Two-stage Synthesis of Palladium (II) Oxide Nanocrystalline Powders for Gas Sensor Application

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Abstract

Two-stage procedure of palladium (II) oxide nano particle powder preparation based on calcination in dry oxygen of previously synthesized palladium (II) nitrate has been developed. The results of XRD analysis testified the homogeneity of PdO samples prepared by calcination at T=400 and 600 ºC. By electron probe microanalysis it has been established that synthesized PdO powders were characterized with excess of oxygen atoms relative stoichiometric ratio of the elements. It has been found that calcination temperature rise from T=400 ºC to T=600 ºC has led to the increase of oxygen atom concentration and to enlargement of nano dimensional crystallites of palladium (II) oxide powders. Results of coherent-scattering region value determination have shown that synthesized palladium (II) oxide powders can be applied at fabrication of gas sensors within the framework of thick film technology.

Introduction

The development of industry, transport, and adoption of high-tech innovations defy new environmental challenges, which were unknown before. At a turn of XX and XXI centuries most of the atmospheric ozone (90%) was in the stratosphere with a maximum concentration between 17 and 25km [1]. However, this situation started to change during last decades. In the atmospheric air the concentration of toxic oxidizing gases with anthropogenic origin has been steadily increased over the last 20 years. Now low-level ozone and nitrogen oxides are the products of industrial and urban pollution, which has been significantly affected by motor transport and jet aircrafts. It has been established that high concentration of this powerful oxidizing gases in the ambient atmosphere is hazardous to human health [2-5]. Thus, the monitoring of ozone and nitrogen dioxide concentration in the atmosphere is a critical task for human health protection [6].

At present, several conventional analytical techniques, such as UV absorbance, optochemical, optical, chemiluminescence, fluorescence, and electrochemical methods, are available for the accurate analysis of ozone concentration in the air [7-10]. However, these methods can be realized as laboratory apparatuses only. For this reason, various types of binary, ternary and quaternary metal-oxide semiconductors have widely been studied for fabrication of portable gas sensor device for individual use [11-15].

Recently it has been shown that palladium (II) oxide thin and ultra-thin films have encouraging prospects for application in oxidizing gas detection [16-18]. At the very initial stage of PdO thin film study the experimental data demonstrated the good sensitivity, operation speed, reproducibility, and stability in time of test results at detection of ozone and nitrogen dioxide threshold limit value (TLV) concentration [17,18]. It is known [19] the values of gas sensor functional parameters depend upon not only the size of nanoparticles but upon the shape of them also. Therefore, it is very interesting to study the gas sensor properties of palladium (II) oxide nanostructures with other morphology, for example, nanocrystalline powders or ceramics.

For this purpose, we have begun to study of the functional parameters of gas sensors based on palladium (II) oxide nanocrystaline powders and results of their chemical composition, phase structure, and morphology investigation.

Experimentation

Synthesis of palladium (II) nitrate powder

According to [20], the main pathways to obtain palladium (II) oxide powders are reactions of thermal decomposition of palladium (II) hydroxide or palladium (II) nitrate (Figure 1). Prima facie, the procedure of palladium (II) oxide preparation seems to be simple and express, if commercial palladium (II) chloride, which is produced in industrial scale, would be used as the initial reagent.
This technique (designated as Technique 1 in Figure 1) can be realized by two sequences of reactions:

**First sequence of reactions:** The precipitation of palladium (II) hydroxide by the reaction of palladium (II) chloride aqua solution with alkali:

\[ \text{PdCl}_2^{aq} + 2\text{NaOH}^{aq} \rightarrow \text{Pd(OH)}_2^{aq} + 2\text{NaCl}^{aq} \]  
(1)

and by further thermal decomposition of the Pd(OH)$_2$ sediment:

\[ \text{Pd(OH)}_2^{aq} \rightarrow \text{PdO}^{aq} + \text{H}_2\text{O}^{gas} \]  
(2)

However, it is known [21,22] that the presence of even a trace amount of chloride anions in metal-oxide semiconductors leads to a sharp decrease in the sensitivity of gas sensors based on these materials. The preparation of Pd(OH)$_2$ according to equation (1) does not guarantee a complete absence of chloride anions in sediments. As a result of thermal decomposition of Pd(OH)$_2$, containing Cl$^-$ anions impurities the powders of PdO can be contaminated with chloride anions too. Palladium (II) oxide with Cl$^-$ anions is not applicable for gas sensitive property investigation. To prevent the contamination of PdO with Cl$^-$ anions the second sequence of reactions can be used.

**Second sequence of reactions:** In this case initial PdCl$_2$ aqua solution reacts with silver nitrate AgNO$_3$ to produce palladium (II) nitrate (Figure 1).

\[ \text{PdCl}_2^{aq} + 2\text{AgNO}_3^{aq} \rightarrow 2\text{AgCl}^{aq} + \text{Pd(NO}_3)_2^{aq} \]  
(3)

Palladium (II) oxide can be obtained by thermal decomposition of Pd(NO$_3$)$_2$:

\[ 2\text{Pd(NO}_3)_2^{aq} \rightarrow 2\text{PdO}^{aq} + 4\text{NO}_2^{gas} + \text{O}_2^{gas} \]  
(4)This technique can provide the formation of pure palladium (II) oxide powders but using of AgNO$_3$ makes it rather expensive.

It is known the method for preparation of palladium (II) oxide powders based on the thermal decomposition of soluble in water palladium (II) salt aerosols [21]. As carrier gas used for aerosol transport to reactor the nitrogen and ambient air have been chosen. The major task of this method was to control over PdO and, finally, Pd particle size and morphology [23]. It is necessary to note, that there are some mistakes in description of PdO particles preparation procedure [23]. For instance, in this technique palladium (II) sulfate PdSO$_4$ and palladium (II) phosphate Pd$_3$(PO$_4$)$_2$, which are insoluble in water at ambient temperature, together with Pd(NO$_3$)$_2$ have been listed as soluble in water palladium (II) salts [23]. Because PdSO$_4$ and Pd$_3$(PO$_4$)$_2$ are insoluble in water, these salts cannot be used for preparation of water aerosols.

The analysis of possible pathways for preparation of palladium (II) oxide powders lets us to suggest (Figure 1) that Pd(NO$_3$)$_2$ is the best precursor for it. In this work two-stage technique of palladium (II) oxide powders synthesis has been used (designated as Technique II in Figure 1). First stage of it is reaction of palladium with concentrated nitric acid:

\[ \text{Pd}^{aq} + 4\text{HNO}_3^{aq} \rightarrow \text{Pd(NO}_3)_2^{aq} + 2\text{NO}_2^{aq} + 2\text{H}_2\text{O} \]  
(5)

and second stage is thermal decomposition of solid Pd(NO$_3$)$_2$ according to Eq. (4).

At this technique the initial substances are palladium with 99.999 per cent purity and concentrated nitric acid prepared from commercial reagent (purity 99.98 per cent) by distillation. The distillation of concentrated HNO$_3$ has been performed in order to eliminate Cl$^-$ anions, which are presented as hydrochloric acid impurity in commercial reagent. The purity of distilled nitric acid has been checked by reaction with AgNO$_3$. 

**Figure 1:** The possible pathways of palladium (II) oxide synthesis.
The synthesis of palladium (II) nitrate was carried out at T = 75-80 °C in porcelain reactor during 100 hours (with interruptions). At temperature T = 40-45 °C the dark brown solution has been evaporated until the formation of Pd(NO$_3$)$_2$ nano dimensional crystals. As in aqueous solution palladium (II) salts are subjected to hydrolysis, an evaporation of palladium (II) nitrate solution has been carried out at pH = 4.0-4.5 values.

**Synthesis of palladium (II) oxide powder**

Nanocrystalline PdO powders have been prepared by Pd(NO$_3$)$_2$ calcination in dry oxygen atmosphere at temperature T$_{\text{calc}}$ = 400 °C and 600 °C. The duration of Pd(NO$_3$)$_2$ calcination procedure was 240 minutes.

**Analysis of palladium (II) nitrate and palladium (II) oxide**

In 2Q region of 20-120 degrees X-ray diffraction (XRD) patterns of Pd(NO$_3$)$_2$ and PdO powders were obtained with filtered CoK$_\alpha$-radiation ($\lambda$=0.179026nm) on a DRON-8 general purpose diffractometer (Bourevestnik, Russian Federation) with 0.02 degree step-by-step movement and sample rotation. XRD patterns were interpreted using the JCPDS database [24]. Using Scherrer equation the values of coherent-scattering region (CSR) have been calculated for palladium (II) oxide powders only because XRD data for palladium (II) nitrate were obtained in 1939 [25] and many reflexes on Pd(NO$_3$)$_2$ are not indexed until now. For calculation of PdO powders CSR average value the profiles of five more intensive reflexes with 0.01degree step-by-step movement have been recorded.

The quantitative elemental composition of powders which were prepared by palladium (II) nitrate calcination in dry oxygen, has been determined by electron probe microanalysis (EPMA) on JEOL JSM 6510 equipped with Oxford Instruments INCA-sight energy dispersion spectrometer (EDS). The EDS analysis has been performed in five different regions of each sample.

**Results and Discussion**

In the present work XRD analysis was used to identify the substance formed by the interaction of metallic palladium with nitric acid. After evaporation of aqua solution, the dark brown powder was obtained. XRD patterns of this powder (Table 1) have been established that interplanar distance $d_{hkl}$ values of this substance are in total compliance with JCPDS database ones for palladium (II) nitrate Pd(NO$_3$)$_2$ [24,25]. Thus, we are convinced that the product of palladium and nitric acid interaction was palladium (II) nitrate.

**Table 1: XRD experimental and JCPDS data [23,24] on palladium(II) nitrate crystal structure.**

| Reflex Number | Miller Index | JCPDS data [23,24] | Experiment |
|---------------|--------------|--------------------|------------|
|               |              | Interplanar Distance $d_{hkl}$ nm | Intensity $I$, arb, Units | Interplanar Distance $d_{hkl}$ nm | Intensity $I$, arb, Units |
| 1             | ---          | 0.5800             | 48         | 0.58841          | 46.2          |
| 2             | ---          | 0.5300             | 24         | 0.53125          | 100          |
| 3             | ---          | 0.4850             | 5          | 0.48322          | 8.38         |
| 4             | ---          | 0.4220             | 100        | 0.42357          | 21.25        |
| 5             | (200)        | 0.3930             | 40         | 0.39419          | 44.83        |
| 6             | (210)        | 0.3580             | 11         | ---              | ---          |
| 7             | ---          | 0.3480             | 11         | 0.34811          | 10.33        |
| 8             | ---          | 0.2950             | 48         | 0.29603          | 22.81        |
| 9             | (220)        | 0.2800             | 64         | 0.28063          | 39.96        |
| 10            | (221)        | 0.2650             | 24         | 0.26521          | 83.24        |
| 11            | ---          | 0.2560             | 16         | 0.25649          | 17.35        |
| 12            | (311)        | 0.2400             | 16         | 0.24182          | 29.82        |
| 13            | (222)        | 0.2300             | 16         | 0.23195          | 8.77         |
| 14            | ---          | 0.2250             | 2          | ---              | ---          |
| 15            | (321)        | 0.2100             | 40         | 0.21062          | 9.75         |
| 16            | ---          | 0.2060             | 16         | 0.20773          | 13.84        |
| 17            | (400)        | 0.1960             | 8          | ---              | ---          |
| 18            | (410)        | 0.1900             | 13         | 0.19256          | 10.72        |
| 19            | (331)        | 0.1820             | 24         | 0.18178          | 9.75         |
| 20            | ---          | 0.1780             | 13         | ---              | ---          |
By calcinations of Pd(NO$_3$)$_2$ in dry oxygen the black powders with dark green blink were prepared. EPMA results of chemical composition of these products are presented in Figure 2 and Table 2. As it can be seen in Figure 2, X-ray spectra within energy interval $E=0$-$10$eV are characterized with palladium and oxygen lines only. The results of quantitative chemical analysis (Table 2) have shown that the molar ratio of the elements meets to the nominal formula PdO-$\text{palladium (II) oxide}$. It is necessary to note that chemical composition of this oxide (Table 2) does not correspond the stoichiometric ratio 1:1 of the elements and is characterized with little excess of oxygen atoms: PdO$_{1+\delta}$. The data in Table 2 have illustrated that deviation from stoichiometric ratio $\delta=0.149\pm0.004$ and $\delta=0.223\pm0.005$ for samples prepared by calcinations at $T_{\text{calc}}=400^\circ\text{C}$ and $T_{\text{calc}}=600^\circ\text{C}$, respectively.

**Table 2:** EPMA results of chemical composition study of the products prepared by Pd(NO$_3$)$_2$ calcination at $T_{\text{calc}}=400^\circ\text{C}$ and $600^\circ\text{C}$

| Temperature of Pd(NO$_3$)$_2$ Calcination $T_{\text{calc}}=400^\circ\text{C}$ | Element | X-ray Line | Mass fraction $\omega$,\% | Mole fraction $x$,\% |
|---|---|---|---|---|
| | Palaldium | L-line | 14.74 | 53.49 |
| | Oxygen | K-line | 85.26 | 46.51 |
| | Total | | 100\% | 100\% |
| Composition | | $^{*}$PdO$_{1.149\pm0.004}$/Pd$_{0.870\pm0.000}$O |
| Temperature of Pd(NO$_3$)$_2$ Calcination $T_{\text{calc}}=600^\circ\text{C}$ | Element | X-ray Line | Mass fraction $\omega$,\% | Mole fraction $x$,\% |
| | Palaldium | L-line | 15.53 | 55.01 |
| | Oxygen | K-line | 84.47 | 44.99 |
| | Total | | 100\% | 100\% |
| Composition | | $^{*}$PdO$_{1.123\pm0.005}$/Pd$_{0.818\pm0.000}$O |

*The statistical error for five EPMA measurements for each sample is presented

![Figure 2](image1.png)

**Figure 2:** EPMA ED spectra and analysis region SEM micrographs of palladium (II) oxide powders prepared by calcination of palladium (II) nitrate at different temperature $T_{\text{calc}}$: 2a: $T_{\text{calc}}=400^\circ\text{C}$, 2b: $T_{\text{calc}}=600^\circ\text{C}$. 
It is known [26,27] that in palladium-oxygen system can be formed metastable palladium (IV) oxide - $\text{PdO}_2$. In order to identify the phase nature of prepared palladium oxide XRD analysis has been performed. The results of XRD analysis depicted in Figure 3 have testified that all samples were homogeneous and consisted of single-phase palladium (II) oxide (tetragonal structure, space group $P4_2/mmc$) [28]. The values of interplanar spacing $d_{hkl}$ are in a good agreement with ones of JCPDS database [25,28].

Using the special software based on solution algorithm of quadric equation’s system with two unobvious parameter the values of $a$ and $c$ lattice parameters of PdO tetragonal structure have been calculated. Nelson-Riley approximation for $2\theta=180$ degrees diffraction angle has been performed for the refinement and more precise determination of $a$ and $c$ lattice parameter values.

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The comparison of calculated $a$ and $c$ parameter values with JCPDS database standard is presented in Table 3. Analyzing the data in Table 3 it is possible to see that values of $a$ and $c$ lattice parameters of synthesized PdO powder samples are very close to JCPDS database ones [24,28].

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The average values of coherent-scattering region (CSR) $\tau$ of PdO powders prepared by calcinations at $T_{\text{calc}}=400$ °C and $T_{\text{calc}}=600$ °C calculated with the help of Scherrer equation are shown in Table 3. As a first approximation at $\tau$ calculation the dimensionless factor $K$ was adopted equal 0.9 that is permissible for crystalline, the shape of which is close to spherical. Probably, sufficiently high values of statistical error $\Delta\tau$ are caused by two reasons. Firstly, PdO nano crystallites have various dimensions for different crystallographic directions. Secondly, the real shapes of PdO particles may differ from spherical. Nevertheless, the data obtained by EPMA and XRD experiment concerning the composition, phase structure, and crystalline sizes of palladium (II) oxide powders allow us to resume that synthesis procedure, which is set forth in this work, can provide the manufacturing of materials applicable for gas sensors fabrication within the framework of thick-film or ceramic technologies.

Comparing of XRD patterns (Figure 3) with the values of $a$ and $c$ lattice parameters and CSR calculation results (Table 3) it is possible to see the obvious tendencies. With the rise of calcination temperature $T_{\text{calc}}$ the microstructure of palladium (II) oxide samples is exposed to changes in three aspects. At first, the quantity of XRD reflexes and their intensities increase. Secondly, the values of $a$ and $c$ lattice parameters of palladium (II) oxide tetragonal structure increase too. Thirdly, the results of CSR calculation have confirmed the enlargement of PdO nano dimensional particles. All these facts give evidence of the advancement of palladium (II) oxide structural perfection degree with the rise of calcination temperature.
It is necessary to discuss the established tendencies of PdO powder microstructure changes together with the results of chemical composition study. It is well-known that EPMA analysis of oxygen atom concentration using ED spectrometers does not provide the absolute accuracy of experimental results. Nevertheless, from the qualitative point of view the EPMA experimental results (Table 2) are in agreement with the hypothesis, which was expressed in previous publications [29,30], that palladium (II) oxide thin films and single crystals are characterized by the deficiency of palladium atoms relative stoichiometric ratio 1:1. The chemical composition formulae of palladium (II) oxide samples (Table 2) can be recalculated as: $\text{PdO}_{1.149\pm0.004}$ or $\text{Pd}_{1.170\pm0.004}^\text{II} O$ and $\text{Pd}\,\text{O}_{1.223\pm0.005}$ or $\text{Pd}_{1.391\pm0.006}^\text{II} O$. Besides, it is interesting to consider the tendency of chemical composition changes with calcination temperature $T_{\text{calc}}$ rise. Evidently, in PdO powder sample the deficiency of palladium atoms or, in other words, the excess of oxygen atoms increases with calcination temperature $T_{\text{calc}}$ (Table 2).

It is necessary to emphasis that until now the nature of the point defects responsible for the nonstoichiometry of palladium (II) oxide is not established finally [30]. Palladium vacancies or oxygen interstitial atoms can be the reason of palladium (II) nonstoichiometry. The solution of this problem demands the additional investigation.

**Conclusion**

Two-stage procedure of palladium (II) oxide nano particle powders preparation based on calcination in dry oxygen of previously synthesized palladium (II) nitrate has been developed. The results of XRD analysis testified the homogeneity of PdO samples prepared by calcination at $T=400$ and $600$ °C. By electron probe microanalysis it has been established that prepared PdO powders were characterized with excess of oxygen atoms relative stoichiometric ratio of the elements. Until now the nature of the point defects responsible for the nonstoichiometry of palladium (II) oxide is not established finally. Palladium vacancies or oxygen interstitial atoms can be the reason of palladium (II) nonstoichiometry. The solution of this problem demands the additional investigation.

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