolutions in data processing. Thus, a subset of convolutions is chosen for the processing stage.

C. Data Processing

Once a feature array has been established from the feature extraction process, it is entered into a principle components analysis (PCA). This process establishes principle components along which variance of the data is maximized.

The first 2 or 3 principle components are then used to generate a data plot. From the degree of separation observed in this plot and variance information obtained from the principle component analysis, the different processing methods can be easily compared.

III. SIMULATION

Utilizing data simulated during the development of the e-mucosa [6], [11], which simulated a 5 s pulse of three analytes (ethanol, toluene, and a 50/50 mixture of the two) passing through the system. The microchannel simulated has a cross-sectional geometry of 0.5 mm $\times$ 0.5 mm and a length of 2.3 m, and was coated with parylene C as the stationary phase. Forty sensors, in four groups of ten different tundra, were placed along the microchannel, as shown in Fig. 3. These simulations were carried out in commercial software, MATLAB using a finite-element model [11].

This simulation would be used to determine strengths and weaknesses of the processing methods outlined previously. In order to provide a meaningful test, noise needed to be added to the e-mucosa simulation data. An e-nose system is subject to many kinds of noise, ranging from environmental factors, variations in analyte concentration, variations in sensor sensitivity, changes in the sensors over time and poisoning of the sensors. These major noise sources were considered as part of three major categories of noise.

A. Additive Noise Signal-to-Noise Ratio (SNR)

Additive noise represents primarily background noise sources – sensor noise, background environmental factors and taints in the environment. This noise source was modeled as white Gaussian noise, with zero mean and a variance based on a percentage of the sensor’s peak magnitude

$$X_{SNR}(t) = X(t) + n(t)$$  \hspace{1cm} (11)$$

where $X_{SNR}(t)$ is the final noisy signal, $X(t)$ the simulation signal and $n(t)$ the noise signal

$$n(t) = n \times \max(X) \times \text{randn}(t)$$  \hspace{1cm} (12)$$

where $n$ is the noise value, and $\text{randn}$ is a number series generated by MATLAB’s random number generation function $\text{randn}$ with a mean of 0 and variance of 1.

B. Sample Variance (SV)

Sample variance (SV) focuses on factors directly affecting the response of the sensors – sensor variation and sample variation. This noise source was modeled as a Gaussian value with a mean of 1 and a variance expressed as a percentage

$$X_{SV}(t) = X(t)N$$  \hspace{1cm} (13)$$

where $X_{SV}(t)$ is the final noisy signal, $X(t)$ the simulation signal, and $N$ the noise value

$$N = 1 + (n \times \text{randn})$$  \hspace{1cm} (14)$$

where $n$ is the noise value and $\text{randn}$ is a number generated by MATLAB’s random number generation function $\text{randn}$ with a mean of 0 and variance of 1.

C. Baseline Drift (BD)

Baseline drift (BD) is a major problem in e-nose systems. This represents variations in the sensor’s baseline, as a result of age, poisoning, and temperature. This noise source was modeled as a Gaussian value with a mean of 0 and a variance based on a percentage of the sensor’s peak magnitude

$$X_{BD}(t) = X(t) + N$$  \hspace{1cm} (15)$$

where $X_{BD}(t)$ is the final noisy signal, $X(t)$ the simulation signal, and $N$ is the noise value

$$N = n \times \max(X) \times \text{randn}$$  \hspace{1cm} (16)$$

where $n$ is the noise value, and $\text{randn}$ is a number generated by MATLAB’s random number generation function $\text{randn}$ with a mean of 0 and variance of 1.

In order to prevent extreme differences in drift occurring in the simulated channel, each pair of equivalent sensors are subjected to the same drift value.

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The e-mucosa data set was replicated 20 times with each of these noise sources and the noise value set to 5%, producing three sets of 20 samples, each set having one noise source added. Another three data sets, also of 20 samples each, were produced with binary combinations of noise sources (SNR + SV, SNR + BD, SV + BD).

The six noisy data sets where then normalized using the three normalization methods presented in the data processing section, producing three groups of six normalized data sets. Each normalized group was then processed using the seven feature extractions presented in the data processing section, producing an array of 42 PCA results in each of the three normalization groups.
TABLE I

<table>
<thead>
<tr>
<th>t-scores</th>
<th>EM-Convolution</th>
<th>EM-Product</th>
<th>EM-Difference</th>
<th>E-Nose</th>
<th>GC+E-Nose</th>
<th>X-Convolution</th>
<th>ZX-Convolution</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>PC 1</td>
<td>PC 2</td>
<td>PC 1</td>
<td>PC 2</td>
<td>PC 1</td>
<td>PC 2</td>
<td>PC 1</td>
</tr>
<tr>
<td>SV</td>
<td>91.30</td>
<td>93.02</td>
<td>167.43</td>
<td>231.76</td>
<td>71.44</td>
<td>0.15</td>
<td>198.13</td>
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<tr>
<td></td>
<td>116.43</td>
<td></td>
<td>231.76</td>
<td></td>
<td>71.44</td>
<td></td>
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</tr>
<tr>
<td>SnR</td>
<td>116.43</td>
<td></td>
<td>231.76</td>
<td></td>
<td>71.44</td>
<td></td>
<td></td>
</tr>
<tr>
<td>BD</td>
<td>116.43</td>
<td></td>
<td>231.76</td>
<td></td>
<td>71.44</td>
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</tr>
</tbody>
</table>

IV. SIMULATION RESULTS

Each processing result was plotted with the first two principle components, examples of which can be seen in Fig. 6.

t-scores between each pair of analytes were calculated for the first two principle components using a separate variance t-test in Unistat 5.5 software. Table I provides a summary of results for the differing processing methods when using auto-scaling normalization.

Overall, our results showed that auto-scaling was the preferable normalization method. In the presence of significant sample variance and SNR noise, the e-mucosa and a convolution or product characteristic signal showed a marked improvement over the traditional e-nose systems, obtaining a higher t-score over the two principle component dimensions. However, these same methods show a weakness to very large amounts of baseline drift in the system during the measurement period. This is actually unlikely as the measurement period is short (minutes) compared with conventional e-nose sensors that need sensor stable over weeks or months. The difference signal performed poorly overall, although was superior to the convolution and product methods in the presence of baseline drift.

V. EXPERIMENTAL DATA

A limited selection of experimental data was available from a prototype e-mucosa system [6]. This data set contains five repeats of data for ethanol, toluene, and a 50/50 mixture of the two at each of five different pulse lengths: 1, 5, 10, 25, and 50 s. While a small sample, attempts were made to analyze these data.

Based on the simulation results, the data were normalized using auto-scaling. The data were then processed using each of the seven processing methods. Fig. 7 illustrates some of the PCA results.

As expected from such a small sample, the results were of limited value. However, it can still be seen that convolution and cross-convolution processing produced PCA plots where the group crossover was significantly reduced.

VI. CONCLUSION

These results show that characteristic signals, such as those produced through combinatorial functions (such as convolution, product and difference) are suitable for use in a data processing and pattern recognition system. The performance of classification systems using these signals was comparable, or better, than a traditional data processing methodology for a classic e-nose device.

However, while being highly resistant to additive SNR noise and sample variance, this system showed a weakness to baseline drift in sensor outputs. The common mode effect and high noise estimate of 5% has significantly overestimated this problem, as sensors are unlikely to drift by such a large amount in the period of seconds or minutes that such tests would normally be carried out for a time-series approach. For example, 2.4% drift per day for a classical e-nose is only 0.01% for a 6 min e-mucosa sample time.

In conclusion, we believe that convolution-based methods of processing can be used to improve the classification accuracy of not only novel e-mucosa type sensor systems but may even be applicable to conventional sensor-based e-nose technologies. In fact, the technique could even be applied to other time-based signals generated by metal oxide and other types of odor sensors.
Finally, our experimental results on simple odors (ethanol and toluene) indicate that using convolution-based methods can provide an improved image for classification, and work utilizing larger data sets will be presented in a separate article. The work extends to more interesting odors, such as essential oils and more complex odors, and ultimately tackling the odor segmentation problem.

REFERENCES


Julian W. Gardner (M’90–SM’02) received the B.Sc. degree from Birmingham University, Birmingham, U.K., in 1979, the Ph.D. degree from Cambridge University, Cambridge, U.K., in 1982, and the D.Sc. degree from Warwick University, Coventry, U.K., in 1997.

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