

HAUSSDORFF AND HELLINGER FOR COLORIMETRIC SENSOR ARRAY CLASSIFICATION

Tommy S. Alstrøm^a, Bjørn S. Jensen^a, Mikkel N. Schmidt^a, Natalie V. Kostesha^b, Jan Larsen^a

^aDept. of Informatics and Mathematical Modeling, Technical University of Denmark
Asmussens Allé 321, 2800 Kgs. Lyngby, Denmark
{tsal,bjje,mns,jl}@imm.dtu.dk

^bDept. of Micro- and Nanotechnology, Technical University of Denmark
Ørsted Plads 345 East, DK-2800, Kgs. Lyngby, Denmark
natalie.kostesha@nanotech.dtu.dk

ABSTRACT

Development of sensors and systems for detection of chemical compounds is an important challenge with applications in areas such as anti-terrorism, demining, and environmental monitoring. A newly developed colorimetric sensor array is able to detect explosives and volatile organic compounds; however, each sensor reading consists of hundreds of pixel values, and methods for combining these readings from multiple sensors must be developed to make a classification system. In this work we examine two distance based classification methods, K -Nearest Neighbor (KNN) and Gaussian process (GP) classification, which both rely on a suitable distance metric. We evaluate a range of different distance measures and propose a method for sensor fusion in the GP classifier. Our results indicate that the best choice of distance measure depends on the sensor and the chemical of interest.

Index Terms— Hausdorff distance, Hellinger distance, chemo-selective compounds, feature extraction, K -nearest neighbor classification, Gaussian Process Classification

1. INTRODUCTION

The development of rapid, reliable, and portable solutions for detection of chemical compounds is an important challenge with many possible applications including screening luggage and packages for explosives, detecting land mines, and monitoring the environment for hazardous compounds. In recent years a number of methods has been developed based on different technologies such as gas chromatography, Raman spec-

We acknowledge the support from the Danish Agency for Science and Technology's, Program Commission on Nanoscience Biotechnology and IT (NABIIT). Case number: 2106-07-0031 - Miniaturized sensors for explosives detection in air. Further we acknowledge assistant professor Raviv Raich, Oregon State University, for lending his code for distance calculation based on Hellinger distances using Parzen windows [1, 2].

trometry, mass spectrometry, ion mobility spectrometry and colorimetric sensors.

Colorimetric sensors can be used to detect a wide range of organic compounds in gas as well liquid phase [3–8]. In previous work a novel colorimetric sensor array developed by Kostesha et al. has been described [9, 10]. The sensor array proved useful for identifying a wide range of explosives as well as other compounds such as amines, cyanides, alcohols, arenes, ketones, aldehydes and acids. A colorimetric sensor contains a chemo-selective compound: A so-called dye which changes color when exposed to a target chemical compound denoted as analyte. The sensor is read by capturing a digital image of the dye before and after exposure to an analyte. Typically, the mean color change is used for detection; however, in earlier work it has been shown that accuracy can be improved by considering the complete distribution of color change for different image pixels [11]. An example of a colorimetric sensor is shown on Fig. 1. In this example the mean value is a particular bad choice as the mean color value is scarcely present in the sensor as shown on Fig. 1C. Furthermore, the sensor array comprises several colorimetric sensors, and detection could likely be improved further by combining information from multiple sensors.

In this paper, we present new methods for analyzing the output of a colorimetric sensor array using the complete distribution of color changes. We hypothesize that using the entire distribution of a sensor one will get an equal or better classification accuracy compared to a single feature representation such as the mean value. To classify a given analyte, we compare the de-facto method K -Nearest Neighbor (KNN) with a Gaussian process (GP) classifier. Both of these approaches rely on a measure of similarity between sensor readings—the KNN through a suitable distance measure and the GP through a covariance function. We motivate and compare several measures of similarity, demonstrating that a

proper choice of similarity measure can significantly improve classification accuracy using a single dye. Finally, we propose a multi kernel method using GP for fusing information from several sensors leading to superior performance compared to using a single sensor.

2. MATERIALS AND METHODS

In this section we describe the colorimetric sensor data and the proposed classification approach.

2.1. Notation

$N = 253$	Number of observations
$L = 3$	Number of channels (RGB)
$M = 31$	Number of dyes/sensors
$N_{\text{pix}} = [49; 1009]$	Pixels per dye/sensor
$\mathbf{x}_{l,n} = [\mathbf{x}_{1,n}^{\top}, \mathbf{x}_{2,n}^{\top}, \mathbf{x}_{3,n}^{\top}]^{\top}$	$1 \times N_{\text{pix}}$ vector of pixel values
$\mathbf{X}_n = [\mathbf{x}_{1,n}^{\top}, \mathbf{x}_{2,n}^{\top}, \mathbf{x}_{3,n}^{\top}]^{\top}$	$3 \times N_{\text{pix}}$ matrix with RGB pixel values
$\mathcal{X} = \{\mathbf{x}_n n = 1 : N\}$	a set of inputs
$\mathcal{Y} = \{y_n; \mathbf{x}_n n = 1 : N\}$	a set of labels for the given inputs

2.2. Colorimetric sensor data

The colorimetric sensor array consists of a number of chemo-selective compounds immobilized onto silica gel resulting in circular spots (Fig. 1A). Each individual sensor was approximately 3 mm in diameter with the total size of the sensor array of approximately 2.5 cm \times 4.0 cm.

The dataset used in this paper has been discussed in detail in earlier work [12] but is summarized here for completeness. The sensor array has been exposed to analytes belonging to various chemical families; acids (45), alcohols (27), amines (42), explosives (56) and other (83). The number in the parenthesis denotes the number of examples measured for the class in question. The group denoted other is not one chemical family but comprises various molecules that is used as reference measurements.

2.2.1. Data acquisition

The sensors were scanned using an ordinary flatbed scanner immediately after immobilization of dyes and then again after exposure of target analytes. The images are then pixel aligned, the dye locations are identified and finally pixel extraction is performed [12, 13]. The pixels are extracted by fitting a circular disc to the entire dye. Due to the chemistry of the sensor array often a distinct ring near the perimeter of the dyes appears (the coffee stain effect) and this area of the dyes is deemed unreliable. In order to accommodate for this effect a smaller area of a dye is used for feature extraction, corresponding to 2/3 of radius of the fitted circular disc [13]. Based on the pixel values features can be calculated, e.g. using the mean value, or distances between measurements.

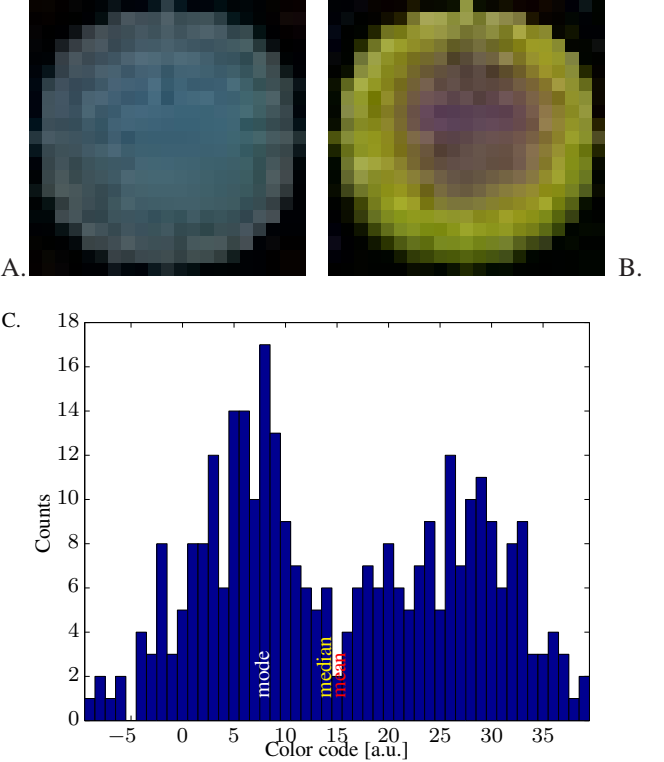


Fig. 1. An example of a specific dye of colorimetric sensor array exposed to the explosive analyte HMX. A: the sensor before exposure. B: the enhanced difference image. C: histogram of difference of the blue channel.

2.3. Classification Methods

2.3.1. K-Nearest Neighbor

The KNN is a simple yet effective classification technique [14] which works as follows. When testing a data point belonging to an unknown class, the distances to all points with known class labels are calculated. The classes of the closest K points are then identified and the unknown point is classified using majority voting. In order to carry out both model selection and estimation of the generalization error, double-cross validation using *leave-one-out* is performed. The inner loop is used to determine the value of K and the outer loop is used to estimate the generalization error.

2.3.2. Gaussian Process Classification

We now turn to a different but equally powerful classification framework based on a non-parametric Bayesian approach. The core of any probabilistic classification methods is a likelihood function modeling the likelihood of observing a specific outcome. Here we consider the cumulative Gaussian

(known as a probit model) defined as

$$p(y_n|f(\mathbf{X}_n)) = \int_{-\infty}^{y_n \cdot f(\mathbf{X}_n)} \mathcal{N}(t|0, 1) dt = \Phi(y_n \cdot f(\mathbf{X}_n)) \quad (1)$$

parameterized by a given a functional value, $f(\mathbf{X}_n)$. Hence, the model or free parameter is the functional value, $f(\mathbf{X}_n)$, and by taking a Bayesian approach, we can directly consider the posterior over the function defined by the finite set of random variables, $\mathbf{f} = [f(\mathbf{X}_1), f(\mathbf{X}_2), \dots, f(\mathbf{X}_N)]^\top$, i.e.,

$$p(\mathbf{f}|\mathcal{Y}, \mathcal{X}) = \frac{p(\mathcal{Y}|\mathbf{f}) p(\mathbf{f}|\mathcal{X})}{\int p(\mathcal{Y}|\mathbf{f}) p(\mathbf{f}|\mathcal{X}) d\mathbf{f}} = \frac{p(\mathcal{Y}|\mathbf{f}) p(\mathbf{f}|\mathcal{X})}{p(\mathcal{Y}|\mathcal{X})} \quad (2)$$

The natural prior for, $f(\cdot)$, is a Gaussian Process (GP) and we denote a function drawn from a GP as $f(\mathbf{X}) \sim \mathcal{GP}(\mathbf{0}, k(\cdot, \cdot)_{\theta_c})$ with a zero mean function, and $k(\cdot, \cdot)_{\theta_c}$ referring to the covariance function with hyper-parameters θ_c , defining the covariance between the random variables \mathbf{f} . The GP can thus be considered a distribution over functions, i.e., $p(\mathbf{f}|\mathcal{X}, \theta_c)$.

Given the probit likelihood model, the posterior over \mathbf{f} needs to be approximated and we resort to Expectation Propagation (EP) which provides a Gaussian approximation to the posterior [15]. The hyper parameters are approximated by point-estimates found by considering the marginal likelihood/evidence which can be optimized in regards to the hyper parameters by gradient methods. This possibility is a clear advantage over non-Bayesian methods.

The prediction for a new input \mathbf{X}^* is obtained by first computing the predictive distribution, $p(f^*|\mathcal{Y}, \mathcal{X}, \mathbf{X}^*)$, which is Gaussian due to the EP approximation. The probability of a given outcome y^* is computed by $P(y|\mathcal{Y}, \mathcal{X}, \mathbf{X}^*) = \int p(y|f^*) p(f^*|\mathcal{Y}, \mathcal{X}, \mathbf{X}^*) df^*$.

2.4. Distances and Covariance Functions

The two classification methods outlined above both require some notion of similarity either in the form of a distance function/metric or in the form of a covariance function. In the following we present and motivate a number of distance measures that can be used for measuring similarities between dyes.

2.4.1. Mean and “inner mean”

The traditional approach when using colorimetric sensors is to calculate the mean value of the response by averaging the color change over all pixels.

Since we have observed that pixel values in the outer edge of the dyes are less reliable, we propose to estimate an optimal radius instead of a fixed radius (typically $r=2/3$) and compute the mean in the central region within this radius. Assuming

that each dye should have a unique color change and such that any variation is due to noise, a reasonable assumption is to choose the radius that estimates the mean most accurately. This is done by choosing

$$\hat{r} = \arg \min_r \frac{1}{\sqrt{N_{\text{pix}}(r)}} \sigma(r), \quad (3)$$

where $N_{\text{pix}}(r)$ and $\sigma(r)$ denote the number of pixels and the standard deviation of pixel values within the central region with radius r . We denote this method “Inner Mean”.

2.4.2. Hellinger Distance

As an alternative to computing the average color change, we can consider the distribution of color change over the dye. The Hellinger distance measures similarity between two probability measures $f_i(\mathbf{x})$ and $f_j(\mathbf{x})$ and is given by

$$d_{\text{He}}(f_i, f_j)^2 = \int \left(\sqrt{f_i(\mathbf{x})} - \sqrt{f_j(\mathbf{x})} \right)^2 d\mathbf{x} \quad (4)$$

Using the Hellinger distance require us to choose how to represent the probability measures. We consider a nonparametric and a parametric approach: The first is a Parzen window Kernel density estimator which have been discussed in earlier work [11]:

$$f_i(\mathbf{x}) = \frac{1}{n_i} \sum_{k=1}^{n_i} K(\mathbf{x} - \mathbf{x}_{ik}) \quad (5)$$

where $K(\mathbf{x}) = 1/(2\pi\sigma^2)^{d/2} \exp(-||\mathbf{x}||^2/2\sigma^2)$ and the kernel width is set to $\sigma = 1$. The second approach is a multivariate normal distribution using a full covariance matrix

$$f_i(\mathbf{x}) = \frac{1}{(2\pi)^{d/2} |\Sigma_i|^{d/2}} \exp \left(-\frac{1}{2} (\boldsymbol{\mu}_i - \mathbf{x})^\top \Sigma_i^{-1} (\boldsymbol{\mu}_i - \mathbf{x}) \right) \quad (6)$$

where $\boldsymbol{\mu}_i$ is the mean and Σ_i the covariance matrix which is estimated by maximum likelihood.

2.4.3. The Hausdorff Distance

The Hausdorff distance measures distance between two point sets, and it is small if all points in each set are close to some point in the other set. This could be useful for comparing dye color changes that are not uniform over the dye. First define the distance between two points \mathbf{x}_i and \mathbf{x}_j as the Euclidean distance $d(\mathbf{x}_i, \mathbf{x}_j) = \|\mathbf{x}_i - \mathbf{x}_j\|_2$. The distance between a point \mathbf{x}_i and a set \mathbf{X}_j is then $d(\mathbf{x}_i, \mathbf{X}_j) = \min_{\mathbf{x}_j \in \mathbf{X}_j} d(\mathbf{x}_i, \mathbf{x}_j)$. The Hausdorff distance is defined as

$$d_{\text{Ha}}(\mathbf{X}_i, \mathbf{X}_j) = \max \left\{ \max_{\mathbf{x}_i \in \mathbf{X}_i} d(\mathbf{x}_i, \mathbf{X}_j), \max_{\mathbf{x}_j \in \mathbf{X}_j} d(\mathbf{x}_j, \mathbf{X}_i) \right\} \quad (7)$$

As an alternative approach one can use the modified Hausdorff distance which is more robust in the presence of noise and outliers

$$d_{\text{MH}}(\mathbf{X}_i, \mathbf{X}_j) = \max \left\{ \frac{1}{N_{\mathbf{X}_i}} \sum_{\mathbf{x}_i \in \mathbf{X}_i} d(\mathbf{x}_i, \mathbf{X}_j), \frac{1}{N_{\mathbf{X}_j}} \sum_{\mathbf{x}_j \in \mathbf{X}_j} d(\mathbf{x}_j, \mathbf{X}_i) \right\} \quad (8)$$

It should be noted that this distance is not a metric as the triangle inequality is not fulfilled [16].

2.4.4. From Distances to Covariance Functions

Given a distance metric $d(\mathbf{X}_i, \mathbf{X}_j)$ we use the distance substitution approach [17] based on a squared exponential kernel, $k(\mathbf{X}_i, \mathbf{X}_j) = \exp\left(-\frac{1}{\sigma_l^2}d(\mathbf{X}_i, \mathbf{X}_j)\right)$. We note that a valid covariance function may be constructed directly for the Hellinger distance by considering the inner product given by the integral which is known as the Probability Product Kernel [18]; however, to make a fair comparisons we treat the Hellinger distance like the other distance measures. Note also, that since the modified Hellinger distance is not a metric, the distance substitution kernel is not positive definite [17].

The use of kernels provides a convenient way of integrating information from different sensors by combining different kernels in a weighted sum. Thus, different dyes can be combined by constructing the following kernel

$$k(\mathbf{X}_i, \mathbf{X}_j) = \sigma^2 \mathbf{I} + \sum_{m=1}^M \alpha_m k(\mathbf{X}_i^m, \mathbf{X}_j^m) \quad (9)$$

where each kernel function is the distance substitution kernel with one of the respective metrics.

To fuse the difference dyes we employ a forward selection method using the following steps. 1) For a given dye, perform a grid search of hyper parameters σ^2 , α_m and σ_l . 2) Find the hyper parameters by optimizing the evidence in GP using the optimal point found in the grid as initial guess. 3) Perform *leave-one-out* cross validation to get classification error. 4) Choose the dye that yields the lowest classification error.

2.5. Evaluation

We apply nearest neighbor classifiers to each dye for each feature extraction technique in a one vs all setting. From earlier work [12] it was shown that the sensor is proficient in detecting among others acids, alcohols, amines and explosives so these are the classes we evaluate. The generalization error is estimated using LOOCV [19]. This scheme result in a total of 124 classifiers of each type per feature extraction method (31 dyes \times 4 classes).

Chemical	M	P	IM	MH	G	E[pFD]	n
Acids	1	2	2	6	0	0.89	11
Alcohols	0	0	0	0	3	0.15	3
Amines	1	0	1	0	0	1.02	2
Explosives	0	3	0	7	1	0.98	11

Table 1. Number of instances the best performing method is significantly better than another method. The methods are Mean, Parzen, Inner Mean, Mod. Hausdorff and Gaussian. E[pFD] is the expected positive false discoveries and n is the number of significant results.

To judge the differences in classification performance we use the McNemar significance test [20]. The McNemar is a paired test which uses the the number of cases where two classifiers disagree about a decision. From this test we calculate p -values for each comparison and use the multiple hypothesis framework proposed by Storey [21]. Based on the p -values we can calculate the expected positive false discoveries (E[pFD]) for our significant differences, that is, the expected quantity of wrongly significant results amount all found significant results.

3. RESULTS AND DISCUSSION

Initially we employ KNN and rank the dyes according to their best performance. Fig. 2 top panels show the dye ranking for KNN. The modified Hausdorff is the best performer for dyes 19 and 30 where it is significantly better than the single feature methods. Further the distribution methods are generally better or on par compared to the single feature methods.

Table 1 shows a summary of the McNemar significance tests. The overall best distance is the modified Hausdorff method although it is noteworthy that this method is not best even once for alcohols whereas is is best for numerous occasions for acids and explosives. Looking into the dyes reveal that the top dyes for acids largely overlap with the top dyes for explosives whereas for alcohols one of the top dyes is dye number 5. For this dye the Inner Mean is significantly better than any of the other methods. It should be noted that the majority of the differences is between one of the distribution methods and the single feature method, i.e. of the 52 significant results, only on 3 instances was the Parzen window worst and only 4 times where the Mod. Hausdorff method worst. Considering we have a total of 6 false discoveries we can only conclude that the distribution methods are significantly better than the single feature methods. To explore the effect of the methods deeper we fuse the dyes using GP.

Initially we want to establish the performance of GP in the same setting as KNN. Fig. 2 show the classification performance for GP when classifying explosives. Again the modified Hausdorff is the top performer for the first two dyes,

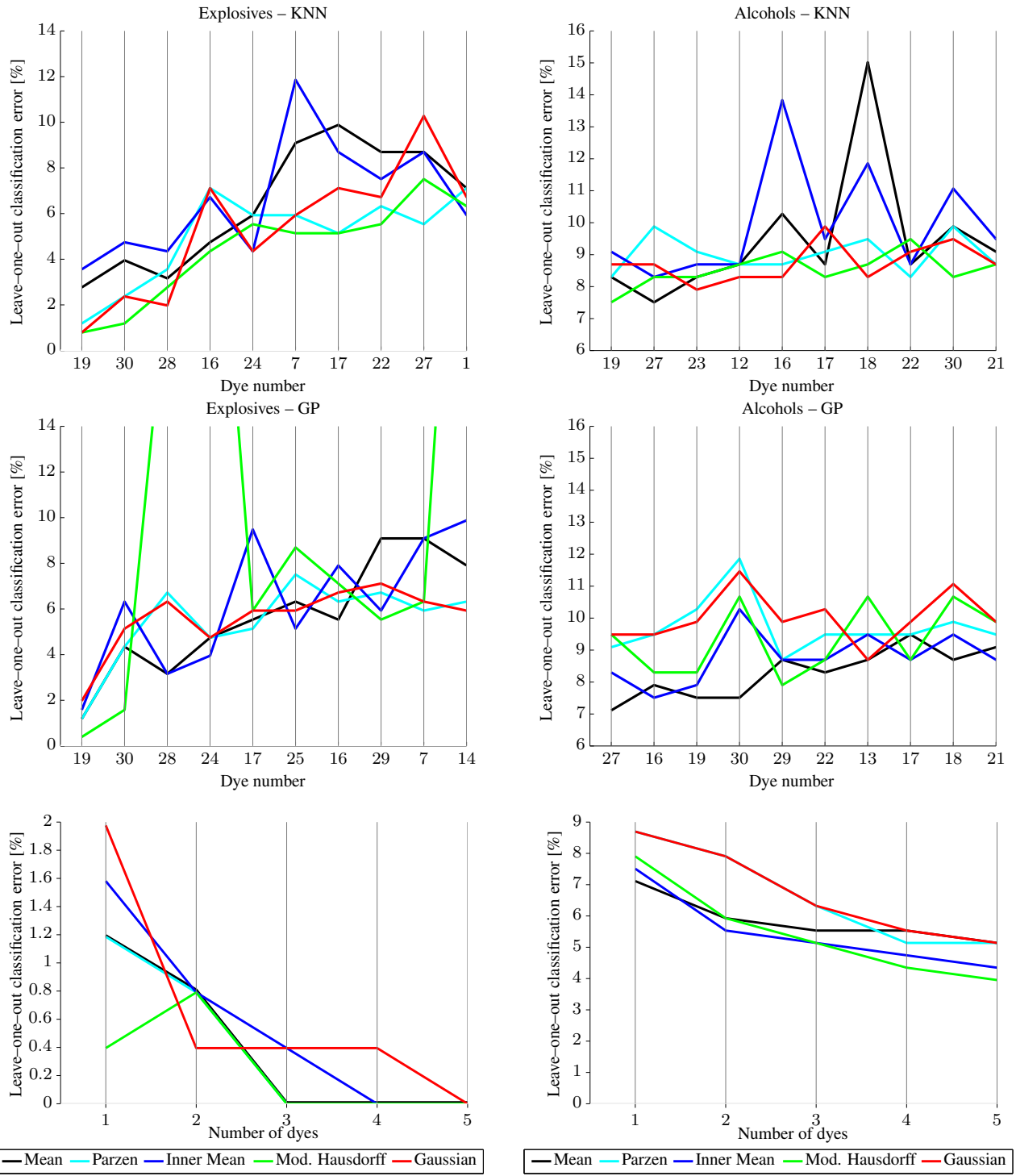


Fig. 2. Classification error for the different measures of similarity for GP and KNN. The top panels show the best performing dyes for explosives and alcohols using KNN. The middle panels show the best performing dyes using GP. The bottom panels shows the result of fusing the dyes together using GP. Bottom left is explosives and bottom right is alcohols.

which is again dye 19 and 30. However for dye 28 and 24 the modified Hausdorff coupled with GP yields an error of 21% and 30% respectively. This might very well be an effect of the fact that the modified Hausdorff is not a valid metric and as such the corresponding kernel might not be psd [16, 17]. But as the case of KNN, the figure does not show a clear indication of which method is superior although it seems that modified Hausdorff is able to capture more information about dye 19 and 30 than the the methods.

When fusing the dyes one by one using forward selection we find no significant results between the methods (Fig. 2). To get a significant result the performance must at least differ 2.4 percent points but already with one dye we are below that margin.

4. CONCLUSIONS

We have proposed three new methods for representation of sensory data in colorimetric sensor arrays, namely the Inner Mean, Hellinger distance using a Gaussian distribution and the Modified Hausdorff. Each method have it's merits. The Inner Mean seem to be particularly proficient for dye number 5 and 22 whereas the modified Hausdorff is especially strong for dye number 19 and 30. This suggests that none of the methods are capable of representing all the dye measurements optimally, although the methods that use the entire histogram are more often superior.

The modified Hausdorff method is the overall best performing method. It is particularly strong for classifying explosives, and since the distance calculation is based on sets of pixels and is parameter-free it can potentially work a lot better for high dimensional data where the Hellinger distance method might be inappropriate.

Finally, we have demonstrated that fusing the sensors can improve the classification error. GP classification effectively identifies which dyes should be included and generally including more dyes reduces the LOOCV error rate. In the fusing scheme the modified Hausdorff method also works well and is the overall best performer. Future work could be to combine all distance methods and learn the combination using Multiple Kernel Learning or evidence optimization instead of minimizing the classification error.

5. REFERENCES

- [1] K. M. Carter, R. Raich, W. G. Finn, and A. O. Hero, “Information preserving component analysis: Data projections for flow cytometry analysis,” *Selected Topics in Signal Processing, IEEE Journal of*, vol. 3, no. 1, pp. 148–158, 2009.
- [2] K. M. Carter, R. Raich, W. G. Finn, and A. O. Hero, “Fine: Fisher information nonparametric embedding,” *IEEE transactions on pattern analysis and machine intelligence*, pp. 2093–2098, 2009.
- [3] C. Zhang and K. S. Suslick, “A colorimetric sensor array for organics in water,” *Journal of the American Chemical Society*, vol. 127, no. 33, pp. 11548–9, Aug. 2005.
- [4] Kenneth S. Suslick, Neal A. Rakow, and Avijit Sen, “Colorimetric sensor arrays for molecular recognition,” *Tetrahedron*, vol. 60, no. 49, pp. 11133–11138, Nov. 2004.
- [5] N. Rakow, A. Sen, M. C. Janzen, J. B. Ponder, and K. S. Suslick, “Molecular recognition and discrimination of amines with a colorimetric array,” *Angewandte Chemie (International ed. in English)*, vol. 44, no. 29, pp. 4528–32, July 2005.
- [6] C. Zhang, D. P. Bailey, and K. S. Suslick, “Colorimetric sensor arrays for the analysis of beers: a feasibility study.,” *Journal of agricultural and food chemistry*, vol. 54, no. 14, pp. 4925–31, July 2006.
- [7] S. H. Lim, L. Feng, J. W. Kemling, C. J. Musto, and K. S. Suslick, “An Optoelectronic Nose for Detection of Toxic Gases.,” *Nature chemistry*, vol. 1, pp. 562–567, Sept. 2009.
- [8] C. Zhang and K. S. Suslick, “Colorimetric sensor array for soft drink analysis.,” *Journal of agricultural and food chemistry*, vol. 55, no. 2, pp. 237–42, Jan. 2007.
- [9] N. V. Kostesha, T. S. Alstrøm, C. Johnsen, K. A. Nielsen, J. O. Jeppesen, J. Larsen, M. H. Jakobsen, and A. Boisen, “Development of the colorimetric sensor array for detection of explosives and volatile organic compounds in air,” in *Proceedings of SPIE*, Apr. 2010, vol. 7673, pp. 76730I–76730I–9.
- [10] N. V. Kostesha, T. S. Alstrøm, C. Johnsen, K. A. Nielsen, J. O. Jeppesen, J. Larsen, A. Boisen, and M. H. Jakobsen, “Multi-colorimetric sensor array for detection of explosives in gas and liquid phase,” in *Proceedings of SPIE*, 2011, pp. 80181H–80181H–12.
- [11] Tommy S. Alstrøm, Raviv Raich, Natalie V. Kostesha, and Jan Larsen, “Feature extraction using distribution representation for colorimetric sensor arrays used as explosives detectors,” in *IEEE International Conference on Acoustics, Speech, and Signal Processing*, 2012.
- [12] Tommy S. Alstrøm, Jan Larsen, Natalie V. Kostesha, Mogens H. Jakobsen, and Anja Boisen, “Data representation and feature selection for colorimetric sensor arrays used as explosives detectors,” *IEEE International Workshop on Machine Learning for Signal Processing*, pp. 1–6, 2011.
- [13] T. S. Alstrøm and J. Larsen, “Feature Extraction and Signal Representation for Colorimetric Sensor Arrays,” Tech. Rep., DTU Informatics, 2011.
- [14] Christopher M. Bishop, *Pattern Recognition and Machine Learning*, Springer, Secaucus, NJ, USA, 2006.
- [15] Carl Edward Rasmussen and Christopher K. I. Williams, *Gaussian processes for machine learning*, Apr. 2006.
- [16] M. P. Dubuisson and A. K. Jain, “A modified hausdorff distance for object matching,” in *12th International Conference on Computer Vision Image Processing*, oct 1994, vol. 1, pp. 566–568.
- [17] Bernard Haasdonk and Claus Bahlmann, “Learning with distance substitution kernels,” in *in Pattern Recognition - Proc. of the 26th DAGM Symposium*, 2004.
- [18] T. Jebara and A. Howard, “Probability Product Kernels,” *Journal of Machine Learning Research*, vol. 5, pp. 819–844, 2004.
- [19] Lars K. Hansen and Jan Larsen, “Linear unlearning for cross-validation,” *Advances in Computational Mathematics*, vol. 5, pp. 269–280, 1996.
- [20] Quinn McNemar, “Note on the sampling error of the difference between correlated proportions or percentages,” *Psychometrika*, vol. 12, no. 2, pp. 153–157, 1947.
- [21] John D. Storey, “A direct approach to false discovery rates,” *Journal of the Royal Statistical Society: Series B (Statistical Methodology)*, vol. 64, no. 3, pp. 479–498, 2002.