A MEMS-Based Approach that Uses Temperature-Dependent Sensing Responses to Recognize Chemical Targets in Untrained Backgrounds

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Abstract—A major practical challenge for solid state microsensors is the detection of trace chemical species over time and in complex gas-phase backgrounds. We describe a MEMS-based, chemiresistive technology that has succeeded in such problems by combining oxide nanomaterials on microscale platforms, acquisition of dense temperature-dependent response data streams, and novel signal processing methods. Unlike the operation and analysis employed with many electronic noses, our higher dimensional approach captures surface electronic implications of changing adsorptive/reactive phenomena caused by rapid thermal cycling. Here, we demonstrate new capabilities for recognizing toxic targets over extended time periods, even in untrained backgrounds that contain aggressive contaminants at higher concentrations. The approach involves a moving-window, correlation-based methodology to identify chemical events and decouple the foreground conditions from the background.

I. INTRODUCTION

Temperature is a critical factor in all of chemistry, including the chemical interactions that occur between sensing materials employed in microsensors and the molecules present within an environment being monitored [1-4]. To benefit from the valuable analytical information contained within the temperature-dependent oxide/ambient interactions, we have designed and fabricated, through top-side micromachining, arrays of individually addressable microhotplates [5-7]. The array microelements, which serve as microsubstrates for multiple chemiresistive sensing oxides, operate with low power consumption and thermal time constants (≈ ms) that are ideal for implementing a variety of temperature programs. Cycling through temperatures rapidly and repetitively generates temporally varying signals that are associated with changing adsorbate populations. Related work [8-10], and our earlier work employing temperature-dependent data streams [11-13], has focused on detecting and recognizing target (volatile organic) chemicals in the presence of known, or pre-trained, backgrounds. Here, we present methodology developed to deal with target species detection in untrained background conditions. The target analytes studied were the toxic industrial chemicals (TICs) NH₃, HCN and Cl₂ at immediate-danger-to-life-and-health (IDLH) concentrations. These molecules were presented to the sensor in 20 °C air with 30% relative humidity (RH) during the training phase. The testing phase sought to detect these chemicals while relatively high concentrations of interference molecules, such as diesel fuel, Windex [14] cleaner and Clorox [14] bleach, were included in the air-based background.

II. EXPERIMENTAL AND ANALYSIS METHODS

A. Device Fabrication and Testing Protocols

MEMS microhotplate arrays have been used for fabricating chemical microsensor devices [2] and as research tools for investigating temperature-dependent phenomena [6,15]. Typically, multiple types of sensing films/materials are used to populate the elements in microsensor arrays in order to gain different information from multiple gas adsorbate-film interactions [12,13]. A 16-element array (see Fig. 1a; a single element is shown at a higher magnification in Fig. 1b) was employed during this study. It was populated with semiconducting films of SnO₂, TiO₂, SnO₂/TiO2, and TiO₂/RuOx. Four copies of equivalent manufacture for each oxide were all produced by self-lithographic chemical vapor deposition techniques, which have been described elsewhere [2,12,16]. For the development work on recognition in untrained backgrounds discussed in this paper, we focused on response signals measured under temperature cycling from a single sensing material (SnO₂). The morphology typical for these SnO₂ films is shown in Fig. 1c. Analytically rich data were acquired by repeatedly cycling the SnO₂ devices through a pulsed mode, 32-temperature program with four different base-level temperature sectors (see Fig. 2). These sectors facilitate the development of differing adsorbate populations as a function of time within each program cycle. Conductance
measurements made on the SnO₂ film elements (at both temperature peaks/ramps and bases) capture the changing surface electronic characteristics to provide information about the composition of the environment being probed.

Metered concentrations of three TICs: NH₃ (300 μmol/mol), HCN (50 μmol/mol) and Cl₂ (10 μmol/mol), were delivered to the microdevice array during the training and testing phases of this study. During the training they were introduced individually in a background of air at 20 °C with 30% RH, controlled using zero-grade dry air and a dew-point generator. These target species were delivered to the microsensor array again during the testing phase, which also included periods when interference vapors of diesel fuel, Windex [14] cleaner and Clorox [14] bleach, were introduced individually into the air-based background at concentrations several orders of magnitude higher than the TICs. The total flow in the experiments therefore consists of contributions from an individual target, an interference, humid air and balance dry air. The small-volume test cell in which the sensor array was mounted is positioned downstream of the gas handling manifold, and includes a controlled exhaust for continuously maintaining a 1 standard liter per minute (slm) flow rate.

B. Data Analysis Methods

Conductance measurements on the SnO₂ sensor elements for the complete cycle of 32 ramp temperatures were concatenated to form a multi-dimensional sensor response that was used in all analyses. To detect the onset of a foreground odor and register it to the most recent baseline/background response, we employ the following algorithm:

1. Compute the correlation coefficient across sensor responses measured at the 32 different temperatures (1024 correlations) using ‘w’ consecutive measurements (t,…,t-w+1) where t denotes the current measurement cycle.

2. If the computed standard deviation of the sensor response correlations between the 32 temperatures during the current measurement cycle (t) and the previous cycle (t-1) are less than ‘thresh’ and the standard deviation for the cycle (t-3) is greater than thresh, then record the average response of the sensor array at cycle t-3,…,t-3-n as the most recent baseline.

3. Measure the response of the sensor device with respect to the most recent baseline. Then the corrected current response is $R_{new} = (R - R_0)$, where $R$ is the current measured response, and $R_0$ is the most recently identified baseline response.

The following parameters were used to generate results described in this manuscript: $w=5; \text{thresh} = 0.1; n =2$. Since both stimulus onsets and offsets were detectable using this approach, we enforced a fixed temporal window between two chemical events to limit baseline registration to chemical onsets.

For visualization purposes, linear discriminant analysis was used to reduce the dimensionality of the multivariate sensor responses.
III. RESULTS AND DISCUSSION

Fig. 3 illustrates the conductance measurements (plotted as log conductance) obtained from one of the SnO$_2$ microsensor films (shown in Fig. 1b, 1c) at three different ramp temperatures. Note that introduction of reducing gases, NH$_3$ and HCN, both produce conductance increases, while the oxidizing species Cl$_2$ produces a decrease in SnO$_2$ conductance, for each of the six exposures included in the sampling set. It can be observed that although data taken over the six-day period reveal repeatable relative conductance changes to the three target analytes, a significant shift occurs in the response baseline. Hence, for successful detection/recognition of the targets, it is necessary to filter the baseline response from the analyte-induced conductance changes.

In order to track critical signal changes and (then) remove contributions to baseline shifts arising from age-induced changes or background compositional changes in an on-line manner, we examined a moving-window event detection approach based on correlation coefficients. Fig. 4 shows an example of correlation coefficients (along vertical axis) against experimental time (horizontal axis) which ranges over three exposures of the target analyte (HCN); high-correlation bands were noticed at both the analyte onsets and offsets. Similar onset/offset banding was also observed for both NH$_3$ and Cl$_2$. The mean correlation was high during "chemical events" such as target onset/offset, and the standard deviation remained low during the TIC exposure period. Thus a correlation mean increase, or a corresponding decrease in the standard deviation, can be used to register the time mark on the baseline response. The contribution of the problematic baseline/background in the signal can be effectively removed by re-computing sensor responses with respect to that most recently tagged baseline. While this method readily registers events that feature sharp chemical onsets/offsets (rapid concentration introduction and withdrawal), we also validated the robustness of this approach on more gradual analyte introductions [17].

A demonstration of the baseline-correction approach on raw conductance data is shown in Fig. 5. The red circles indicate the baseline just before the identified chemical events (i.e. - HCN introductions). By re-computing the sensor response with respect to the most recently registered baseline, the offset contribution is effectively removed (Fig. 5b).

Fig. 6 shows the multivariate sensor responses over a period of days after LDA dimensionality reduction for all targets. Only the training data for the three analytes were used to determine the projection vectors. Clearly the three analytes form well-defined, distinguishable clusters indicating that sufficient analytical information is available from the temperature-programmed sensor responses to both identify and distinguish the three targets in a 30% RH background.

Next, we examined the ability of the approach to remove the contribution of an untrained background/interference. For this purpose, we introduced the targets in backgrounds that included vapors of either Windex [14], diesel or Clorox [14]. Here again, we adopted the same baseline correction approach to register and remove the sensor response to the background chemical. Fig. 7 shows an example of the response of the sensor during training (30% RH background) and testing phases (here for the Windex-containing [14] background, with 30% RH) after dimensionality reduction. It is worth noting that the projection vectors were obtained using the response of the targets in 30% RH background (training samples) alone. Clearly, the three targets form well-defined, separable clusters.

Figure 3. Selected illustrative responses of the SnO$_2$ sensor (plotted as isotherms) at three different temperatures shown over a six-day data-collection period.

Figure 4. Event detection with temperature programmed sensors. Correlation coefficient between the 32 ramp-temperature measurements (1024 combinations; vertical axis) are shown over time (along the horizontal axis). The six high correlation bands correspond to alternating HCN onsets and offsets.
after the background contribution is removed from the sensor response.

IV. CONCLUSION

We have demonstrated an event-detection and baseline-correction methodology for separating foreground signals representative of a target species from background signals that can be strongly influenced by interfering compounds. In case studies involving temperature-modulated, chemiresistive microsensors, the approach succeeded in identifying IDLH levels of NH$_3$, HCN and Cl$_2$ even in air-based backgrounds that contained high concentrations of fuel and cleaning compounds that had not been present in prior training trials. During the course of development, we also showed that the baseline corrections can be effective in dealing with aging effects of the sensors. Ongoing work seeks to extend the capabilities of our methods to dynamic changes in backgrounds with unknown interferences, and also demonstrate approaches for reducing baseline differences in the electrical characteristics of microdevices fabricated under equivalent manufacturing procedures.

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REFERENCES


[14] Mention of these and any other commercial products is strictly for provision of proper experimental definition, and does not constitute an endorsement by the National Institute of Standards and Technology.

