# Chemical Science



## **EDGE ARTICLE**

View Article Online
View Journal | View Issue



Cite this: Chem. Sci., 2018, 9, 2469

# Bifunctional CO oxidation over Mn-mullite anchored Pt sub-nanoclusters *via* atomic layer deposition†

Xiao Liu, <sup>1</sup>D<sup>a</sup> Yuanting Tang, <sup>b</sup> Meiqing Shen, <sup>1</sup>D<sup>c</sup> Wei Li, <sup>d</sup> Shengqi Chu, <sup>e</sup> Bin Shan <sup>1</sup>D\*<sup>b</sup> and Rong Chen <sup>1</sup>D\*<sup>a</sup>

CO oxidation is a widely used model system for understanding fundamental aspects of heterogeneous catalysis. While platinum (Pt) continues to be a reference material for CO oxidation catalysis, poisoning of Pt catalysts presents a critical issue that blocks reaction sites and impedes subsequent reaction steps. Fabrication of CO poison-free Pt catalysts remains a great challenge due to its CO-philic nature. Herein, we report a Pt based catalyst to effectively tackle CO poisoning by tightly anchoring Pt sub-nanoclusters onto Mn-mullite oxide (SmMn<sub>2</sub>O<sub>5</sub>) *via* atomic layer deposition. Superior CO oxidation activity has been observed with a significantly lowered light-off temperature and apparent activation energy. *In situ* diffuse reflectance infrared Fourier transform spectroscopy analysis, oxygen isotope experiments and density functional theory calculations confirm that the low-temperature activity originates from active oxygen atom sources at the bifunctional interface structure.

Received 28th December 2017 Accepted 26th January 2018

DOI: 10.1039/c7sc05486f

rsc li/chemical-science

#### Introduction

Catalytic oxidation of CO over platinum (Pt) metal has been the subject of extensive investigations since the classical work of Langmuir, and has found important technological applications in automotive exhaust emission control and fuel cells. Overall, Pt has shown excellent performance as well as stability against oxidation and compound formation, making it the *de facto* standard reference catalyst. However, one of the critical problems that plague Pt catalysts is the well-known CO poisoning effect that limits its activity by the blockage of surface active sites and subsequent reaction steps under low temperatures. Such phenomena are quite general considering that CO will be used as a reactant or generated as a reaction intermediate in many catalytic processes such as CO oxidation and

water–gas shift reactions, and in fuel cells.  $^{7-9}$  In view of the high dispersion (smaller Pt nanoparticles) required for catalyst applications, CO poisoning will be only more pronounced, wherein a large fraction of coordinatively unsaturated edge and vertex sites over-bind CO molecules.  $^{10-13}$  Iglesia *et al.* have reported that the CO molecules binding on Pt atoms with low coordination numbers are stronger than those on Pt (111), due to the greater electron donation of Pt atoms on nanocluster surfaces. The average CO binding energy on Pt nanoclusters at full coverage is also greater than that on Pt (111), due to the weaker repulsive interactions among adsorbed CO molecules.  $^{13}$  Overall, the high CO coverage will hinder the adsorption and dissociation of  $\mathrm{O}_2$  molecules on Pt facets, which agrees well with our theoretical predictions.  $^{14}$ 

Considerable efforts have been devoted to eliminating the CO poisoning problem, primarily via electronic or bifunctional mechanisms. In the electronic mechanism, Pt metal is alloyed with other transitional metals to form multi-component/coreshell nanoparticles that have modified electronic structures and weakened CO binding strengths.15-17 However, the element distribution of the alloy catalysts is difficult to control and competitive adsorption cannot be completely suppressed, especially under the circumstance that other reactants interact more weakly with Pt than CO. The bifunctional mechanism, on the other hand, is a highly efficient way of mitigating CO poisoning by providing spatially separated sites for oxygen species. This is typically achieved by utilizing reducible transition metal oxide supports like CeO<sub>2</sub>, FeO<sub>x</sub> and Co<sub>3</sub>O<sub>4</sub>, where the interfacial lattice oxygen of Pt-oxide composites therein can participate in the redox reactions. 18-22 The oxide support must

<sup>&</sup>lt;sup>a</sup>State Key Laboratory of Digital Manufacturing Equipment and Technology, School of Mechanical Science and Engineering, Huazhong University of Science and Technology, Wuhan 430074, Hubei, People's Republic of China. E-mail: rongchen@mail.hust.edu.

bState Key Laboratory of Materials Processing and Die and Mould Technology, School of Materials Science and Engineering, Huazhong University of Science and Technology, Wuhan 430074, Hubei, People's Republic of China. E-mail: bshan@mail.hust.edu.cn 'School of Chemical Engineering and Technology, Tianjin University, Tianjin 300072, People's Republic of China

<sup>&</sup>lt;sup>d</sup>General Motors Global Research and Development, Chemical Sciences and Materials Systems Lab, 3500 Mound Road, Warren, Michigan 48090, USA

<sup>&#</sup>x27;Institute of High Energy Physics, Chinese Academy of Sciences, Beijing 100049, People's Republic of China

<sup>†</sup> Electronic supplementary information (ESI) available: Catalyst preparation and characterization details, supplemental figures and tables. See DOI: 10.1039/c7sc05486f

possess high bulk oxygen mobility so that oxygen species at the interface can be quickly consumed and replenished to sustain the reaction. Unfortunately, the sluggish oxygen diffusion rate would seriously limit the activity of the composite catalysts, and

the oxygen transport channel may be poisoned by products or intermediates, which can lead to performance degradation. 23-26 Therefore, design of a bifunctional catalyst that can supply oxygen species not limited by the bulk oxygen diffusion opens up new opportunities in the development of high-performance CO oxidation catalysts.

Herein, we report a novel Pt based composite catalyst that has the unique capability of directly dissociating gas phase oxygen molecules on the oxide site, overcoming the bulk oxygen diffusion limit for supplied oxygen species in conventional bifunctional catalysts. The composite catalyst is synthesized *via* atomic layer deposition (ALD) of Pt clusters on SmMn<sub>2</sub>O<sub>5</sub> (SMO) mullite-type oxides, which show excellent low temperature CO oxidation activity and low apparent activation energy. In situ diffuse reflectance infrared Fourier transform spectroscopy (DRIFTS), <sup>18</sup>O isotope-labelling experiments and density functional theory (DFT) calculations confirm that the active oxygen supplied by the bifunctional interface is the origin of the low temperature CO oxidation activity.

#### Results and discussion

**Chemical Science** 

The highly dispersed Pt sub-nanoclusters are synthesized via ALD on SMO mullite-type oxides, which are in the shape of irregular ellipsoids with an average size of about 50 nm (Fig. S1†). As shown in Fig. 1a, the Pt clusters of the as-prepared Pt/SMO catalysts are on a sub-nano scale of 0.5-0.9 nm. The bright spots in the high-angle annular dark-field scanning

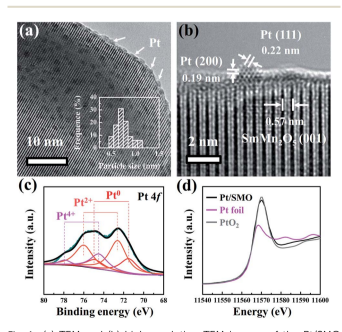


Fig. 1 (a) TEM and (b) high resolution TEM images of the Pt/SMO catalyst are presented. The size distribution of the Pt clusters has been inserted in (a). (c) The Pt 4f XPS spectrum of the Pt/SMO catalyst. (d) The normalized Pt L<sub>III</sub>-edge XANES of Pt/SMO, Pt foil and PtO<sub>2</sub>.

transmission electron microscopy (HAADF-STEM) images represent the uniformly distributed Pt clusters on the SMO supports (Fig. S2†), which agrees well with the lack of an appreciable Pt diffraction peak in the XRD pattern of Pt/SMO (Fig. S3†). The lattice fringes with a d-spacing of 0.57 nm in Fig. 1b are assigned to the SMO (001) planes. The lattice fringes with a d-spacing of 0.22 nm and 0.19 nm are assigned to the Pt (111) and (200) planes, respectively.

The XPS results (Fig. 1c) show that the main state of the Pt clusters can be attributed to oxidized Pt (Pt2+, 56% and Pt4+, 20%) and the other Pt species are in the state of metallic Pt atoms (Pt<sup>0</sup>, 24%). According to the morphology of Pt clusters, we assigned the oxidized Pt species to the interfacial Pt atoms bound to surface oxygen atoms and the metallic Pt atoms to the Pt atoms far away from the interface. The large amount of oxidized Pt indicates the high dispersion of Pt clusters and strong interfacial interactions. Consistent with the XPS analysis, the Pt L<sub>III</sub>-edge X-ray absorption near-edge structure (XANES) spectra (Fig. 1d) show that the Pt/SMO catalyst has a large amount of oxidized Pt species, with a white line intensity similar to that of the PtO2 sample, which further implies the strong chemical anchoring of Pt clusters on the SMO oxides. The Fourier transform extended X-ray absorption fine structure (EXAFS) spectrum (Fig. S4†) shows a strong peak at about 1.7 Å, which can be attributed to the Pt-O bonds. The fitted Fourier transform EXAFS spectra show that the fitted Pt-O bond length of Pt/SMO is about 2.007 Å and the coordination number is about 3.51 (Table S2†). The shorter Pt-O bond length and smaller Pt-O coordination number compared to those of the PtO<sub>2</sub> sample also indicate the Pt/SMO interfaces as the origin of the Pt-O bonds.

A CO oxidation activity test has been performed to evaluate the CO-tolerant performance of Pt/SMO catalysts at low temperature. As shown in Fig. 2a Pt/SMO exhibits excellent low temperature CO oxidation activity with a  $T_{50}$  (50% CO

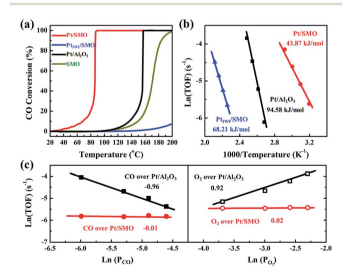


Fig. 2 (a) The CO conversion of Pt/SMO, Pt<sub>IWI</sub>/SMO, Pt/Al<sub>2</sub>O<sub>3</sub> and SMO as a function of reaction temperature. (b) The Arrhenius plots of CO oxidation rates of Pt/SMO, Pt<sub>IWI</sub>/SMO and Pt/Al<sub>2</sub>O<sub>3</sub>. (c) The reaction orders of CO and O2 over Pt/SMO and Pt/Al2O3.

**Edge Article** 

conversion temperature) of about 86 °C, which is much lower than that of pure SMO (171 °C). It also shows excellent structural stability in multi-cycle repeated activity tests (Fig. S5†). As a control experiment, the incipient wetness impregnation method has also been used to construct the Pt/SMO composite catalyst (Pt<sub>IWI</sub>/SMO, Fig. S6†). Pt<sub>IWI</sub>/SMO exhibited poorer activity than pure SMO. The Mn 2p XPS spectra show that the concentrations of Mn<sup>3+</sup> and Mn<sup>4+</sup> ions of Pt<sub>IWI</sub>/SMO are quite different to that of pure SMO and Pt/SMO, indicating the change of surface active sites during Pt impregnation (Fig. S7†). Moreover, Pt<sub>IWI</sub>/SMO has a large amount of Pt<sup>4+</sup> oxidation states (0.88) and the H<sub>2</sub> reduction treatment can slightly enhance its activity by yielding more Pt2+ and Pt0 states (Fig. S8 and S9†). However, the enhanced activity of Pt<sub>IWI</sub>/SMO is still much lower than that of Pt/SMO, implying the key role of the interface structure to the high activity of the Pt/SMO catalyst. The Pt clusters have also been deposited on Al<sub>2</sub>O<sub>3</sub> supports as a reference sample (Pt/Al<sub>2</sub>O<sub>3</sub>) by the same ALD process as for Pt/SMO. The TEM images (Fig. S10†) show that the average size of deposited Pt clusters is about 1.19 nm and there is also no characteristic Pt peak in the XRD pattern of Pt/Al<sub>2</sub>O<sub>3</sub>, indicating a high dispersion of small Pt clusters as well. The Pt clusters on Al<sub>2</sub>O<sub>3</sub> supports are mainly in the metallic Pt state (0.62) per the Pt 4d XPS spectrum shown in Fig. S11,† which is a result of the weaker interactions between the Pt clusters and Al<sub>2</sub>O<sub>3</sub> supports. The measured  $T_{50}$  of Pt/Al<sub>2</sub>O<sub>3</sub> is about 157 °C, which is due to the CO poisoning effect at low temperatures on Pt facets.

The intrinsic behavior of Pt clusters for CO oxidation on both SMO and Al<sub>2</sub>O<sub>3</sub> supports has been further investigated by performing kinetic tests to eliminate the thermal and diffusion effects (Fig. S12†). Arrhenius plots of reaction rates for Pt/SMO, Pt<sub>IWI</sub>/SMO and Pt/Al<sub>2</sub>O<sub>3</sub> are shown in Fig. 2b. The calculated apparent activation energy of Pt/SMO (43.87 kJ mol<sup>-1</sup>) is smaller than that of Pt<sub>IWI</sub>/SMO (68.21 kJ mol<sup>-1</sup>) and about half of that for Pt/Al<sub>2</sub>O<sub>3</sub> (94.58 kJ mol<sup>-1</sup>). Despite the high gas hourly space velocity of 120 000 mL g<sup>-1</sup> h<sup>-1</sup> used, Pt/SMO catalysts exhibit a much lower  $T_{50}$  and apparent activation energy compared with other oxide supported Pt catalysts in previous studies (Table S3†). The reaction orders of Pt/SMO and Pt/Al<sub>2</sub>O<sub>3</sub> with respect to CO and O<sub>2</sub> are presented in Fig. 2c. The reaction orders of CO and O<sub>2</sub> over Pt/Al<sub>2</sub>O<sub>3</sub> are close to −1 and 1, suggesting CO inhibition in the low temperature region and the typical Langmuir-Hinshelwood reaction mechanism.27,28 The active oxygen species during CO oxidation over Pt/Al<sub>2</sub>O<sub>3</sub> are supplied by the O2 dissociation on the surface of Pt clusters, which will be inhibited by the strongly bound CO molecules at low temperatures. The reaction orders of CO and O2 over Pt/SMO are close to zero, indicating negligible competitive adsorption between CO and O2 during CO oxidation over Pt/SMO. Thus, the supply of active oxygen species for CO oxidation over Pt/SMO may be independent of the Pt sites covered by CO molecules.

In order to clarify the origin of the active oxygen species in the CO oxidation over Pt/SMO, the CO adsorption and  $O_2$  dissociation behaviors have been investigated. As shown in Fig. 3a, the *in situ* DRIFTS spectra show that there are three sets of peaks after CO molecular adsorption (black lines) on Pt/SMO at room temperature (25 °C), which are assigned to the CO

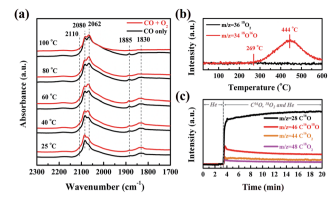


Fig. 3 (a) In situ DRIFTS spectra of Pt/SMO with CO adsorption (black lines) and oxidation (red lines) at different reaction temperatures. (b) Changes of m/z=34 and 36 as a function of temperature for the  $^{18}{\rm O}_2$  pretreated Pt/SMO catalyst during the TPIE experiment. (c) Changes of m/z=28, 44, 46 and 48 during CO oxidation with  ${\rm C}^{16}{\rm O}$  and  ${\rm ^{18}O}_2$  at 80 °C as a function of time for the Pt/SMO catalyst after  ${\rm ^{16}O}_2$  pretreatment.

molecules adsorbed at the bridge sites (1830 cm<sup>-1</sup> and  $1885 \text{ cm}^{-1}$ ), the top sites (2060 cm<sup>-1</sup> and 2080 cm<sup>-1</sup>, as well as the broad shoulder (1950-2050 cm<sup>-1</sup>)) and the oxidized Pt clusters (2110 cm<sup>-1</sup>).<sup>29-31</sup> Only the bridge-bonded CO molecules (1830 cm<sup>-1</sup>) will be desorbed as the temperature increases to 100 °C. Upon introduction of the oxygen gas feed, the bridgebonded CO molecules can be observed to react with the injected O<sub>2</sub> at 80 °C (red line), while the linear-bonded CO molecules are very stable on the Pt clusters. Moreover, after all of the bridge-bonded CO molecules have reacted with O2 at 100 °C, large amounts of linear-bonded CO molecules remain on Pt clusters. Therefore, it can be deduced that the bridge-bonded CO molecules at the interface are the primary active source for CO oxidation at low temperature. As a comparison, the in situ DRIFTS spectra of CO oxidation over Pt/Al<sub>2</sub>O<sub>3</sub> (Fig. S13†) show that the injected O2 will not react with the adsorbed CO molecules when the temperature is below 140 °C. When the temperature reaches 160 °C (higher than  $T_{50}$ ), the bridgebonded and linear-bonded CO molecules on the Pt clusters both disappear rapidly, showing no apparent selectivity for the oxidation of these two types of CO molecule.

To further identify the source of active oxygen during CO oxidation, an isotope-labelling method was implemented. The temperature programmed isotope exchange (TPIE) experiment for  $^{18}O_2$  pretreated Pt/SMO shows that the initial temperature of hetero-exchange between the gas phase and solid surface is about 269 °C (Fig. 3b). There is no obvious hetero-exchange at low temperatures, indicating the low activity of lattice oxygen species for low temperature reactions. Specifically, we have utilized an isotope-labelling method to study the source of active oxygen for CO oxidation over the  $^{16}O_2$  pretreated Pt/SMO catalyst at 80 °C. As shown in Fig. 3c, after  $C^{16}O$  and  $C^{18}O_2$  are injected into the system, the mass intensity of  $C^{16}O$  will be sharply increased. Simultaneously, the increased mass intensity of  $C^{16}O_2$ ,  $C^{16}O^{18}O$  and  $C^{18}O_2$  (m/z=44, 46 and 48) can be related to the CO oxidation reaction at 80 °C. Among the three

product species, the trace amount of  $C^{18}O_2$  is almost negligible. Depending on the source of oxygen atoms,  $C^{16}O_2$  and  $C^{16}O^{18}O$  can be assigned to the process of  $C^{16}O$  reacting with lattice oxygen ( $^{16}O$ ) and dissociated oxygen ( $^{18}O$ ), respectively. Fig. 3c demonstrates that  $C^{16}O^{18}O$  is the dominant species in the  $CO_2$  products, confirming that dissociated oxygen rather than lattice oxygen is the main source of active oxygen for low temperature CO oxidation. As most Pt sites on Pt clusters have been poisoned by CO molecules, the dissociation takes place on the active sites of the SMO surface at the Pt/SMO interface.

**Chemical Science** 

DFT calculations were employed to elucidate the dissociation process of O2 at the Pt/SMO interface. The pyramidal shaped Pt<sub>10</sub> cluster has been selected considering that it is the most stable cluster in the gas phase.32,33 Interface geometry relaxation of the structure shows that Pt<sub>10</sub> has been effectively anchored on the SMO (010) facet with an exposed Mn<sub>2</sub> dimer (Fig. S14†), which is a thermodynamically stable species and has been shown to possess high activity towards O<sub>2</sub> dissociation in our previous theoretical study.34 CO adsorption on the exposed Mn2 dimer is very weak (-0.31 eV) and the binding strength of CO on Pt at the interface is much weaker than that on Pt sites far away from the interface (Fig. S15†). This can be attributed to the more-vacant d orbitals of interfacial Pt atoms due to outward charge transfer to the SMO support (Fig. S16†), which is consistent with our XPS results and other Pt/oxide systems in previous studies.31,35 Considering the strong poisoning of CO (<-2 eV) on Pt atoms away from the interface, we have constructed a CO covered Pt/SMO model (Fig. S17†) as the initial state for the catalytic cycle calculation. Competitive adsorption energy analysis of CO and O<sub>2</sub> at the Pt/SMO interface (Fig. S17†) shows that CO adsorption prefers the bridged sites of interfacial Pt, and O<sub>2</sub> prefers the Mn<sub>2</sub> dimer, creating an effective spatially separated adsorption for the two reactants. The complete

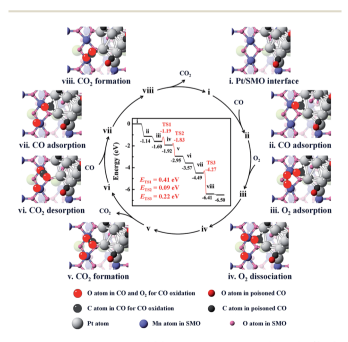


Fig. 4 The energetic route of CO oxidation at the interface of Pt/SMO. The catalytic cycle can be summarized as  $2CO + O_2 \rightarrow 2CO_2$ .

reaction route of CO oxidation at the Pt/SMO interface is illustrated in Fig. 4. The CO molecule initially adsorbs on the Pt interfacial site with an adsorption energy of −1.14 eV and the O<sub>2</sub> molecule adsorbs on the Mn2 dimer with an adsorption energy of -0.45 eV. Subsequently, the adsorbed oxygen will dissociate into active oxygen atoms on the Mn2 site with a low barrier energy of 0.41 eV. CO molecules can react with active oxygen atoms sequentially and form CO2 molecules with barrier energies of 0.09 eV and 0.22 eV, respectively. During the whole reaction route, there is no competitive adsorption between CO and O2 at the interface, and the CO adsorbed on the Pt sites cannot suppress the O2 adsorption and dissociation steps on the Mn dimers, which is consistent with the zero-order rate results for the Pt/SMO catalyst. Overall, the O2 dissociation step on the Mn<sub>2</sub> component is the rate determining step (RDS) of CO oxidation over the Pt/SMO interface with a barrier energy of  $0.41 \text{ eV} (39.51 \text{ kJ mol}^{-1})$ , which is comparable to the experimentally measured apparent activation energy. Comparing the barrier energies of O2 dissociation and CO2 formation of Pt/ SMO to that at the interface of Pt clusters and other oxides in previous studies (Table S4†), Pt/SMO shows a relatively low reaction barrier and can be more active towards CO oxidation. This indicates that the Pt/SMO interface plays a key role in bifunctional CO oxidation catalysis.

#### Conclusions

In conclusion, a bifunctional Pt/SMO interfacial structure is synthesized by tightly anchoring Pt sub-nanoclusters on SMO oxides, which provides spatially separated sites for CO and O<sub>2</sub>. With CO binding to Pt and effective O<sub>2</sub> dissociation on the Mn<sub>2</sub> dimer, the interface serves as an efficient poison-free CO oxidation site. The proposed catalytic reaction mechanism of CO oxidation over Pt/SMO shows that the high activity of O<sub>2</sub> dissociation at the bifunctional interface is the key to the low temperature CO oxidation activity. Our work on the bifunctional catalyst removes the lattice oxygen diffusion limit and sheds light on the design of new poison-free CO oxidation catalysts.

#### Conflicts of interest

There are no conflicts to declare.

# Acknowledgements

This work is supported by the National Natural Science Foundation of China (51575217 and 51572097). R. C. acknowledges the Thousand Young Talents Plan, the Recruitment Program of Global Experts, and the Program for Changjiang Scholars and Innovative Research Team in University (IRT13017). We would also like to acknowledge the technology support from the Analytic Testing Center and Flexible Electronics Research Center of HUST. The calculations have been performed at the Texas Advanced Computing Center (TACC) at The University of Texas at Austin (http://www.tacc.utexas.edu).

**Edge Article Chemical Science** 

## Notes and references

- 1 I. Langmuir, Trans. Faraday Soc., 1922, 17, 621-654.
- 2 M. Bowker, Chem. Soc. Rev., 2008, 37, 2204-2211.
- 3 M. K. Debe, Nature, 2012, 486, 43-51.
- 4 Q. S Chen, F. J. Vidal-Iglesias, J. Solla-Gullón, S. G. Sun and J. M. Feliu, Chem. Sci., 2012, 3, 136-147.
- 5 H. Falsig, B. Hvolbæk, I. S. Kristensen, T. Jiang, T. Bligaard, C. H. Christensen and J. K. Nørskov, Angew. Chem., Int. Ed., 2008, 47, 4835-4839.
- 6 S. B. Vendelbo, C. F. Elkjær, H. Falsig, I. Puspitasari, P. Dona, L. Mele, B. Morana, B. J. Nelissen, R. van Rijn, J. F. Creemer, P. J. Koovman and S. Helveg, Nat. Mater., 2014, 13, 884-890.
- 7 M. A. Newton, D. Ferri, G. Smolentsev, V. Marchionni and M. Nachtegaal, Nat. Commun., 2015, 6, 8675.
- 8 Q. Fu, H. Saltsburg and M. Flytzani-Stephanopoulos, Science, 2003, 301, 935-938.
- 9 S. Alayoglu, A. U. Nilekar, M. Mavrikakis and B. Eichhorn, Nat. Mater., 2008, 7, 333-338.
- 10 A. D. Allian, K. Takanabe, K. L. Fujdala, X. H. Hao, T. J. Truex, J. Cai, C. Buda, M. Neurock and E. Iglesia, J. Am. Chem. Soc., 2011, 133, 4498-4517.
- 11 M. Moses-DeBusk, M. Yoon, L. F. Allard, D. R. Mullins, Z. L. Wu, X. F. Yang, G. Veith, G. Malcolm Stocks and C. K. Narula, J. Am. Chem. Soc., 2013, 135, 12634-12645.
- 12 S. Bonanni, K. Aït-Mansour, W. Harbich and H. Brune, J. Am. Chem. Soc., 2012, 134, 3445-3450.
- 13 K. L. Ding, A. Gulec, A. M. Johnson, N. M. Schweitzer, G. D. Stucky, L. D. Marks and P. C. Stair, Science, 2015, 350, 189-192.
- 14 B. Shan, N. Kapur, J. Hyun, L. G. Wang, J. B. Nicholas and K. Cho, J. Phys. Chem. C, 2009, 113, 710-715.
- 15 J. L. Liu, F. R. Lucci, M. Yang, S. Lee, M. D. Marcinkowski, A. J. Therrien, C. T. Williams, E. C. H. Sykes and M. Flytzani-Stephanopoulos, J. Am. Chem. Soc., 2016, 138, 6396-6399.
- 16 Y. Hsieh, Y. Zhang, D. Su, V. Volkov, R. Si, L. J. Wu, Y. M. Zhu, W. An, P. Liu, P. He, S. Y. Ye, R. R. Adzic and J. X. Wang, Nat. Commun., 2013, 4, 2466.
- 17 P. P. Fang, S. Duan, X. D. Lin, J. R. Anema, J. F. Li, O. Buriez, Y. Ding, F. R. Fan, D. Y. Wu, B. Ren, Z. L. Wang, C. Amatore and Z. Q. Tian, Chem. Sci., 2011, 2, 531-539.
- 18 G. N. Vayssilov, Y. Lykhach, A. Migani, T. Staudt, G. P. Petrova, N. Tsud, T. Skála, A. Bruix, F. Illas,

- K. C. Prince, V. Ma-tolín, K. M. Neyman and J. Libuda, Nat. Mater., 2011, 10, 310-315.
- 19 L. Q. Liu, F. Zhou, L. G. Wang, X. J. Qi, F. Shi and Y. Q. Deng, J. Catal., 2010, 274, 1-10.
- 20 J. Ke, W. Zhu, Y. Y. Jiang, R. Si, Y. J. Wang, S. C. Li, C. H. Jin, H. C. Liu, W. G. Song, C. H. Yan and Y. W. Zhang, ACS Catal., 2015, 5, 5164-5173.
- 21 K. An, S. Alayoglu, N. Musselwhite, S. Plamthottam, G. Melaet, A. E. Lindeman and G. A. Somorjai, J. Am. Chem. Soc., 2013, 135, 16689-16696.
- 22 F. Wang, W. Li, X. L. Feng, D. P. Liu and Y. Zhang, Chem. Sci., 2016, 7, 1867-1873.
- 23 B. T. Qiao, A. Q. Wang, X. F. Yang, L. F. Allard, Z. Jiang, Y. T. Cui, J. Y. Liu, J. Li and T. Zhang, Nat. Chem., 2011, 3,
- 24 A. Holmgren, B. Andersson and D. Duprez, Appl. Catal., B, 1999, 22, 215-230.
- 25 M. Boaro, F. Giordano, S. Recchia, V. Dal Santo, M. Giona and A. Trovarelli, Appl. Catal., B, 2004, 52, 225-237.
- 26 C. Li, Y. Sakata, T. Arai, K. Domen, K. Maruya and T. J. Onishi, J. Chem. Soc., Faraday Trans. 1, 1989, 85, 929-
- 27 M. Cargnello, V. V. T. Doan-Nguyen, T. R. Gordon, R. E. Diaz, E. A. Stach, R. J. Gorte, P. Fornasiero and C. B. Murray, Science, 2013, 341, 771-773.
- 28 X. Liu, Q. Q. Zhu, Y. Lang, K. Cao, S. Q. Chu, B. Shan and R. Chen, Angew. Chem., Int. Ed., 2017, 56, 1648-1652.
- 29 V. J. Cybulskis, J. Wang, J. H. Pazmiño, F. H. Ribeiro and W. Nicholas Delgass, J. Catal., 2016, 339, 163-172.
- 30 M. J. Kale and P. Christopher, ACS Catal., 2016, 6, 5599-5609.
- 31 L. DeRita, S. Dai, K. Lopez-Zepeda, N. Pham, G. W. Graham, X. Q. Pan and P. Christopher, J. Am. Chem. Soc., 2017, 139, 14150-14165.
- 32 K. Bhattacharyya and C. Majumder, Chem. Phys. Lett., 2007, 446, 374-379.
- 33 C. R. Yin, F. R. Negreiros, G. Barcaro, A. Beniya, L. Sementa, E. C. Tyo, S. Bartling, K. Meiwes-Broer, S. Seifert, H. Hirata, N. Isomura, S. Nigam, C. Majumder, Y. Watanabe, A. Fortunelli and S. Vajda, J. Mater. Chem. A, 2017, 5, 4923-4931.
- 34 Z. Z. Chen, X. Liu, K. Cho, R. Chen and B. Shan, ACS Catal., 2015, 5, 4913-4926.
- 35 S. C. Ammal and A. Heyden, J. Phys. Chem. C, 2011, 115, 19246-19259.