Active Temperature Programming for Metal-Oxide Chemoresistors

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Abstract—Modulating the operating temperature of metal-oxide (MOX) chemical sensors gives rise to gas-specific signatures that provide a wealth of analytical information. In most cases, the operating temperature is modulated according to a standard waveform (e.g., ramp, sine wave). A few studies have approached the optimization of temperature profiles systematically, but these optimizations are performed offline and cannot adapt to changes in the environment. Here, we present an “active perception” strategy based on Partially Observable Markov Decision Processes (POMDP) that allows the temperature program to be optimized in real time, as the sensor reacts to its environment. We characterize the method on a ternary classification problem using a simulated sensor model subjected to additive Gaussian noise, and compare it against two “passive” approaches, a naïve Bayes classifier and a nearest neighbor classifier. Finally, we validate the method in real time using a Taguchi sensor exposed to three volatile compounds. Our results show that the POMDP outperforms both passive approaches and provides a strategy to balance classification performance and sensing costs.

Index Terms—Active sensing, hidden Markov models, metal-oxide (MOX) sensors, partially observable Markov decision processes (POMDP).

I. INTRODUCTION

Metal-Oxide (MOX) gas sensors are robust, inexpensive, and highly sensitive, but have poor specificity. Interestingly, modulating the temperature of a MOX sensor during exposure to volatile chemicals gives rise to gas-specific temporal signatures that may be used to enhance selectivity [1]. The idea of temperature modulation for MOX sensors has been around for more than two decades. Researchers have used a wide array of temperature waveforms, but most of these studies have been empirical, see, e.g., [2] and [3]. To our knowledge, only two studies have proposed systematic approaches to optimizing temperature profiles [4], [5], though both methods require that the optimization be performed offline.

In this paper, we propose an active-sensing approach that can optimize the temperature profile online, that is, as the sensor collects data from its environment. Our approach consists of modeling the dynamic response of a sensor to a sequence of temperature pulses by means of an Input-Output Hidden Markov Model (IOHMM) [6], an extension of the Hidden Markov Models commonly used in speech recognition; see [7] for a gentle introduction to these models. IOHMMs can be used to learn a dynamic mapping between two data streams, an input (temperature in our case) and an output (sensor conductance).

Given a learned IOHMM model, we approach temperature optimization process as one of sequential decision making under uncertainty where the system must balance the cost of applying additional temperature steps against the risk of (mis)classifying the chemical analyte based on the available information. This is solved through a partially observable Markov Decision process (POMDP); see [8] for a good introduction. In a previous conference paper [9], we evaluated this active-sensing approach on a toy binary classification problem using a simulated MOX sensor model. Here, we thoroughly characterize the approach through a series of simulations, compare it against two passive classification approaches, and finally validate it experimentally on a commercial gas sensor.

II. RELATED WORK

Over a decade ago, Kunt et al. [4] proposed a systematic approach for optimizing the temperature profile of MOX sensors. The approach consisted of two stages: modeling and optimization. During the modeling stage, a wavelet network was used to learn the sensor dynamics from experimental data. The model was able to predict the next conductance value of the sensor from its previous values and from the previous and next temperature values. During the second stage, an optimization routine was then used to find a program that maximized the distance between the temperature-modulated sensor responses to two target gases, as predicted by the wavelet network model. The optimization was subject to a smoothness constraint to avoid drastic changes between consecutive temperatures. When applied to the discrimination of methanol and ethanol using a microhotplate device, the optimization method returned a temperature program such that the sensor response to the two analytes was nearly orthogonal.

More recently, Vergara et al. [10] proposed a system-identification method for determining suitable temperature-modulation frequencies. Their method was based on pseudorandom binary sequences (PRBS) and maximum length sequences (MLS), square-wave signals with several interesting properties: they are repeatable, which ensures reproducibility; they have a flat power spectrum over a large frequency range, which makes them suitable for system identification; and they have maximum length, so the impulse response of the system can be estimated from the cross-correlation. In this method, a PRBS-MLS is used to drive the sensor heater while the sensors are exposed to various target compounds. For each individual target, the impulse response \( h(t) \) is computed as the cross-correlation between the
excitation signal (PRBS) and the sensor response, and the spectral components are computed from the Fast Fourier Transform of $h(t)$. Finally, each individual frequency is ranked based on its information content (measured as the ratio of between- to within-class scatter), and a subset of the most informative frequencies is selected. Using this procedure, a single sensor at three modulating frequencies was sufficient to discriminate and quantify various gases and their mixtures.

Both of these landmark studies, however, require that the temperature profiles be optimized offline. Here, we propose an optimization approach that operates online, that is, as the sensor collects data from its environment. Our method borrows from literature in the fields of robotics and computer vision, where active sensing has had a long tradition [11]–[13].

### A. Active Sensing

Active sensing strategies are inspired by the fact that perception [14] is not a passive process but an active one in which an organism controls its sensory organs in order to extract information from the environment (see Fig. 1). A recurrent theme in computer vision has been the recognition of three-dimensional (3-D) objects from 2-D images, a problem where active vision may be used to select camera views to reduce ambiguities introduced by the imaging process [15], [16]. An early theoretical result [12] shows that active vision provides tractable solutions to several problems (e.g., shape from shading, contour, and texture, and structure from motion) that are ill-posed with a passive vision strategy. Other applications of active vision include object recognition, target tracking, and scene reconstruction [17]. Within the robotics community, active sensing strategies are commonly used in localization and navigation tasks, where there is a tradeoff between the immediate rewards of actions (e.g., bringing the robot closer to its goal) and long-term effects (e.g., gathering information to avoid getting lost in the way) [18]–[20].

Active sensing can simplify the detection/recognition part of sensing, but requires a strategy to select an appropriate sequence of sensing actions. A number of action selection methods have been proposed, which can be broadly grouped into two categories: probabilistic and behavioral [21]. Probabilistic approaches view active sensing as a state estimation problem: the system is to keep a belief state (i.e., a probability distribution over all possible states) and select a suitable sequence of actions from it [16], [22]. Behavioral methods, on the other hand, do not differentiate between belief updates and action selection, and instead learn a controller (e.g., a neural network) that is optimized to solve the task at hand.

In contrast with the wealth of work in robotics and vision, active sensing has only received minimal attention in chemical sensing. In one of the earliest studies, Nakamoto et al. [23] developed a method for active odor blending, where the goal was to reproduce an odor blend by creating a mixture from its individual components. The authors developed a control algorithm that adjusted the mixture ratio so that the response of a gas sensor array to the mixture matched the response to the odor blend. More recently, Priebe et al. [24] developed a statistical pattern recognition method for active sensing based on the concept of Integrated Sensing and Processing (ISP) [25]. Given a feature vector, the method builds a decision tree that partitions feature space hierarchically; nodes close to the root of the tree select features based on their ability to cluster examples regardless of class labels, whereas nodes at the leaves select sensors based on their ability to discriminate examples from different classes. The model was evaluated on an array of 19 optical sensors exposed to trichloroethylene (a carcinogenic industrial solvent) in complex backgrounds. The ISP method was able to reduce misclassifications by half, while requiring only 20% of the sensors.

Active sensing has also received attention in recent years as a resource-allocation method in multitarget and multisensor applications [26]. Here, one seeks to find an optimal schedule for a sensor (or group of sensors) that minimizes feature acquisition costs and maximizes target identification accuracy. The work proposed here follows this problem formulation; namely, we seek to minimize the number of temperature steps applied to a MOX sensor, while maximizing the accuracy with which each specific target analyte is classified.

### III. METHODS

Consider the problem of classifying an unknown gas sample into one out of $M$ known categories $\{\omega^{(1)}, \omega^{(2)}, \ldots, \omega^{(M)}\}$ using a MOX sensor with $D$ different operating temperatures $\{\rho_1, \rho_2, \ldots, \rho_D\}$. To solve this problem, one typically measures the sensor’s response at each temperature, and then analyzes the complete feature vector $x = [x_1, x_2, \ldots, x_D]^T$ with a pattern-recognition algorithm [27]. Though straightforward, this “passive” sensing approach is unlikely to be cost-effective because only a fraction of the measurements is generally necessary to classify the chemical sample. Instead, in active classification we seek to determine an optimal sequence of actions $a = [a_1, a_2, \ldots, a_T]$, where each action corresponds to setting the sensor to one of the $D$ possible temperatures (or terminating the process by assigning the sample to one of the $M$ chemical classes). More importantly, we seek to select this sequence of actions dynamically, based on accumulating evidence. This process is illustrated in Fig. 2.

#### A. Modeling the Sensor Dynamics

Following [28], we model the sensor’s steady-state response at temperature $\rho_i$ to chemical $\omega^{(c)}$ with a Gaussian mixture:

$$p(x_i | \omega^{(c)}) = \sum_{m_i=1}^{M_i} \sigma_{i,m_i}^{(c)} N(x_i | \mu_{i,m_i}^{(c)}, \Sigma_{i,m_i}^{(c)})$$  \hspace{1cm} (1)

where $M_i$ is the number of Gaussians, and $\sigma_{i,m_i}^{(c)}, \mu_{i,m_i}^{(c)}, \Sigma_{i,m_i}^{(c)}$ are the mixing coefficient, mean vector, and covariance matrix of each Gaussian for class $\omega^{(c)}$, respectively. Given a sequence of actions $[a_1, a_2, \ldots, a_T]$, we assume that the sensor progresses
that leads to an at state, but to produce an observation. With the exception of the cost, these POMDP parameters can be obtained directly from the IOHMM as follows:
- Initial belief: \( b_0(s) = p(\omega^c)\pi(0)(s) \), for \( s \in S^c \).
- State transition: \( T(s'|s,a) = \tau(s') \), for \( s,s' \in S^{(u)} \); zero otherwise.¹
- Observation function: \( \Omega(o|s) = \varphi(0)(o|s) \), for \( s \in S^c \).

The POMDP stores information about the system in a belief state \( b_T(s) \), a probability distribution (across states from all the IOHMMs) given the initial belief \( b_0(s) \) and the history of actions \( [a_1 \ldots a_T] \) and observations \( [o_1 \ldots o_T] \):

\[
b_T(s) = p(s|a_1 \ldots a_T, o_1 \ldots o_T, b_0) = p(s|a_T, a_T, b_T-1).
\]

Since \( b_T(s) \) is a sufficient statistic, it can be updated incrementally from its previous estimate \( b_T-1(s) \) by incorporating the latest action \( a_T \) and observation \( o_T \):

\[
b_T(s') = \frac{p(o_T|s', a_T, \sum_{s''} s''|a_T, b_T-1) b_T-1(s'')}{p(o_T|a_T, b_T-1)}
\]

where the denominator \( p(o_T|a_T, b_T-1) \) can be treated as a normalization term to ensure that \( b_T(s') \) sums up to 1, and all terms in the numerator are known from the POMDP model.

Using this POMDP formulation, the active-classification problem becomes one of finding a function that maps belief states into actions (\( b_T(s) \rightarrow a \)) so as to minimize the expected risk; this function is normally referred to as a policy. We consider two types of actions and their associated costs:
- Sensing actions \( (a = \rho_i) \), which correspond to applying temperature \( \rho_i \). The cost of sensing actions is \( c(s,a = \rho_i) = c_{\rho_i} \), which reflects the fact that certain temperatures may be more expensive (e.g., draw more power).
- Classification actions \( (a = \hat{\rho}_u) \), which assign the sample to a particular class; classification actions are terminal. The cost of classification actions is \( c(s,a = \hat{\rho}_u) = c_{\text{class}}(\forall s \in S^{(u)}) \), which represents a misclassification penalty whenever \( u \neq v \).

C. Finding a Sensing Policy
Finding an exact solution for a POMDP policy is P-SPACE complete (i.e., the hardest class among all problems that can be solved in polynomial space) and therefore intractable for most applications [30]. Moreover, for a standard POMDP solution repeated actions are permissible (i.e., measuring the sensor response at the same temperature multiple times), which is undesirable in our case. For these reasons, we employ a myopic policy [28] that only takes sensing action if the cost of sensing \( c_{\rho_i} \) is lower than the expected future reduction in Bayes risk.

¹This ensures that transitions across IOHMMs are not allowed, since we assume that the chemical stimulus does not change over time.
Given belief state $b_T(s)$, the expected risk of a classification action is
\[
R_C(b_T(s)) = \min_u \sum_v c_{av} \sum_{s \in S(v)} b_T(s)
\]
where $u$ represents the class with minimum Bayes risk, and $\sum_{s \in S(v)} b_T(s)$ is the belief across all states that belong to chemical $v$. In turn, the expected risk of a sensing action is
\[
R_S(b_T(s), a) = \sum_{\forall \alpha} \min_u \left( \sum_v c_{av} \sum_{s' \in S(v)} p(a | s', a)p(s' | s, a)b_T(s) \right)
\]
which averages the minimum Bayes risk over all observations that may result from the sensing action. Hence, the utility of sensing action $a$ can be computed as
\[
U(b_T(s), a) = [R_C(b_T(s)) - R_S(b_T(s), a)] - c_a.
\]
If $U(b_T(s), a)$ is negative for all sensing actions, then the cost of sensing $c_a$ exceeds the expected reduction in risk $[R_C(\cdot) - R_S(\cdot)]$, and a classification action is taken. Otherwise, the action with maximum utility is taken.

D. Continuous Observation Space

The expected Bayes risk as computed in (5) is applicable only for discrete observation spaces. In the case of continuous features, the summation in (5) becomes a difficult integration [28], so instead we discretize the observation space at each temperature using $k$-means clustering. This results in a finite set of bins with sorted centers $\{\bar{\Phi}_{pi,1}, \bar{\Phi}_{pi,2}, \ldots, \bar{\Phi}_{pi,k}\}$ for each sensing action $a = p_i$, from which the likelihood of $p_{i, ij}$ at state $s'$ in $S$ is approximated as
\[
p(a | s', a) = p(\bar{\Phi}_{pi, a} | s') = \int_{(\bar{\Phi}_{pi, j-1} + \bar{\Phi}_{pi, j} + 1)/2}^{(\bar{\Phi}_{pi, j} + \bar{\Phi}_{pi, j+1})/2} N(x | \mu_{s'}, \Sigma_{s'})
\]
where $\mu_{s'}$ and $\Sigma_{s'}$ are the mean and variance of the Gaussian distribution of state $s' \in S$. This probability is then used in (5) to estimate the expected risk of sensing actions. The integration is performed numerically with a trapezoidal rule.

IV. MODEL CHARACTERIZATION ON SYNTHETIC DATA

We first validated the active sensing model using a simulated MOX sensor. This allowed us to characterize the model with different levels of additive noise. Experimental validation on an actual MOX sensor is described in Section V.

A. Simulated Sensor Model

The response of MOX sensors is influenced by various factors such as operating temperature, concentration of oxygen species, humidity, film thickness, grain size, doping, and catalyzers [31, 32]. A variety of nonlinear models have been proposed to estimate the sensor conductance as a function of temperature, assuming all other influential parameters are fixed [33]–[35]. Here, we follow a phenomenological model developed by Wlodek et al. [36], where the change in conductance in response to a temperature cycle is approximated as a sum of Gaussian densities; this is a sensitive choice since Gaussian mixture models are universal approximation functions. Following [37], we also model the sensor dynamics using a low-pass filter, resulting in
\[
G(T(t)) = k_1G(T(t - 1)) + (1 + \alpha)(k_2e^{-(T(t) - T_0)/\sigma})^2 + k_2T(t) + G_0
\]
where $T(t)$ is the sensor temperature at time $t$, and $G(T(t))$ is its conductance at $T(t)$. $T_0$ is the temperature at which the sensor conductivity is maximum, $k_1$, $k_2$ and $\sigma$ are parameters that capture the steady-state properties of the sensor, $G_0$ is the baseline measurement (usually made when the sensor is exposed to air) and $\alpha$ captures history effects.

B. Classification Performance Versus Sensing Costs

We simulated a scenario with three chemical targets and a sensor with 20 different temperature settings. Sensor parameters were set as follows: $\alpha = 0.25$, $k_1 = 6.0$ and $k_2 = 3.5$ for all classes; $\sigma = 6$ and $T_0 = 9$ for $\omega^{(1)}$; $\sigma = 6$ and $T_0 = 11$ for $\omega^{(2)}$; and $\sigma = 8$ and $T_0 = 10$ for $\omega^{(3)}$; $G_0$ was fixed at 1.0. The temperature-dependent response of the sensor to the three chemicals is shown in Fig. 3(a). These responses were obtained by running the sensor with a random temperature sequence and recording the corresponding responses. Thus, the spread at each temperature illustrates the effect of the sensor dynamics (i.e., history effects).

Training data for each analyte was generated using 20 random temperature sequences, each sequence containing 20 temperature pulses. Three IOHMMs (one per analyte) were trained on this data; the number of Gaussian components per temperature in (1) was set to $M_f = 4$. The model was tested on 60 samples, 20 from each class. Each sample was generated by randomly selecting an initial temperature $T(0)$ unknown to the system, and initializing the sensor response to
\[
G(T(0)) = k_1e^{-(T(0) - T_0)/\sigma})^2 + k_2T(0) + G_0
\]
To test the POMDP, we assumed a zero-one loss function for classification costs ($c_{av} = 1$ if $u \neq v$; zero otherwise) and varied sensing costs from $c_i = 0.025$ to $c_i = 1$ in increments of 0.025; we also initialized each belief state $b_T(s)$ with a constant value. Fig. 3(b) shows the average classification rate and length of the temperature sequence as a function of feature acquisition costs. For the lowest sensing cost ($c_i = 0.025$), the

![Fig. 3. (a) Conductance versus temperature for three chemical classes. For visualization purposes, the conductance values of chemicals 2 and 3 were shifted along the x-axis by 0.1 and -0.1, respectively. (b) Classification rate and average sequence length as a function of feature acquisition costs.](image-url)
The system achieves 100% classification accuracy with an average sequence of 2.3 temperatures, but the same classification rate can be achieved with a shorter sequence (an average of 1.8 steps) by increasing the sensing costs to $c_i = 0.30$. The classification rate gradually decreases to 83.33% for $0.30 < c_i \leq 0.475$, after which classification rates settle in the range of 83.33%, or essentially what can be achieved with a single temperature. Finally, for sensing costs greater than $c_i = 0.675$, the system acquires no features and produces a chance-level classification rate of 33%; at this point, sensing costs have become too high compared to misclassification costs.

C. Comparison With a Passive Sensing Approach

We evaluated the POMDP using sensor measurements with additive white Gaussian noise, and compared it against two “passive” classification schemes: a nearest neighbor classifier (NN) [38] and a naïve Bayes (NB) classifier [39]. The latter was obtained directly from the IOHMM as

$$p(\omega^{(c)}|o_1 \ldots o_T) = p(\omega^{(c)}) \prod_{t=1}^{T} \sum_{j=1}^{m} p(\omega^{(c)}(o_t)|s_j)$$ (10)$$

where $m$ is the total number of states in the IOHMM for class $\omega^{(c)}$. We estimated the average classification rate of the three classifiers at increments of 0.1 in noise variance; feature acquisition costs were set to $c_i = 0.025$. At each noise level, we ran the NN and NB classifiers multiple times with the same number of temperatures so that (on average) they used the same number of temperatures as those selected by the POMDP at the corresponding noise level; this ensured a fair comparison among the three classifiers. Features for the NN classifier were obtained through sequential forward feature subset selection [40]; the individual feature subsets were evaluated based on classification performance, i.e., a wrapper approach [41]. In contrast, features for the NB classifier were determined by ranking each temperature according to its information content (i.e., a filter approach), measured as the largest eigenvector of a Fisher’s linear discriminant analysis (LDA). As an example, the distribution of LDA eigenvalues shown in Fig. 4(a) indicates that temperatures 4, 7, and 17 have the most discriminatory information, respectively; these three temperatures would then be selected when training a NB classifier with three features. All the models (IOHMM, feature subset selection, NB, and LDA) were trained on noise-free data (60 sample vectors, 20 from each class), and then tested on noisy data (300 test cases, 100 from each class). Each training and test case was generated by initializing the sensor to a randomly generated temperature, as described in (9).

Results are shown in Fig. 5(d). As expected, classification performance for the three classifiers degrades with increasing levels of noise. The active classifier consistently outperforms the two passive classifiers, and is also able to maintain near-optimal performance (100%) for low noise levels. Though both passive classifiers always use the most informative features (as estimated from training data), they are unable to adapt when observations become noisy. In contrast, the active classifier selects features adaptively, and therefore is able to select features that complement evidence accumulated from previous noisy measurements.

D. Active Feature Selection

To further understand how the active classifier was able to outperform two passive classifiers, we analyzed the feature selection strategy of the POMDP. Fig. 5(a)–(c) shows the probability of each temperature being selected regardless of the chemical input at three noise levels (0, 0.5, and 1.0). In the first iteration, the POMDP selects the action with the lowest expected risk, which is computed as the average of the minimum Bayes risk over all observations that may result from the sensing action [see (5)]. Therefore, the first temperature selected by the POMDP is fixed and independent from the measurement noise. Based on LDA, we expected feature 4 to be the first temperature to be selected by the POMDP, since it has the maximum discriminatory information. In contrast, the POMDP selected feature 19 in the first iteration. Inspection of the variance at each temperature, shown in Fig. 4(b), indicates that feature 19 has very low variance for chemical 3. This suggests that the POMDP uses this feature to discriminate chemical 3 from the other two. While the NB classifier is constructed from the IOHMM emission densities and therefore may be able to exploit variance information as well, its features are selected based on LDA, which only captures discriminatory information in the mean of the data but not in its variance.

We also analyzed the feature selected by the POMDP in the second iteration, i.e., after it has acquired feature 19. When no noise is added to the measurements and chemical 1 is present, the POMDP selects feature 7 with 92% probability in the second
iteration; chemical 1 has the lowest variance at that temperature. When chemical 2 is present, the POMDP selects feature 4 with 94% probability in the second iteration. Though chemical 2 has a high variance at feature 4, the average sensor response to chemical 2 is clearly distinct from that to the other two chemicals; see Fig. 3(a). When chemical 3 is present, the POMDP selects feature 4 with 100% probability in the second iteration; chemical 3 has the lowest variance at that temperature. These results show that the POMDP selects the second feature depending on evidence acquired from the first measurement. Similar interesting trends may be observed in later iterations. When observations are subjected to noise with variance of 0.5, the POMDP selects features 19, 7, 4, and 17 most often, but also selects other less informative temperatures to complement the inaccurate evidence obtained from the noisy measurements. Similar trends are observed at higher noise levels. Features 9, 10, and 11 are the least informative since the sensor response to the three chemicals is similar at these temperatures [see Fig. 3(a)]; in consequence, these temperatures are rarely selected by the POMDP.

V. EXPERIMENTAL VALIDATION

We validated the active sensing model using a commercial MOX sensor (TGS2600; Figaro USA, Inc.) exposed to three different analytes (acetone, ammonia, and isopropyl alcohol). As an approximation, we used the heater voltage $V_{TH}$ as a proxy for the sensor’s operating temperature. The sensor was introduced in 30 ml glass vials through a tight aperture on the cap and allowed to equilibrate with the static headspace of 10 ml of the analyte. The sensor was interfaced through a National Instruments data acquisition card (USB 6009) controlled using Matlab. Each analyte was serially diluted (1/3 dilution ratio) to determine a set of dilutions at which the sensor provided similar isothermal responses for all the analytes; this ensured that the analytes could not be discriminated without temperature modulation. To obtain the isothermal response, the sensor was exposed to each analyte for 100 seconds under a constant heater voltage of 5 V. The final dilution factors were 1/3 for ammonia (one serial dilution), 1/243 for acetone (five serial dilutions), and 1/2187 for isopropyl alcohol (seven serial dilutions).

We used eight heater voltages (i.e., sensing actions) ranging from 1 to 8 V. Fig. 6(a) shows the sensor’s transient response to the three analytes for a random temperature sequence; each step was 40 s long. Information from each transient was reduced to a single observation by computing its integral response [42]. Fig. 6(b) shows the final temperature-conductivity profile, obtained by recording the sensor response to ten random voltage sequences (eight different voltage pulses per sequence) and computing the average response at each temperature. Therefore, the spread at each temperature reflects history effects.

We trained a separate IOHMM for each analyte. Fig. 7(a) shows IOHMM prediction results when the sensor was exposed to acetone and driven with a random temperature sequence; prediction results for the other two targets were comparable. The IOHMMs explain a significant proportion of the variance in the response of each sensor: 90.4% for acetone, 92.1% for ammonia, and 90.1% for ISP. IOHMM predictions were shown to be less accurate at lower voltages: 1 and 2 V for acetone, 1–3 V for ammonia, and 1 V for ISP. These results are consistent with the fact that the sensor response has higher variance at low temperatures; see Fig. 6(b). Moreover, IOHMM prediction errors were significantly higher for large negative temperature steps ($\Delta \leq -5$ V), a reasonable result considering that cooling is passive (e.g., limited by heat dispersion). Despite these estimation errors, the POMDP selects meaningful temperature sequences that predict the chemical class accurately, as we will see next.

To evaluate the classification accuracy of the POMDP, we assumed uniform misclassification costs ($c_{uv} = 1$ if $u \neq v$; zero otherwise) and varied sensing costs in the range from $0 < C_s \leq 0.5$ in increments of 0.05. We ran the active-sensing algorithm five times for each analyte, resulting in 15 test cases at each cost setting. Fig. 7(b) shows the classification rate and average number of temperature steps selected by the POMDP as a function of sensing costs. Results are consistent with those obtained on the simulated sensor model: as feature acquisition costs increase, the POMDP selects fewer temperature steps at the expense of classification rate. In contrast with the simulation results in Fig. 3(b), however, these results show a more graceful degradation in classification performance as a function of the sensing costs.

VI. DISCUSSION

We have presented a method to optimize the temperature program of MOX sensors in real time, as the sensor reacts to its environment. The approach consists of building a dynamical model of the sensor by means of an IOHMM, and casting the resulting model into a POMDP. Our results show that IOHMMs can capture the conductance-temperature dependence and thermal dynamics of MOX sensors, and that the POMDPs
outperforms two “passive” classification approaches, a NN rule and a NB rule. The latter result is significant considering that the two passive classifiers were aided by a feature subset selection stage that identified the best features, whereas the POMDP operated on the entire feature space (when comparing methods, the average number of features used by each classifier was the same). The NB classifier was obtained directly from the IOHMM densities; thus, the main difference between NB and POMDP is that the latter takes state transition probabilities into consideration, which allows it to account for the sensor dynamics. This factor, combined with its adaptive feature selection capabilities, explains why the POMDP is able to consistently outperform the NB classifier (as well as the NN classifier). To this end, our results indicate that the POMDP not only selects features with high discriminatory information but also features that have low variance. As a result, the POMDP is able to exploit information both in the mean and in the variance of each class.

IOHMM predictions on experimental data indicate that the model is less accurate at lower temperatures, when features are more sensitive to environmental influences. Though it is possible that the number of Gaussian components in our implementation ($m = 4$) was insufficient to model the sensor dynamics, prediction errors could also be due to insufficient training data, since our training set consisted of only ten random temperature sequences (each temperature occurring once in a sequence), and did not contain every possible temperature transition. Similar problems with data sparsity are quite common in speech recognition, the domain where HMMs are most commonly used.

A. Future Work

Our work has focused on real-time prediction, but the models may also be used to generate temperature profiles offline in a more principled fashion than through trial-and-error of various waveforms (e.g., sine waves, saw tooth), and more along the line of signal exploitation methods in [4] and [10]. IOHMMs are an example of a dynamic Bayesian network with one discrete random variable (input voltage) and one continuous random variable (sensor response), but the model could be extended to incorporate additional variables. As an example, chemical concentrations could be added as a continuous random variable. The resulting Bayesian network could then model conductance-temperature and conductance-concentration relations simultaneously. At the time of this writing, we are working on extensions of the model to handle sensor drift, using a dynamic Bayesian network, drift can be modeled as a hidden variable and predicted using probabilistic inference. Interference from other gases and the effects of humidity may be incorporated in a similar fashion.

In this study, we assumed uniform feature acquisition costs regardless of operating temperature, but variable sensing costs could be used to penalize higher temperatures and thus reduce power consumption. There is a great potential for deploying chemical sensors on large-scale networks [43], but such deployments are constrained due to the lack of power-aware, autonomous sensors. The active sensing approach proposed here provides a strategy for implementing energy-aware chemical sensing networks using low-cost commercial sensors.

REFERENCES


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