High-performance varistors simply by hot-dipping zinc oxide thin films in Pr$_6$O$_{11}$: Influence of temperature

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High-performance ZnO-Pr$_6$O$_{11}$ thin-film varistors were fabricated simply by hot-dipping oxygen-deficient zinc oxide thin films in Pr$_6$O$_{11}$ powder. The films had a composition of ZnO$_{0.81}$ and a thickness of about 200 nm, which were deposited by radio frequency magnetron sputtering a sintered zinc oxide ceramic target. Special attention was paid on the temperature dependence of the varistors. In 50 min with hot-dipping temperature increased from 300–700 °C, the nonlinear coefficient ($\alpha$) of the varistors increased, but with higher temperature it decreased again. Correspondingly, the leakage current ($I_L$) decreased first and then increased, owing mainly to the formation and destroying of complete zinc oxide/Pr$_6$O$_{11}$ grain boundaries. The breakdown field ($E_{1mA}$) decreased monotonously from 0.02217 to 0.01623 V/nm with increasing temperature (300–800 °C), due to the decreased number of effective grain boundaries in the varistors. The varistors prepared at 700 °C exhibited the optimum nonlinear properties with the highest $\alpha = 39.29$, lowest $I_L = 0.02736$ mA/cm$^2$, and $E_{1mA} = 0.01757$ V/nm. And after charge-discharge at room temperature for 1000 times, heating at 100 or 250 °C for up to 100 h, or applying at up to 250 °C, the varistors still performed well. Such nanoscaled thin-film varistors will be very promising in electrical/electronic devices working at low voltage.

Nowadays technological development aims for designing and manufacturing various kinds of super-large-scale integrated circuits, which require increasing quantity of miniaturized devices including various varistors working at low voltage. Up to now, most of the commercially applied varistors have been ZnO-Bi$_2$O$_3$ based electronic ceramic devices, which are widely used in various electrical circuits, electronic devices and electrical power systems to counteract against dangerous over-voltage surges, due to their highly nonlinear current-voltage ($I$-$V$) characteristics and outstanding energy handling capabilities$^{1,2}$. For conventional ZnO-Bi$_2$O$_3$ based ceramic varistors, they are prepared by sintering ZnO powder with a small amount of the so-called varistor forming oxide (VFO) Bi$_2$O$_3$, which is essential for imparting the nonlinearity of ZnO ceramics, while other oxides, such as nickel oxide, manganese oxides, rare metals oxides and so on, are also added in small amount to further enhance the non-linearity of the varistors$^{3,4}$. It is well-known that their nonlinear $I$-$V$ characteristics are controlled by the well-recognized double Schottky barrier mechanism, which is triggered by the contacting layer between ZnO grains (the core) and the VFO Bi$_2$O$_3$ thin films (the shell) on them formed during sintering$^5$. Thus, to meet the requirement of miniaturized low-voltage devices, complicated and relatively costly processes such as tape casting, higher temperature sintering and micro machining/manufacturing have to be developed, attempting to promote the growth of ZnO grains and/or reduce the thickness of the electronic components. But owing to the complicated processes and low product consistence during preparing low-voltage ZnO chip varistors by ceramic methods, the alternative films deposition methods aiming at reducing the thickness of the components are more desirable$^6$. To date, several physical and chemical methods have been proposed to prepare ZnO film varistors, such as magnetron sputtering, molecular beam epitaxy, pulsed laser deposition, chemical vapor deposition, spray pyrolysis, sol-gel process, and so on$^{7-10}$. Among them, magnetron sputtering is preferable due to its relatively high deposition rate, low substrate temperature, high film thickness uniformity, good controllability and high repeatability$^{11}$. And many composite ZnO-Bi$_2$O$_3$ and multilayered ZnO/Bi$_2$O$_3$ thin-film varistors were reported$^7,8$. But to prepare such varistors, dual (Zn/ZnO and Bi/Bi$_2$O$_3$) or various composite ZnO-Bi$_2$O$_3$ sputtering targets, and even more complicated precursors are needed. And because

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Pr6O11 powder at the designed conditions, the elemental composition and chemical state of the obtained samples in Pr6O11 at 700 °C for 50 min in air are shown in Fig. 3a–d. It can be seen that Zn and O atoms are distributed helped the grain in the films to grow.

energy. Furthermore, with increasing hot-dipping temperature, the mean grain size of the samples increased revealing that the hot-dipping promoted the reforming of the grains in the films, reaching their lower state of films turned out to be smoother and denser, and the grains in the films became more round (see Fig. 2b–g), with flaky (of obvious edges) and unevenly sized grains. But after the films were hot-dipped, the surface of the characteristic of low-temperature deposition, effective double Schottky barriers at the grain boundaries6. However, such thin-film varistors will still suffer from low high-temperature stability in applications, because at high temperature, the core oxygen-deficient zinc oxide films are easily oxidized and the shell Bi2O3 thin films are highly volatile.

Moreover, just because Bi2O3 is used as the VFO, the conventional ZnO-Bi2O3 based ceramic varistors have a few drawbacks due to the high volatility and reactivity of Bi2O3 during liquid sintering12. The high volatility of Bi2O3 will change the varistors’ characteristics due to the variation of inter-composition ratio of the additives. The high reactivity of Bi2O3 will destroy the multi-layer structure of the chip varistors and deteriorate the surge-absorption capabilities due to the decrease of effective grain boundary number13. To overcome these problems, ZnO ceramic varistors containing different VFOs have been actively studied, in which ZnO-Pr6O11 based varistors are most promising in application, due to their advantages over ZnO-Bi2O3 based varistors in relatively simple two-phase microstructure of ZnO grains and praseodymium oxide intergranular phases. Such structure can reduce the quantity of compositional materials, improve the electrical properties of the ceramics, and keep the composition stable at high-temperature sintering due to the high melting point of Pr6O1114.

Therefore, in this work, we use Pr2O3 to replace Bi2O3 in the hot-dipping of oxygen-deficient zinc oxide thin films. Special attention was paid on the temperature dependence of the composition, microstructure and electrical properties of the samples. Surprisingly, ZnO-Pr2O3 thin-film varistors with a recorded high nonlinear coefficient (39.29) were fabricated. And because of the high-temperature stability of Pr2O3, the obtained ZnO-Pr2O3 thin-film varistors presented a very high performance even in elevated temperature applications.

Elemental composition and chemical state
X-ray photoelectron spectroscopy (XPS) analysis indicated that, only Zn 2p and O 1s peaks could be detected in the applied oxygen deficient zinc oxide thin films, which had a composition of ZnO0.81. After hot-dipping in Pr2O3 powder at the designed conditions, the elemental composition and chemical state of the obtained samples were also examined by XPS. Typical results on the sample hot-dipped in Pr2O3 in air at 700 °C for 50 min are presented in Fig. 1. From the full spectrum as shown in Fig. 1a, Zn 2p, Pr 3d and O 1s peaks could be identified, indicating that some Pr atoms were composite with the zinc oxide film after hot-dipping. Moreover, from Fig. 1b, it can be seen that the Zn 2p narrow spectrum of the hot-dipped samples displays two symmetric peaks. The peaks located at 1022.2 ± 0.1 eV and 1045.2 ± 0.1 eV can be assigned to Zn 2p3/2 and Zn 2p1/2 of Zn2+ oxidation state, respectively15. In addition, as seen from Fig. 1c, the asymmetric Pr 3d5/2 narrow spectrum can be deconvoluted into two peaks. The strong peak located at 977.1 ± 0.1 eV is corresponding to Pr6+ oxidation state, while the weak peak at 971.5 ± 0.1 eV can be assigned to Pr3+ oxidation state16. And from this spectrum, the percentages of Pr4+ and Pr3+ oxidation states were further evaluated as 66.7% and 33.3%, respectively. Thus the mean valence of praseodymium atoms in this sample was calculated as 3.67. In other words, the chemical composition of Pr with O is Pr2O3, which is exactly in agreement with that of the originally applied pure Pr2O3 powder, implying that Pr2O3 presents mainly as a separate phase in the sample. Meanwhile, as shown in Fig. 1d, the narrow spectrum of O 1s of the samples could be fitted into two components. The first component has a binding energy of 529.8 ± 0.1 eV, which can be attributed to the oxygen atoms in ZnO17 and Pr2O318. The second one has a binding energy of 531.4 ± 0.1 eV, which is owing to the adsorbed oxygen (such as O2) on the film surface19. All these results indicate that the as-presented hot-dipped sample is a ZnO-Pr2O3 composite. And considering that the originally applied zinc oxide thin films were oxygen deficient with a composition of ZnO0.81, such results revealed that the zinc oxide thin films were further oxidized during the hot-dipping.

Microstructure and elemental distribution
Figure 2 presents typical field-emission scanning electron microscopy (FE-SEM) images on the surfaces of the prepared thin films, revealing that the hot-dipping imparts a significant impact on the morphology of the samples. As is seen from Fig. 2a, the surface of the as-deposited ZnO0.81 films without hot-dipping is relatively rough with flaky (of obvious edges) and unevenly sized grains. But after the films were hot-dipped, the surface of the films turned out to be smoother and denser, and the grains in the films became more round (see Fig. 2b–g), revealing that the hot-dipping promoted the reforming of the grains in the films, reaching their lower state of energy. Furthermore, with increasing hot-dipping temperature, the mean grain size of the samples increased gradually from 49.37 to 121.68 nm (see the inset of each image and Fig. 2h), indicating that the hot-dipping helped the grain in the films to grow.

The elemental composition and distribution of all the hot-dipped samples were further examined by energy dispersive X-ray spectroscopy (EDX). Typical results on the fresh surface of the sample prepared by hot-dipping in Pr2O3 at 700 °C for 50 min in air are shown in Fig. 3a–d. It can be seen that Zn and O atoms are distributed quite uniformly in the samples, but it seems that Pr atoms are scattered mainly at the grain boundary and it is very difficult to estimate if the added Pr2O3 diffuses into the oxygen-deficient zinc oxide grains during the
hot-dipping. This result indicates that the hot-dipping could drive the added Pr$_6$O$_{11}$ selectively spread into the grain boundaries of the zinc oxide films, constructing the typical structure of a ZnO-Pr$_6$O$_{11}$ varistor. In addition, with increasing hot-dipping temperature, the measured content of Pr in the samples increased initially and decreased somewhat later (see Fig. 3e). When the samples were hot-dipped from 300 to 700 °C, the content of Pr increased from 1.73 to 5.11 at.%. But when the temperature was further increased up to 800 °C, the content of Pr was reduced to 4.36 at.%. For this phenomenon, it can be explained as follows. Because the Pr$_6$O$_{11}$ distribution is controlled by thermal diffusion process, so a moderate increase of the hot-dipping temperature is beneficial for its diffusion into the zinc oxide film. When it does not exceed 700 °C, with the increase of hot-dipping temperature, the diffusion speed of Pr$_6$O$_{11}$ increases, resulting in increased content of Pr in the samples. However, when it is higher than 700 °C, with further increasing temperature, the content of Pr in the samples decreases, possibly because Pr$_6$O$_{11}$ separates out from the samples at enhanced temperature. Moreover, typical results on the cross-section of the sample prepared by hot-dipping in Pr$_6$O$_{11}$ at 700 °C for 50 min in air are presented in Fig. 4. It can be seen that the Zn and O atoms are distributed quite homogeneously on the cross-section of the films as they have done on the fresh film surface; and the Pr element also presents a similar non-uniform distribution on the cross-section of the films with that on the film surface. And, Pr has already diffused throughout the whole film without any gradient along the film thickness. The afore-mentioned results indicate that the elemental composition and distribution of the hot-dipped samples are consistent throughout the films.

From the microstructure examination on the film cross-section, it should be noted that all the hot-dipped film samples have a quite homogeneous thickness of about 200 nm, which is almost equal to the original thickness of the as-deposited ZnO$_{0.81}$. This result implies that the hot-dipping would not change the thickness of the samples, which is beneficial to keeping the size of the varistor components.

Phase composition and defect state

The phase composition and defect state of the film samples were examined by X-ray diffraction (XRD). Figure 5a shows the XRD patterns of the samples prepared by hot-dipping in Pr$_6$O$_{11}$ for 50 min in air at different temperatures. To make a comparison, the result on the as-deposited ZnO$_{0.81}$ thin films is also presented. As is seen from this figure, all the samples only display a relatively strong (002) peak and a much weak (103) one, both of which can be indexed to the hexagonal wurtzite structure zinc oxide phase (JCPDS card no. 65–3411). In other word, only ZnO phase can be identified without any other extra phases, no matter whether they are hot-dipped or not (also see Extended Data Fig. 1a). This phenomenon is resulted in probably by the fact that, under the applied hot-dipping conditions, the content of Pr$_6$O$_{11}$ diffusing into the films is low, which cannot be identified in the XRD detection limit. Considering the high stability of Pr$_6$O$_{11}$ (which is stable in air at a temperature of up to
Figure 2. SEM results. FE-SEM surface images of the as-prepared thin films samples hot-dipped in Pr$_6$O$_{11}$ for 50 min in air at different temperatures: (a) without hot-dipping, (b) 300, (c) 400, (d) 500, (e) 600, (f) 700 and (g) 800 °C. The inset in each image displays the grain size distribution in the films. (h) The calculated grain size from the FE-SEM images as a function of the hot-dipping temperature.
800 °C, as is seen from the Extended Data Fig. 2) and high melting point of Pr$_6$O$_{11}$ (2042 °C), the existed praseodymium oxide in the film should be in the form of Pr$_6$O$_{11}$ phase. And this result is in accordance with the EDX data as shown in Fig. 3e.

To evaluate the defect state in the samples, the lattice constant $c$ of the zinc oxide grains in the prepared samples was calculated from the recorded XRD patterns, and the result is presented in Fig. 5b. For comparison, the result on the as-deposited ZnO$_{0.81}$ film is also presented. The lattice constant of the as-deposited ZnO$_{0.81}$ thin film is 5.1668 Å, smaller than that of stoichiometric ZnO (5.2066 Å)\(^{20}\), implying that the applied ZnO$_{0.81}$ thin film is an oxygen-deficient one with lots of oxygen vacancies\(^{11,21}\). However, after hot-dipping at the designed temperatures, the lattice constant for the samples increased, although all of them were still less than 5.2066 Å (more or less). This result implies that the oxygen-deficient ZnO$_{0.81}$ thin films would be oxidized during hot-dipping because oxygen atoms are easy to diffuse into the films, resulting in zinc oxide films with less oxygen vacancies; but they had not been oxidized completely under the present conditions and thus there were still certain amount of oxygen vacancies in the film samples. In particular, with the hot-dipping temperature increased from 300–600 °C, the lattice

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**Figure 3.** EDX mapping results on the fresh surface of a typical sample prepared by hot-dipping ZnO$_{0.81}$ thin film in Pr$_6$O$_{11}$ at 700 °C for 50 min in air. (a) EDX scanning area; and (b–d) EDX spectra of Zn, O and Pr, respectively. (e) The measured content of Pr in all the hot-dipped samples as a function of the hot-dipping temperature.
constant promptly increased, indicating that the films were being oxidized quickly, resulting in samples with zinc oxide grains of less and less oxygen vacancies. When the hot-dipping temperature increased from 600–800 °C, the lattice constant increased more and more slowly, approaching to 5.2066 Å (the value of stoichiometric ZnO), implying that the samples were gradually oxidized completely with a composition of zinc oxide close to the stoichiometric ZnO. And all these results are similar with the lattice constant change of the as-deposited ZnO$_{0.81}$ film annealed at the same temperature as in hot-dipping (see Extended Data Fig. 1b). Moreover, although the radii of both Pr$^{3+}$ (99 Å) and Pr$^{4+}$ (85 Å) ions are bigger than that of Zn$^{2+}$ ion (74 Å)$^{22}$, the lattice constant of the hot-dipped samples is still smaller that of stoichiometric ZnO, implying that the Pr ions would not diffuse into the zinc oxide grains during the hot-dipping. This result indirectly confirms that Pr$_6$O$_{11}$ is mainly distributed at the grain boundary in the process of hot-dipping, which is in agreement with the results recorded by XPS and SEM mapping.

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**Figure 4.** EDX mapping results on the cross-section of a typical sample prepared by hot-dipping ZnO$_{0.81}$ thin film in Pr$_6$O$_{11}$ at 700 °C for 50 min in air. (a) EDX scanning area; and (b–d) EDX spectra of Zn, O and Pr, respectively.

**Figure 5.** XRD results. (a) XRD patterns for the films samples prepared by hot-dipping in Pr$_6$O$_{11}$ for 50 min in air at different temperatures. (b) Lattice constant c calculated from the recorded XRD patterns for the zinc oxide grains as a function of the hot-dipping temperature. For comparison, the results on the as-deposited ZnO$_{0.81}$ film are also presented.
Varistor properties

Figure 6 presents the electric field vs current density (E-J) characteristic curves of the zinc oxide film varistors prepared by hot-dipping ZnO_{0.81} thin films in Pr_{6}O_{11} for 50 min in air at different temperatures. It is well-known that the as-deposited oxygen-deficient ZnO_{0.81} film will display an Ohmic behavior, because there is no double Schottky barrier between their grain boundaries. But after hot-dipping in Pr_{6}O_{11} under the designed conditions, all the samples exhibit excellent nonlinear characteristics, indicating they are promising varistors. This result also implies that through hot-dipping in Pr_{6}O_{11}, an intergranular Pr_{6}O_{11}-rich phase, which separates the semiconducting ZnO grains, has been well formed, because the presence of electrically active grain boundaries can produce the double Schottky barriers, inducing the non-linear characteristics of the varistor samples.

From Fig. 6, the basic electrical parameters of the obtained varistors were calculated, in which nonlinear coefficient (\(\alpha\)) is the most important one. Figure 7 illustrates the corresponding nonlinear coefficient of the varistors as a function of the hot-dipping temperature. As is seen from this figure, with the increase of hot-dipping temperature, the nonlinear coefficient initially increased and then decreased, reaching a maximum of 39.29 (much higher than those of the ZnO-based bulk ceramic varistors) when the varistor was prepared by hot-dipping ZnO_{0.81} thin film in Pr_{6}O_{11} at 700 °C for 50 min in air. With increasing hot-dipping temperature when it was less than 700 °C, the nonlinear coefficient of the prepared varistors increased gradually. This is because in the process of hot-dipping, Pr_{6}O_{11} would gradually permeate into the zinc oxide grain boundaries in the films, forming an insulating layer at the grain boundaries, finally triggering the nonlinear behavior of a varistor. And because the formation of the intergranular insulating Pr_{6}O_{11} layer is determined by thermal diffusion process, as discussed in Section 3.2, a moderate increase of the hot-dipping temperature is beneficial for the diffusion of Pr_{6}O_{11} into the zinc oxide grain boundaries in the films. As is seen in Fig. 3e, the content of Pr in the films continuously increases till the hot-dipping temperature increases up to 700 °C. Therefore, with increasing hot-dipping temperature up to 700 °C, the structure of the intergranular insulating layer will become more and more completed. As a result, the nonlinear coefficient of the samples increased. However, when the hot-dipping temperature was higher than 700 °C, the nonlinear coefficient of the samples would gradually decrease with further increasing temperature. This is because Pr_{6}O_{11} separates out from the intergranular insulating layer of the zinc oxide film varistors at enhanced temperatures with decreasing content of Pr in the films, destroying the
completeness of the intergranular layer, reducing the intergranular insulation resistance, finally leading to the
decrease of the nonlinear coefficient of the samples.

Correspondingly, with the increase in the hot-dipping temperature, the leakage current ($I_L$) initially decreased
and then increased, achieving a minimum of 0.02736 mA/cm² for the varistors prepared by hot-dipping ZnO₀.₈₁
thin film in Pr₆O₁₁ at 700 °C for 50 min in air (see the blue line marked with balls in Fig. 7). With the increase of
hot-dipping temperature, the leakage current of the samples shows a trend against the nonlinear coefficient. The
high nonlinear coefficient for a varistor would lead to low leakage current because of the formation of a relatively
high tunnel current, while the low nonlinear coefficient would result in high leakage current due to the formation
of higher thermal excitation radio current²⁵.

The varistor voltage ($E_{1mA}$) and voltage per grain boundary ($V_g$) of the zinc oxide film varistors as a func-
tion of hot-dipping temperature are illustrated in Fig. 8. The varistor voltage sharply decreased monotonously
from 0.02217 to 0.01623 V/nm with the increase of hot-dipping temperature (black line marked with stars). It
is well-known that the breakdown voltage (varistor voltage) of zinc oxide varistors depends on the size of zinc
oxide grains in the samples and the applied voltage per grain boundary²⁶. But from Fig. 8, it can be seen that,
with increasing hot-dipping temperature, the applied voltage per grain boundary changed in a very narrow range
(1.35–1.72, see the blue line marked with balls). And the varistor voltage of the present zinc oxide film varistors
is in inverse proportion with the size of the zinc oxide grains in the films (see Fig. 2h). Thus, the sudden drop of
varistor voltage with increasing hot-dipping temperature is attributed to the decrease in the number of effective
grain boundaries caused by the increase in the ZnO grain size.

In summary, when the samples were hot-dipped in Pr₆O₁₁ at 700 °C for 50 min in air, the prepared zinc
oxide thin-film varistor presents the maximum nonlinear coefficient of 39.29, the minimum leakage current of
0.02736 mA/cm², and a varistor voltage of 0.01757 V/nm.

Electrical conducting mechanism

Impedance spectroscopy was used to characterize the different microstructure regions of the zinc oxide films
samples prepared by hot-dipping in Pr₆O₁₁ for 50 min in air at different temperatures. Typical complex imped-
ance spectra for the present varistor materials, $Z''$ versus $Z'$, are displayed in Fig. 9a. From this figure, the grain
and grain boundary resistances of the zinc oxide film samples hot-dipped at different temperatures are shown in
Fig. 9b. As is seen from this figure, with the increase in hot-dipping temperature, the grain resistance of the sam-
pies gradually increased, reaching a plateau when the temperature was higher than 600 °C (black line marked with
stars), while the grain boundary resistance initially increased and then decreased, achieving a maximum when the temperature was 700 °C (blue line marked with balls).

As discussed in Sections 3.2 and 3.3, microstructurally, the hot-dipping of oxygen-deficient zinc oxide thin films in Pr$_6$O$_{11}$ will give rise to a structure consisting of semiconducting zinc oxide grains surrounded by very thin insulating intergranular Pr$_6$O$_{11}$ layers. Accordingly, as shown in Fig. 9a, the negative impedance diagram describes the grain resistance and grain boundary resistance. With the increase in the hot-dipping temperature, the oxidation of oxygen-deficient zinc oxide films would become more and more completed, which would result in less oxygen vacancies in the films and thus increase the grain resistance (see Fig. 9b). Because of the existence of oxygen vacancies in the oxygen-deficient zinc oxide films, under the action of applied electric field, the adjacent electrons will fill the vacancies, leaving new spaces in the original positions of electrons. After that, other electrons would transfer to the newly created spaces. As a result of the whole process, a certain amount of charges were transferred (conducting). This process is called vacancy conduction. Since the number of defects (oxygen vacancies) decreased with the increase of hot-dipping temperature, electrons would be difficultly transferred via vacancy, thus reducing the charges transferred. Consequently, the electrical resistivity of the zinc oxide films would increase with increasing hot-dipping temperature. As for the plateau of grain resistance when the temperature was higher than 600 °C, it is because the samples were gradually oxidized completely with a composition of zinc oxide close to the stoichiometric ZnO, which possesses the highest grain resistance in all forms of oxygen-deficient zinc oxides.

With regards to the change of grain boundary resistance with the hot-dipping temperature (see the blue line marked with balls in Fig. 9b), it can be explained as follows. When it was lower than 700 °C, the grain boundary resistance increased gradually with increasing hot-dipping temperature. This is because Pr$_6$O$_{11}$ permeated into the boundary of the zinc oxide grains in the film varistor samples during the hot-dipping, forming an insulating layer at the grain boundary, and with increasing hot-dipping temperature, the insulating layer became more and more completed, thus resulting in increased grain boundary resistance. But if it was higher than 700 °C, with further increased hot-dipping temperature, the grain boundary resistance of the samples decreased gradually. This phenomenon is correlated with the separation of Pr$_6$O$_{11}$ from the grain boundaries in the varistors at enhanced temperatures, which will destroy the completeness of the intergranular insulating Pr$_6$O$_{11}$ layer, thus decreasing the electrical resistance of the grain boundary. When the film was hot-dipped in Pr$_6$O$_{11}$ at 700 °C for 50 min in air, the resultant sample would present a maximum value of grain boundary resistance, the maximum contribution of grain boundary resistance to the total resistance, because in such case an intergranular insulating layer with perfect structure was formed. Because the nonlinearity of a varistor is strongly dependent on the resistivity of the grains and grain boundary, low value of grain resistance and high value of grain boundary resistance will lead to a high nonlinear coefficient$^{27}$. Therefore, the sample prepared by hot-dipping ZnO$_{0.81}$ thin films in Pr$_6$O$_{11}$ at 700 °C for 50 min in air presented the optimum nonlinear performance, although its grain resistance is relatively large.

**Varistor stability**

There are many factors affecting the stability of ZnO varistors. When a varistor is working, it would always bear a certain steady-state voltage. Thus, as a resistor, the number of charging-discharging cycle and working temperature it suffers from often present a significant impact on its nonlinear characteristics. In order to investigate the varistor stability for the present ZnO-Pr$_6$O$_{11}$ thin film samples, with the samples prepared under the optimum condition (by hot-dipping ZnO$_{0.81}$ thin films in Pr$_6$O$_{11}$ in air at 700 °C for 50 min), their E-J curves were recorded after the samples were charged-discharged at room temperature for 1000 times, heated at 100 or 250 °C for up to 100 h, and applied at up to 250 °C, respectively. The results are presented in Fig. 10.

Figure 10a represents the E-J characteristic curves of a typical sample measured during repeating charge-discharge cycles. It is seen from this figure that the nonlinearity of the present ZnO-Pr$_6$O$_{11}$ thin-film varistor is recoverable after charge-discharge for 1000 times, indicating that the samples have good stability for repeating charge-discharge. Figure 10b and c displays the E-J plot of the samples at an operation temperature of 100 and 250 °C for up to 100 h, respectively. This plot indicates that at enhanced application temperature up to 250 °C, the nonlinearity of the present ZnO-Pr$_6$O$_{11}$ thin-film varistor is steady even after applying for 100 h, revealing that the samples have good stability for application at enhanced temperature. And Fig. 10d exhibits the E-J characteristic curves of a typical sample measured at varying temperature. It can be seen that the increase of application temperature up to 250 °C almost presents no effect on the nonlinearity of the present ZnO-Pr$_6$O$_{11}$ thin-film varistor, indicating that the samples have good stability for application at varying temperature.

When a ZnO varistor is working, it should withstand a certain steady-state voltage, because the heat produced when a current passes through a resistor can’t dissipate easily, and the temperature of the varistor will rise rapidly. After maintaining the initial steady state, traditional ZnO varistors quickly reached the possible heat breakdown conditions, the so-called aging. Therefore, the stability of a ZnO varistor at high-temperature operation is very important. In the present work, after charge-discharge at room temperature for 1000 times, heating at 100 or 250 °C for up to 100 h, or applying at up to 250 °C, the varistors still performed well. Therefore, such nanoscaled thin-film varistors will be very promising in electrical/electronic devices working at low voltage.

In summary, high-performance ZnO-Pr$_6$O$_{11}$ thin-film varistors were fabricated simply by hot-dipping oxygen-deficient zinc oxide thin films in Pr$_6$O$_{11}$ powder. The applied zinc oxide films had a composition of ZnO$_{0.81}$, and a thickness of about 200 nm, which were deposited onto conducting silicon chips by radio frequency magnetron sputtering of a sintered zinc oxide ceramic target. Then the films were hot-dipped in Pr$_6$O$_{11}$ at a temperature from 300 to 800 °C for an optimized time of 50 min in air. With the increase of hot-dipping temperature, the varistor voltage of the samples decreased from 0.02217 to 0.01623 V/nm, owing to the decreased number of effective grain boundaries in the varistors. The nonlinear coefficient increased initially and then decreased, presenting a recorded maximum of 39.29, when the temperature was 700 °C; correspondingly, the leakage current density first decreased and then increased, reaching a minimum of 0.02736 mA/cm$^2$, owing mainly to the
formation and destroying of complete zinc oxide/Pr₆O₁₁ grain boundaries. And after charge-discharge at room
temperature for 1000 times, heating at 100 and 250 °C for up to 100 h, or applying at up to 250 °C, the varistor
performance of the samples was still well kept. Due to its low sensitive voltage, high nonlinear coefficient, low
leakage current, and high stability, the present nanoscaled thin-film varistors would be very promising in elec-
tronic/electrical devices working at low voltage.

Methods

Samples preparation. The applied zinc oxide films had a composition of ZnOₐ₀.₈₁ and a thickness of about
200 nm, which were deposited onto conducting silicon chips by reactive RF magnetron sputtering of a sintered
zinc oxide ceramic target at ambient temperature. For details, please check them in refs 6 and 11. Then the as-de-
posited ZnOₐ₀.₈₁ thin film samples were buried in Pr₆O₁₁ powder in a half-covered alumina crucible, and heated
in a muffle furnace at a temperature of 300, 400, 500, 600, 700 and 800 °C, respectively, for an optimized time
of 50 min (see Extended Data Fig. 3). After the hot-dipping, the samples were cooled down naturally to room
temperature simply by shutting down the electricity of the furnace. Finally, silver paste was daubed at room tem-
perature on the chip surface as electrodes for the measurement of room-temperature electrical properties. But for
the varistor stability tests, the silver paste was toasted at 500 °C onto the samples in order to avoid the diffusion of
the electrode materials during the measurement.

Materials characterization. The elemental composition and chemical state of the samples were measured
by a Thermo Fisher X-ray photoelectron spectroscope (non-monochromated Al Kα radiation, photon energy
1486.7 eV). And the spectrometer was calibrated by the binding energy of C1s line (284.6 eV). The microstruc-
ture of the samples was examined on their surfaces and fractural cross-sections by FE-SEM (LEO-1530). From
the obtained SEM images, the grain size and film thickness of the samples were evaluated by the Nano Measurer
software. The elemental distribution in the samples was analyzed by an attached EDX system. The phase com-
position of the samples was identified by grazing incidence XRD (GI-XRD, D/αmax-RB, Cu Kα radiation, and
λ = 1.5418 Å) through a continuous scanning mode at a speed of 6°/min with an X-ray incidence angle of 0.5°.

Measurements of electrical properties. The E-J characteristics of the samples were recorded by a Keithley 2410 Multimeter. Through the recorded E-J curves, the nonlinear coefficient (α) of the samples was calculated according to the following equation:

Figure 10. E-J characteristic curves of the zinc oxide film varistors during stability tests. (a) when it was
measured at room temperature for 1000 times of repeating charge-discharge cycles; (b,c), when it was applied
at 100 and 250 °C for up to 100 h, respectively; and (d) when it was applied at a temperature up to 250 °C, during
which at each temperature, the sample was soaked at least for 5 min before measurement. All the samples were
prepared under the optimum conditions of hot-dipping ZnOₐ₀.₈₁ thin films in Pr₆O₁₁ at 700 °C for 50 min in air.
\[ \alpha = \frac{1}{\log E_{1mA} - \log E_{0.1mA}} \]

where the \( E_{1mA} \) and \( E_{0.1mA} \) are the electric fields corresponding to the current densities of 1.0 and 0.1 mA/cm\(^2\), respectively. The electric field at the current density of 1.0 mA/cm\(^2\) was defined as the varistor voltage \( (E_{1mA}) \), and the leakage current \( (I_L) \) was determined at 0.75\( E_{1mA} \). Excepting the varistor stability tests, all the \( E-J \) curves were measured at room temperature with each sample for one-time testing. But for the varistor stability tests of the optimum samples, their \( E-J \) curves were recorded after the samples were charged-discharged at room temperature for 1000 times, heated at 100 and 250 °C for up to 100 h, and applied at up to 250 °C, respectively. The applied voltage per grain boundary \( (V_{gb}) \) was calculated using:

\[ V_{gb} = E_{1mA} \cdot \frac{d}{D} \]

where \( E_{1mA} \) is the varistor voltage of the prepared film varistors, \( d \) is the average size of zinc oxide grains, and \( D \) is the thickness of the varistor thin films. The complex impedance spectra of the samples were recorded with an electrochemical workstation (CHI660E, CH Instrument Company, China) under the frequency ranging from 1 to 10\(^6\) Hz at an amplitude voltage of 20 mV. The impedance data were analyzed with the ZSimpWin program.

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Author Contributions
Y.W. and Q.W. prepared the ZnO-Pr₆O₁₁ thin-film varistors and characterized the electrical properties of the samples. C.B.W. examined the composition and structure of samples. Y.W., Z.J.P. and X.L.F. designed the experiments and interpreted the results. Y.W., Z.J.P. and X.L.F. wrote the main manuscript text, and all authors participated in the review of the manuscript. Z.J.P. and X.L.F. supervised the whole projects.

Additional Information

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