CeO$_2$ facets control: from single (100) to multiple†

Wenzhe Si, a,b Yu Wang, a,c Yue Peng, a Jianjun Chen a,* a and Junhua Li a

The synthesis of high energy facets and tunable low energy facets was achieved in CeO$_2$ films on a SrTiO$_3$ substrate by pulsed laser deposition. Three different facets of CeO$_2$ with distinctive morphologies appeared consecutively. Firstly, the (100) facet of CeO$_2$, which is the highest energy surface, was grown on the SrTiO$_3$ (STO) (100) substrate. The surface energy was decreased gradually with the increase of the laser pulse shots, and the (110) and (111) LEF appeared consecutively. The three different facets presented distinctive morphologies and the interface between each facet could be easily observed by the cross section of Transmission Electron Microscope (TEM) imaging. The interface between the (100) and (110) facet may exhibit excellent oxygen storage capacity, even better than the (100) facet.

1. Introduction

Oxide facets with distinctive crystallographic features possess varied atomic terminating characters, which are found to exhibit significant differences in magnetism, sensing and especially catalysis. 1–3 Rational design and orientable control of crystal facets are commonly adopted to optimize their performances. 4–7 High energy facets (HEF) that have abundant unsaturated active sites usually exhibit excellent properties. 8,9 But they are often unstable, and difficult to obtain by traditional chemical methods. 10–12 The exploration of advanced synthetic approaches to oxides with well-defined HEF is in urgent demand and of great importance.

In addition, some particular combinations of HEF and low energy facets (LEF) have been proven to show outstanding performances, even better than HEF. 1–3 Yu et al. found that the combination of TiO$_2$ anatase (001) (ca. 70%) and (101) (ca. 30%) facets was more active than the clean (001) which is the traditional highest energy surface of TiO$_2$. 9 Pan et al. synthesized (001) and (230) facets of BaTiO$_3$ sensitized with CdSe quantum dots, which exhibited better photocatalytic activity than (001) facet. 1 The superior activities were due to the charge separation and electronic reduction activities resulting from the synergy effect of (001) and (230) facets. However, controllable synthesis of the oxides with dominant HEF and tunable LEF is still a great challenge.

Here a facile method to controllably synthesize and tune the oxides facets by pulsed laser deposition (PLD) was reported (Scheme 1). CeO$_2$, which is an important material utilizing in fuel solar cells, gates for metal–oxide semiconductor devices, phosphors and three-way catalysts, was selected as a target. 13–18 Firstly, (100) facet of CeO$_2$, which is the highest energy surface, was grown on the SrTiO$_3$ (STO) (100) substrate. 19,20 The high laser power induced CeO$_2$ to grow epitaxially along (100) facet. The surface energy was decreased gradually with the increase of the laser pulse shots, and (110) and (111) LEF appeared consecutively. The three different facets presented distinctive morphologies and the interface between each facet could be easily observed by the cross section of Transmission Electron Microscope (TEM) imaging. The terminated surface could be tuned by the thickness control by PLD. This method may provide a new sight for the controllable synthesis of the oxides with tunable facets, especially for those oxides with distinctive morphologies on different facets.

Scheme 1 Synthesis route of the CeO$_2$ films on SrTiO$_3$ substrate.
2. Materials and methods

2.1 Chemicals

CeO$_2$ powder was purchased from Shanghai Yuelong Company of analytical grade purity and used without any further purification. STO (100), STO (110) and Si (100) substrates were purchased from Hefei Kejing Materials Technology Company, LTD. The CeO$_2$ powder was finely grounded and then pressed into a pellet (diameter is 2 cm, and thickness is 0.8 cm). Afterwards, the resulting pellet was calcined to target at 1000 K for 10 h in the air.

2.2 Preparation of CeO$_2$ films

The CeO$_2$ films can be achieved on STO (100), STO (110) and Si (100) substrates by the PLD method using a KrF excimer laser with a wavelength of 248 nm, a laser fluence of 1.5 J cm$^{-2}$ and a repetition rate of 3 Hz. The distance between a ceramic target and a substrate is 8 cm. Here the CeO$_2$ films were deposited on STO (100) for different pulse shots: 1000, 5000, 10 000, and 50 000, respectively. STO (110) and Si (100) for 5000 shots. The temperatures of films growing are 800 K. Then the films were annealed maintained at the growth temperature for 0.5 h. The temperature was subsequently lowered at a rate of 1 K min$^{-1}$. The films with different deposition pulse shots for 1000, 5000, 10 000, and 50 000 are abbreviated as S1, S5, S10, and S50, respectively.

2.3 Characterization

The crystal structure of the CeO$_2$ films were identified by X-ray diffraction (XRD, Rigaku D/MAX 2500 V/PC) with Cu K$_\alpha$ radiation (\(\lambda = 1.5418 \text{ Å}\)) at 40 kV and 30 mA at room temperature. Scanning electron microscopy (SEM) images were acquired using a focused ion beam (Helios Nanolab 600i). Since the CeO$_2$ films were poorly conductive, the conductive adhesive was used to fix the film on the specimen holder and avoid charge accumulation when ion beam irradiation.

The surface morphology was measured by an atomic force microscopy (AFM, SPI 3800 N Probe Station + SPA 400 unit system), which was manufactured by SII NanoTechnology Inc. DFM (Dynamic Force Mode) scanning mode was selected in the experiments mentioned above, while the vibration frequency of cantilevers is about 150 KHz.

Transmission electron microscopy (TEM) was carried out using a JEM-2100F at 200 kV. X-ray photoemission spectroscopy (XPS, Thermo ESCAlab 250) experiment was performed using monochromatic Al K$_\alpha$ radiation (1486.6 eV). TEM specimens were extracted from the cross-section of CeO$_2$ film by the FIB lift-out technique. To prevent damage to the film from focused Ga$^{3+}$ ion beam, W(CO)$_6$ gas [W(CO)$_6$(IBD)] was injected to the interested region with electron beam deposition to form a thin protective layer of tungsten composite. After W(CO)$_6$(IBD), focused ion beam deposition was continued in case of damage from milling process. The lift-out specimen was mounted onto a crescent copper grid with FIB deposition of W(CO)$_6$ [W(CO)$_6$(EBD)] and then mill it to the right thickness. After a low voltage bombard process, amorphous layer, which formed during the milling process, were removed and the specimen was transferred to TEM characterization.

X-ray photoelectron spectroscopic (XPS) was used with Mg K$_\alpha$ (\(h\nu = 1253.6 \text{ eV}\)) X-ray source to determine the Ce 3d binding energies (BEs) of surface cerium species. The C 1s signal (BE = 284.6 eV) of contaminant carbon was employed to calibrate the BE values of Ce 3d.

3. Results and discussion

3.1 XRD results

The X-ray diffraction (XRD) patterns of CeO$_2$ films with different pulse shots deposited on STO (100) substrates are shown in Fig. 1. All the peaks have been normalized. For the S$_1$ sample, only (200) plane of CeO$_2$ cubic fluorite phase can be observed. When the pulse shots increased up to 5000, (400) plane appeared and the peak intensity of (200) was enhanced. Based on the results of S$_1$ and S$_5$, which represent the initial growing stage, the CeO$_2$ film has a preferred orientation along the (100) direction. For the S$_{10}$ sample, (200) and (400) planes of CeO$_2$ still existed and were enhanced in intensity, compared to S$_1$ and S$_5$.

3.2 Morphologies

Then Scanning Electron Microscope (SEM) and Atomic Force Microscope (AFM) were carried out to show the morphologies of these three different facets, and detect the surface where (110) and (111) planes grow on (Fig. 2). The conductive adhesive was used when the SEM images were acquired. The role of
Conductive adhesive is usually divided into two parts: one role is to avoid charge accumulation. It is a common approach to improve the SEM resolution. Another role is to fasten the sample. It can be seen that the shape of S1 was 3D linen-like nanorods. However, its morphology did not show good uniformity. For the S2 sample, the grid structure became more regular (Fig. 2b). The nanorods with two different directions were almost equal in length and diameter, and the size of the pores (20–50 nm) was stacked by the nanorods, which may create large surface area in the framework. For Fig. 2c, a few nanosheets (shown in the red circles) with triangle form grew on top of the nanorods. Correlating with the XRD results that (100) and (110) facets coexisted for S10, it can be inferred that the triangle-like nanosheets with (110) facet was grown up from the nanorods, which could be attributed to the decrease of surface energy. For the S50 sample (Fig. 2d), new structure consisted of the star-like nanosheets appeared. What’s more, single (110) and (111) facets of CeO2 films were synthesized separately on STO (211) and Si (001) substrates (Fig. S3 and S4†). It can be seen that the morphologies were corresponding to what we obtained on STO (001) by tuning the thickness of films. The results suggested that three different structures with distinctive morphologies could be formed on S50 sample simultaneously, and each structure was oriented along to the different directions.

### 3.3 TEM results

For local crystal lattice determination, a small cross section of S50 was prepared by Focused Ion Beam (FIB) for TEM imaging. Different morphologies of CeO2 facets and the transformation position of each structure can be easily observed from Fig. 3a. The region 1, 2 and 3 represented the linen-like nanorods, triangle-like nanosheets and star-like nanosheets, respectively.

**Fig. 2** SEM images of CeO2 films which are labeled as (a, b, c, and d) for sample S1, S5, S10 and S50, respectively; the insets are corresponding AFM images.

**Fig. 3** (a) TEM characterization of cross section of S50. (b and c) HRTEM of region a and b in (a), respectively.

Ce 3d X-ray Photoelectron Spectroscopy (XPS) spectra of CeO2 films with different pulse shots are shown in Fig. 4. The peaks labeled U are due to 3d3/2 spin–orbit states, and those labeled V are the corresponding 3d5/2 states. The U3/V3 doublet is due to the primary photoemission from Ce4+–O2. The U/V and U’/V’ doublets are shakedown features resulting from the transfer of one or two electrons from a filled O 2p orbital to an empty Ce 4f orbital. The U1/V1 doublet is due to photoemission from Ce3+ cations. This shakedown feature gives rise to an additional doublet, which is labeled as U0/V0 in the spectrum. Ce3+ ratio showed a downward trend by increasing the pulse shots, except for S10. The oxygen storage capacity of CeO2 mainly relies on the intrinsic oxygen vacancies of the surface, which is attributed to the rapid redox cycles of Ce4+ ↔ Ce3+. In CeO2 structure, oxygen storage capacity is related with the Ce3+/Ce4+ ratio. The more the amount of Ce3+ exists, the more the oxygen vacancies exists. The material can adsorb more oxygen with the
increase of the amount of oxygen vacancy. As the interface between the (100) and (110) facets possessed the largest amount of Ce$^{3+}$ ratio (25.99%), it is indicated that it may possess the highest oxygen storage capacity.

4. Conclusions

In summary, a facile PLD method was presented for the synthesis and tuning of the exposed facets of CeO$_2$ films. First, (100) HEF of CeO$_2$ was grown on the STO (100) substrate. With increasing the pulse shots, (110) and (111) LEF appeared consecutively resulting from the decrease of surface energies. Each facet had distinctive morphologies and the positions of the interfaces between each facet can be directly observed from the cross-section of the CeO$_2$ film by TEM. The interface between (100) and (110) facet showed highest Ce$^{3+}$ ratio, which led to the excellent oxygen storage capacity. This work provides a simple way to tune CeO$_2$ film from single HEF to multiple, and shows potential possibility to controllably synthesize the oxides with dominant HEF and tunable LEF.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This work was financially supported by the National Natural Science Foundation of China 21876093, and the National Key Research and Development Program (2017YFC0210700 and 2018YFC0213400).

References

1. D. Zhong, W. Liu, P. Tan, A. Zhu, Y. Liu, X. Xiong and J. Pan, Appl. Catal., B, 2018, 227, 1–12.
2. K. X. Yao, X. M. Yin, T. H. Wang and H. C. Zeng, J. Am. Chem. Soc., 2010, 132, 6131–6144.
3. Q. Xiang, K. Lv and J. Yu, Appl. Catal., B, 2010, 96, 557–564.
4. Y. Luan, L. Jing, Y. Xie, X. Sun, Y. Feng and H. Fu, ACS Catal., 2013, 3, 1378–1385.
5. L. Ruan, H. Ramezani-Dakhel, C.-Y. Chiu, E. Zhu, Y. Li, H. Heinz and Y. Huang, Nano Lett., 2013, 13, 840–846.
6. X. Xie, Y. Li, Z.-Q. Liu, M. Haruta and W. Shen, Nature, 2009, 458, 746–749.
7. B. Lim, M. Jiang, P. H. C. Camargo, E. C. Cho, J. Tao, X. Lu, Y. Zhu and Y. Xia, Science, 2009, 324, 1302–1305.
8. M. Leng, M. Liu, Y. Zhang, Z. Wang, C. Yu, X. Yang, H. Zhang and C. Wang, J. Am. Chem. Soc., 2010, 132, 17084–17087.
9. Z.-Y. Zhou, N. Tian, J.-T. Li, I. Broadwell and S.-G. Sun, Chem. Soc. Rev., 2011, 40, 4167–4185.
10. H. B. Jiang, Q. Cuan, C. Z. Wen, J. Xing, D. Wu, X.-Q. Gong, C. Li and H. G. Yang, Angew. Chem., Int. Ed., 2011, 50, 3764–3768.
11. J. Yin, Z. Yu, F. Gao, J. Wang, H. Pang and Q. Lu, Angew. Chem., Int. Ed., 2010, 49, 6328–6332.
12. X. Han, M. Jin, S. Xie, Q. Kuang, Z. Jiang, Y. Jiang, Z. Xie and L. Zheng, Angew. Chem., Int. Ed., 2009, 48, 9180–9183.
13. Q. Fu, H. Saltsburg and M. Flytzani-Stephanopoulos, Science, 2003, 301, 935–938.
14. E. P. Murray, T. Tsai and S. A. Barnett, Nature, 1999, 400, 649–651.
15. A. Corma, P. Atienzar, H. Garcia and J.-Y. Chane-Ching, Nat. Mater., 2004, 3, 394–397.
16. J. Ke, J.-W. Xiao, W. Zhu, H. Liu, R. Si, Y.-W. Zhang and C.-H. Yan, J. Am. Chem. Soc., 2013, 135, 15191–15200.
17. A. H. Morshed, M. E. Moussa, S. M. Bedair, R. Leonard, S. X. Liu and N. El-Masry, Appl. Phys. Lett., 1997, 70, 1647–1649.
18. X. Wang, D. Liu, S. Song and H. Zhang, J. Am. Chem. Soc., 2013, 135, 15864–15872.
19. P. W. Tasker, J. Phys. C: Solid State Phys., 1979, 12, 4977–4984.
20. J. Conesa, Surf. Sci., 1995, 339, 337–352.
21. Y. Xu and R. Li, Appl. Surf. Sci., 2018, 455, 997–1004.
22. P. Min, S. Zhang, Y. Xu and R. Li, Appl. Surf. Sci., 2018, 448, 435–443.