Selective Gas Detection by a Single MOX-Sensor †

Aleksei Shaposhnik 1*, Pavel Moskalev 1, Alexey Zviagin 1, Kristina Chegereva 1, Stanislav Ryabtsev 2, Alexey Vasiliev 3 and Polina Shaposhnik 4

1 Voronezh State Agrarian University, Voronezh, Russia; moskalefff@gmail.com (P.M.); a.a.zviagin@rambler.ru (A.Z.); kristina.chegereva@yandex.ru (K.C.)
2 Voronezh State University, Voronezh, Russia; ryabtsev@phys.vsu.ru
3 NRC Kurchatov Institute, Moscow, Russia; A-A-Vasiliev@yandex ru
4 Moscow State University, Moscow, Russia; polinashaposhnik@gmail.com
* Correspondence: a.v.shaposhnik@gmail.com; Tel.: +7-920-225-1024
Published: 25 August 2017

Abstract: Selective analysis of CO gas, hydrogen and hydrogen sulfide was carried out using single metal oxide sensor. Non-stationary sensor heating mode allowed discovering individual peculiarities of each gas in question. Resistance vs. time curves were converted into multidimensional data arrays and processed by chemometrics methods; as a result, qualitative and quantitative analysis of single-component gas systems was successfully carried out.

Keywords: MOX-sensor; selectivity; nonstationarity; data arrays; chemometrics

1. Introduction

Utilization of inexpensive and compact gas analyzers based on chemical sensors is usually confined to non-selective analysis of the systems with known composition. There is a clear need to develop relatively cheap and simple gas analyzers performing not only qualitative, but also quantitative analysis. It was previously supposed that this role could be played by multi-sensor systems, so called e-noses. However, reliability of these devices is low because a breakdown of one of the sensors leads to the failure of the whole device. Therefore, it is important to develop a device, which allows carrying out selective gas analysis using only one or two low-selective sensors. This task can be solved by means of non-stationary modes of device operation.

2. Materials and Methods

Sensitive material was prepared by sol-gel method. Surplus of hydrogen peroxide was added to the solution of tin acetate in the glacial acetic acid. Sol of α-tin acid was formed when small amount of ammonia add. Sol was precipitate by centrifuge. Tin dioxide nanopowder was formed as a result of tin acid calcination. Average size of SnO2 granules was 4–6 nm [1].

Glycol was added to nanopowder. Paste was placed onto special dielectric substrate with platinum electrodes. After calcination sensor was formed. In the present work we applied SnO2-based sensor with palladium and platinum additives (3% and 0.5% respectively).

Process analysis was organized using cycles with 15 s period. At the beginning of each period the sensor was heated up to 450 °C during 2 s and cooled to 100 °C during 13 s.
3. Results

3.1. Responses of Sensor

Time-resistance dependences during one period are shown in Figure 1 (for different hydrogen concentrations), Figure 2 (different concentrations of CO), Figure 3 (different concentrations of H₂S). One can mention that the shape of the curves can characterize both the nature of the gas and its concentration.

**Figure 1.** Time-resistance dependence for different hydrogen concentrations during one measurement period.

**Figure 2.** Time-resistance dependence for different carbon monoxide concentrations during one measurement period.
Figure 3. Time-resistance dependence for different hydrogen sulphide concentrations during one measurement period.

3.2. Data Processing

Multi-dimensional data arrays obtained as a result of the experiments, after an appropriate mathematical processing, can be used for selective analysis of other gas systems, which differ from the initial ones by the composition. The values of the sensor resistance $R(j)$ were read with a 50 Hz frequency. Each of the curves presented in Figures 1–3 was obtained by averaging the results of 45 of sensor heating and cooling cycles. Thus, each values of $R(j)$, where $j = 1, 2, ..., 750$, can be considered as a point in 750-dimensional space. Figure 4 shows that after the transformation of the presented data $R(j)$ by the principal component method (PCA), the dimension of the initial problem was reduced to two, with a loss of approximately 2% of the dispersion fraction.

Figure 4. Principal component analysis of carbon monoxide, hydrogen and hydrogen sulfide.
The Figure 4 shows as well that the MOX-sensor demonstrates good selectivity and reasonable monotone behavior at analyte concentrations higher that 5 ppm. This suggests that the approach under discussion is promising for qualitative and quantitative analysis of other gas systems.

All results were obtained using the chemometrics package [2], and software environment for statistical computing R [3]. All specified software is distributed under the terms of GPL licenses, versions 2 and 3.

4. Discussion

There are different reasons for a change of electrical resistance of sensor, when its temperature is changed during gas analysis procedure. First, resistance of semiconductor material depends on temperature, even if this semiconductor is placed into vacuum or into inert gas. Second, chemisorptions of oxygen and other components of air under analysis is also temperature dependent. Third, temperature affects the kinetics of chemical interaction of analytes (reducing gases) and chemisorbed oxygen. Fourth, temperature determines parameters of the desorption of products of chemical interaction of reducing gases with chemisorbed oxygen. The first factor that is the sensing material resistance itself as a function of temperature is not influenced by the character of analyte, however other three factors are strongly affected by the composition of the gas under analysis.

Figures 1–3 demonstrate that the shape of the curves presenting electrical resistance of the sensor as a function of temperature at temperature variation is considerably different for different analytes; this enables qualitative and quantitative analysis of gas using single semiconductor gas sensor. Multi-dimensional data treated using chemometrics approach permitted us to determine selectively the concentrations of carbon monoxide, hydrogen, and hydrogen sulfide.

5. Conclusions

It is shown that metal oxide semiconductor gas sensor characterized by restricted intrinsic selectivity can be used for selective analysis of gas. This approach makes possible the fabrication of simple and not very expensive gas analyzers with enhanced ability to carry out not only qualitative, but also quantitative gas analysis using temperature modulation and chemometrics approach.

Acknowledgments: Authors are thankful to the Russian Scientific Foundation (RSF) for financial support of this activity, project no. 14-13-01470.

Conflicts of Interest: The authors declare no conflict of interest.

References


© 2017 by the authors. Licensee MDPI, Basel, Switzerland. This article is an open access article distributed under the terms and conditions of the Creative Commons Attribution (CC BY) license (http://creativecommons.org/licenses/by/4.0/).