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A Vehicle Exhaust NO_x Electrochemical Sensor Based on Au-Yttria Stabilized Zirconia Nanocomposite

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Mixed potential sensor which is built on Yttria Star red Zirconia (YSZ) with a plain planar structure Au/YSZ/Pt has actual benefit among different virus of *Sx* sensors. To further sensing abilities, au The fact that adding YSZ into the Au electrode composites electrodes were take earch of reduced in air the polarization en demonstrated by electrochemical impedance ce has cesist cz com spectroscopy. The proposed osite electrodes in a YSZ-based sensor have more oroù old electroc positive effect than pure based on the excellent performance. The proposed YSZ-based sensor could linearly centration between 50 to 400 ppm with a low detection limit O_2 in the c tec of 20 ppm.

Keywords: NOx periodemical sensor; Gold, Yttria stabilized zirconia; Nanocomposite

1. INTRO ÚCTIC

Various narmful gases, for example, hydrocarbons (HCs), nitrogen oxides (NOx), carbon monoxide (CO), and hydrogen (H₂) are emitted by the combustion of fossil fuels or the leakage from gas tanks in automobiles and power plants by accident. NOx is especially harmful that it causes adverse effect to respiratory system of human body and air pollution in urban area. Hence, to obtain appropriate control of combustion catalysts [1], there are urgent demands for high performance gas sensors monitoring NOx sensitively and selectively [2].

In the last decade there have been plenty of studies and reports on different kinds of solidelectrolyte-based electrochemical NO*x*-sensing devices since conditions were very harsh for operating

a practical sensor (multi-component gas mixtures, high humidity, high temperature). Among which the construction, overall survey and design of stabilized-zirconia-based NOx sensors associated with metallic- or metal oxide-based sensing electrodes (SEs) is attracted lots attentions. Zirconia-based sensors is supposed to be highly sensitive and selective against NO₂ at operating temperatures above 800 °C since temperature of engine may be even 800–900 °C by chance when accelerating vehicle. It was found that a mixed-potential-type yttria-stabilized-zirconia (YSZ)-based sensor with a NiO-SE responded sensitively to NO₂ at temperatures of 800–850 °C [3, 4] when detecting a metal oxide-SE recently. The NiO-SE is supposed to be spruced at comparatively high temp tures of 1300-1400 °C [5] to be more sensitive and selective against NO₂like this. To achiev the NiOreduction SE's temperature when sintered, several later attempts came into being by ddition d second [6-10]. A component(dopant)to obtain the high performing capacity towards NQ -dopedsamar ceria (SDC) [11] or the β -alumina (2Na₂O-11Al₂O₃) [12] were taken referen o use a ctrode.

To screen for NOx, for instance, the couples Pt-(Auxide) [] $Pt-MnOr_2O_3[14]$, or Pt-Au [15], the two electrodes have to display varying electro c of the formation ic activity. ecaus of a mixed potential on the sensing electrode [14], the signal of a electrode otentiometric sensor demonstrates non-Nernstian behaviors in a complex This is latter associated to the as environment. duplication of reactions of parallel electrode coming up in the triple phase boundaries (TPB) [14, 16], for example, the electrochemical reduction(oxidatio of $NO_2(N\Phi)$ and oxygen electrode reaction. NO₂ is entitled with higher sensitivity than that of Q. Besides, NO_2 and NO shows opposite responses by adopting the Au/Pt couple of elic es associated with YSZ, negative for NO₂[15, 17-20] and positive for NO.

A catalytic filter set on te e electroles [21] can help to avoid cross sensitivity to expel p 01 atalytic larger is able to completely bring the molecules into gases like hydrocarbons and Co The CO₂by oxidation without detected To achieve the thermodynamic equilibrium, the the xidation into NO₂. Hence, the NO/NO₂ ratio only relies on catalytic layer can also vze the N pressure of oxygen. Furnished with a catalytic filter, we are to create a the temperature of xed partia detecting 1 sensitively. Then, concentrations of overall NOx, NO₂ and NO at potentiometric ensor oxygen partial. pera re and constant temperature could be brought by the system.

This paper shows the implementation of NO₂ detection of YSZ-based solid state planar sensors with au-composites the sensing electrodes and Pt a reference electrode. Its aim is to improve the quantity of T2Bs of the sensing electrodes and the porosity, which are made up of Au mixed with ZrO_2 later. Dectrochanical capabilities for the reaction of oxygen electrode were detected by impedance spletroscopy and made a comparison with those of pure Au sensing electrode. At last, sensing abilities for NO₂ at a temperature of 800 °C are also assessed.

2. EXPERIMENTS

2.1. Materials and apparatus

For creation of the sensor, a tape-casted YSZ (8 mol.%-doped Y_2O_3) plate(10 mm × 10 mm, 0.2 mm in thickness) was employed. Screen-printing (200 mesh) from a home-made ink which

contains solvent (ESL 404) 1 g mixed in a three-roll mill (Exakt 80E) for 10 mins, YSZ powder4 g and organic binder 1.7 g (ESL V400A) established the functional and porous layer 5 mm × 5 mm of solid electrolyte (YSZ: $(ZrO_2)_{0.92}(Y_2O_3)_{0.08}$ brought by superconductive). After that, the samples were exsiccated at 120 °C in air for 15 min and spruced at 1100 °C for 2 h with a ramp of 5 °C/min. Adopting a commercial paste of Au (ESL 8880-H) as well as Pt (ESL 5545), gold and platinum electrodes of 4 mm length and 1 mm width were screen-printed on the surface of the YSZ layer. The electrodes was 2 mm near each other. The stored electrodes were desiccated at 120 °C further for 15 min and then sintered with a ramp of 5 °C/min for 2 h in air at 800 °C. The othelectrodes was labelled the sensing (Au-SE) electrode and the platinum electrodes was known as the reference (RE) electrode.

2.2. Sensor fabrication

The Au commercial paste and α -terpineol organic kinds, were mixed together, the result of which constituted an Au-SE, about 4 μ m thick on average by screect printed onto the front side of the YSZ. Then the Au-printed YSZ plates were spruced in airat 500 °C, 800 °C and 1100 °C for 2 h.

After calcination in airat 850 °C, specific area of YSZ powder, which was featured by adopting BET measurements with nitrogen (ASAP 2020) and laser granulometry (MASTERSIZER 2000 MALVERN), was 15 m²/g but some area of VO_2 was $3 \times v^2/g$. Mean particle size of agglomerates for ZrO₂andYSZ respectively was 0.288 µm and 16 µm.

In a quartz tube, the fabricated planar errors, tiked to a conventional gas-flow apparatus, e. Features of the sensors for sensing gas were to be tested at were established, set in an electric furn 500 °C. The Pt-RE as well 2 L was uncertered to the sample gas or to wet base gas at the s th Au-S same time(5 vol.% O_2 + N₂ban, e). And each of parent gas (CO, CH₄, C₃H₈, NO and vol.% H₂ NO_2) with the base diluted to ake the sample gas. All of the concentrations of each gas ppm in the sample gas for the selectivity tests. 100 ml/min was the tested were kept stant at 4 tion range of 50–400 ppm, the dependence of the NO_2 sensitivity i the NO_2 concent gas-flow rate. gnal electrometer (R8240, Advantest Corp., Japan), output of the electromotive was assessed. W ing signal, which lies between the Pt-RE and the Au-SE of the sensor. force (as a ser vas tak The of the se In the base gas (*emf*_{base}) and *emf* value in the sample gas (*emf*_{sample}) differ *af* valu ensitive (Δemf) they are to each gas. By adoption of an electrochemical in that analyzer(PCNTAT30, AutoLab[®], The Netherlands), measurements of curves of current-voltage be displayed. Through adoption of a two-electrode configuration, the polarization (polarization) curves were gauged at a constant scan rate of 2 mV/min between -20 as well as +100 mV in potentiodynamic mode.

3. RESULTS AND DISCUSSION

Figure. 1A display that the YSZ layer is extremely porous, probably due to the presence of evaporated binder and solventin the initial paste. However, the surface of this layer evidences

percolation between micrometric YSZ grains ensuring bulk ionic conductivity. Figure 1B shows the XRD pattern of the prepared YSZ.



Assessment of gas-sensing ability at different temp ra tention to by the s is paid current study. Figure 1 demonstrates the cross sensitivities of vario gases for the sensors under the wet condition at 500 °C adopting Au-SEs assembled from the nano-structured precursor before sprued at respectively 1100 °C, 500 °C and 800 °C. Every tested YSZ-based sensor linked to the Au-SEs showed comparatively high NO₂ sensitivity; but to arge extent the results for selectivity against 20 °C-s NO_2 differ. The sensor adopting the s-porous rued Au-SE brought the best NO_2 sensitivity, but the reaction to NO_2 left pices for a usable application. A relatively good reaction to CH₄ accompanied good sensitivity Spite of slightly lowering NO₂ sensitivity 10_2 . investigated, the NO₂ selectivity of the nsor referred to was obtained by the increase of the NiO-SE's sintering temperature to 800 °C fected to the dense and partially melted NiO-The sensor col tivity SE1100 °C-sintered show ch tested gases. d good se



Figure 2. Cross sensitivities to different gases(400 ppm each) for the YSZ-based sensor under wet condition(5 vol.% water vapor) at 500 °C adopting all (A) 500 °C-, (B) 800 °C- and (C) 10yo00 °C-sintered Au-SEs.

Figure 3 displays a distinctive Nyquist plot of the AuY/AuY, AuZ/AuZ and Au/Au in air by 500 °C. For Au/Au sample, two specific semicircles that can be divided and discomposed are exhibited

by impedance spectra. Because of the obstruction of the conduction by the pores [22], in the high frequency (HF) domain, the first semicircle stands for the resistivity of the YSZ solid electrolyte layer (R_{SE}). In the low frequency (LF) scope (from 10^2 to 10^{-3} Hz), at the two symmetrical electrodes, the second contribution confirms with the whole polarization resistance (R_{POL}) connected with the kinetic electrode reactions [23, 24]. The electron transfer resistance (Rct) was measured to be 422 Ω , 457 Ω and 466 Ω at the bare Au/Au, AuZ/AuZ and AuY/AuY, respectively, after the optimization of the equivalent and calculation. A significant increase of Rct was observed when depositing AuY surface.



Figure 3. Certain Nyquist plot under experiment of Autom scalple in air at 500 °C(numbers with arrow indicating \log_{10} of the frequency for a suring).

The apacitance YSZ 4A), resistance of the layer (Figure the properly fits for the SE semi-circle. The YSZ parallel $R_{\rm SE}C_{\rm SE}$ element w n R_{CE} an SET $10^6 \Omega$ at 500 °C and is similar for three sensors (Figure 3B), layer's resistance is around 2 n method, high repeatability. Reports in the past studies [22] showed which emphasizes repara ae i Id be associated with the screen printed YSZ layer's high porosity. lower value that this one, which Around 1.2 ev the a vation energy, which is alike to the three symmetrical sensors, which accords with the value in t porous YSZ membranes infused with 1 wt% Pt. The Arrhenius plot of reports d the Jariza on resis air from 500 to 800 °C is demonstrated by Figure 4B. Generally, resista be YSZ layer is a order of magnitude higher than polarization resistance of the two symmetric electroes. AuZ/AuZ and Au/Au, the two samples, resemble with each other in values of stances and in the apparent activation energy. Hence, adding an insulating material to polarization r enhance the electrode porosity, for example, zirconia appears to change the oxygen electrode kinetic significantly. It is found that the presence of the oxide phase suppresses the cathodic electrode reaction.



Figure 4. (A) Arrhenius plots of the solid electrolyte existance (R_{SE}) (L) and the polarization resistance (R_{POL}) of the three symmetric sensors A Y/AuY, A Z/AuZ and Au/Au.

ion of time (Figure 5) at 500 °C for evolution. In the base gas, OCV was monitored as a fur n the Au-SE and the Pt electrode. These responses go in accordance with potential different es betwe Because of the temperature regulation, initia al varies upruptly. OCV values of the Au sensor increase greatly in the first 2 h, after that it will by step. However, it cannot get to a steadyris state forever. The Pt/Au sensor's bund to be +61 mV after 20 h on stream. The OCV alue is between Pt electrodes and Au en atmos here often results from the disagreement in the oxy O_2 electrode kinetic rate c ygen activity. Behaviors of the two Au composites nneeted are alike, starting from OCV and decaying gradually. The AuZ and AuY sensors' crease of together v xth +3 mV respectively after on stream5 and 12 h. responses get to a



Figure 5. Sensor Pt/YSZ/SE baseline in 1.5% H₂O, 12% O₂ at 450 °C.

Figure 6 shows different reactions as well as recovery times of the three sensors after putting in 100 ppm NO₂ at 500 °C. AuY's response time is approximately 20 s, the shortest one, but the Au sensor spends about 40 s to respond and the sensor of AuZ needs 15 mins to react, the slowest one with

instable signal. To reassume the reaction at the beginning after the NO₂ exposure ends under 500 °C, several minutes are necessary, whatever the electrode is Au–Y composite could strongly cut thin recovery time for AuZ and Au which is 7 min instead of 40 min. A good adsorption of NO₂ onto the electrodes is likely to cause these long recovery time. The recovery time is reasonable on AuY and Au at a temperature of 550 °C with a lower concentration of NO₂ (20 ppm). The linear response concentration of proposed AuY is between 50 to 400 ppm.



Figure 6. Responses (ΔV) of AuY, Au and AuZ sense with 12% O₂ and 1.5% H₂O at 500 °C to 100 ppm of NO₂.

4. CONCLUSIONS

research on the gold composite electrodes' impact on the The study way d to take Au/YSZ/P sensor. The competence of sensor referred to on response capacities of a mix otenti toward NO₂ wa °C. The fact that the polarization resistance is greatly decreased tored at AV-YSZ electrode is showed by electrochemical impedance under the c of msta es demonstrates that due to delocalization of YSZ's ionic conductivity to electrode measurements, wh ectro e volume, the electrochemical features of the oxygen electrode response whole respo se in ized for the composite electrode of Au-YSZ only. Thus, it can be concluded that can be ptir electrode, in a YSZ-based sensor porous Au–YSZ composite electrodes display compared pure g a beneficial e

References

- 1. S. Capone, A. Forleo, L. Francioso, R. Rella, P. Siciliano, J. Spadavecchia, D. Presicce and A. Taurino, *Journal of Optoelectronics and Advanced Materials*, 5 (2003) 1335
- Y. Li, X. Li, Z. Tang, J. Wang, J. Yu and Z. Tang, Sensors & Actuators B Chemical, 223 (2015) 365
- 3. P. Elumalai and N. Miura, Solid State Ionics, 176 (2005) 2517
- 4. P.A. Rasheed, T. Radhakrishnan, P.K. Shihabudeen and N. Sandhyarani, Biosensors &

bioelectronics, 83 (2016) 361

- 5. P. Elumalai, J. Wang, S. Zhuiykov, D. Terada, M. Hasei and N. Miura, *Journal of The Electrochemical Society*, 152 (2005) H95
- V.V. Plashnitsa, T. Ueda and N. Miura, *International journal of applied ceramic technology*, 3 (2006) 127
- P. Elumalai, V.V. Plashnitsa, Y. Fujio and N. Miura, *Journal of The Electrochemical Society*, 156 (2009) J288
- S. Sundaravel, B. Venkatachalapathy, L. Sujatha, N.M. Sudharsan, T.S. Rao and T.M. Sridhar, "Yttria stabilized Zirconia sensors for biomedical detection of hydrogen sulphide", Indo – Australian Conference on Biomaterials, Tissue Engineering, Drug Delivery Systems, Regenerative Medicine, 2015.
- 9. L.Y. Woo, R.S. Glass, E.L. Brosha, R. Mukundan, F.H. Garzon, W.J. Buttne, M.B. Post, Rivkin and R. Burgess, *Ecs Transactions*, 45 (2013) 19
- 10. M.J. Lee, J.H. Jung, K. Zhao, B.H. Kim, Q. Xu, B.G. Ahn, S.H. Kim and S.Y. Kim Journal of the European Ceramic Society, 34 (2014) 1771
- 11. C. Pijolat, G. Tournier and J.-P. Viricelle, Sensors and Actuators 8: Connacal, 14 (2009) 7
- 12. N. Guillet, R. Lalauze, J.-P. Viricelle, C. Pijolat and L. Mortmaro, *Materials Science and Engineering: C*, 21 (2002) 97
- J. Zosel, D. Tuchtenhagen, K. Ahlborn and U. Guth. *Casors and cituators B: Chemical*, 130 (2008) 326
- 14. C. Yin, Y. Guan, Z. Zhu, X. Liang, B. Wang, Q. Lao, H. Zhang, J. Ma, F. Liu and Y. Sun, Sensors and Actuators B: Chemical, 183 (2013) 474
- 15. J. Gao, J.-P. Viricelle, C. Pijolat, P. Breuil, P. Vernot, A. Boreave and A. Giroir-Fendler, *Sensors and Actuators B: Chemical*, 154 (2011)
- 16. N. Miura, S. Zhuiykov, T. Ono, M. Hasei and T. Yamazoe, Sensors and Actuators B: Chemical, 83 (2002) 222
- 17. J. Zhu, C.R. Pérez, T.S. Oh, B. Ku, as and J.A. Vohs, *Journal of Materials Research*, 30 (2014) 357
- 18. M. Fee, S. Ntais, A. Wert and A. Josenova, *Journal of Solid State Electrochemistry*, 18 (2014) 2267
- 19. C.H. Hua and C.C. Cho. Jpn. j. appr. bys, 55 (2016) 080302
- 20. S. Yu, C. Han, P. L. H. D. S. He and Z. Guo, International Journal of Hydrogen Energy, 39 (2014) 562
- 21. J.-C. Yang and P.K. Dutta, Sense's and Actuators B: Chemical, 123 (2007) 929
- 22. M. Kleitz and M. Steil, Journal of the european ceramic society, 17 (1997) 819
- 23. L. Betel, P. Veneux, F. Gillard, C. Roux and E. Siebert, Solid State Ionics, 176 (2005) 793
- 24. T Wang, E.F. Nova and R.E. Soltis, Sensors and Actuators B: Chemical, 77 (2001) 132
- 25. W. 7 Zhang, P. Schmat-Zhang and U. Guth, Solid State Ionics, 169 (2004) 121

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