**"** 

# 维化学报

## **Chinese Journal of Catalysis**

主编 林励吾

2013

Editor-in-Chief LIN Liwu

Vol. 34 No. 7





中国化学会催化学会会刊

Transaction of the Catalysis Society of China

#### In This Issue



**對面**: 张静等利用石墨型氮化碳(C<sub>3</sub>N<sub>4</sub>)和氨硼烷(NH<sub>3</sub>BH<sub>3</sub>, AB)球磨制备了 AB-C<sub>3</sub>N<sub>4</sub> 体系. 由于氮化碳的加入, 氨硼烷的放氢反应温度明显降低, 放氢诱导期缩短, 同时抑制了副产物无机苯的生成. 该材料有望作为储氢材料为氢能汽车提供氢源. 见本期第1303-1311页.

Cover: By ball milling carbon nitride  $(C_3N_4)$  and ammonia borane  $(NH_3BH_3, AB)$ , the  $AB-C_3N_4$  system was successfully synthesized, as described by Zhang and coworkers in the Article on pages 1303–1311. The dehydrogenation temperature of  $C_3N_4$ -modified ammonia borane was reduced to lower region with depression of byproduct and shorter induction period compared with pristine ammonia borane. Thus, it is expected that the  $C_3N_4$ -modified ammonia borane can be used as a hydrogen storage material for the hydrogen fuel car in the future.

#### About the Journal

Chinese Journal of Catalysis is an international journal published monthly by Chinese Chemical Society, Dalian Institute of Chemical Physics, Chinese Academy of Sciences, and Elsevier. The journal publishes original, rigorous, and scholarly contributions in the fields of heterogeneous and homogeneous catalysis in English or in both English and Chinese. The scope of the journal includes:

- New trends in catalysis for applications in energy production, environmental protection, and production of new materials, petroleum chemicals, and fine chemicals;
- Scientific foundation for the preparation and activation of catalysts of commercial interest or their representative models;
- Spectroscopic methods for structural characterization, especially methods for in situ characterization;
- New theoretical methods of potential practical interest and impact in the science and applications of catalysis and catalytic reaction;
- Relationship between homogeneous and heterogeneous catalysis;
- Theoretical studies on the structure and reactivity of catalysts.
- The journal also accepts contributions dealing with photo-catalysis, bio-catalysis, and surface science and chemical kinetics issues related to catalysis.

#### **Types of Contributions**

- Reviews deal with topics of current interest in the areas covered by this journal. Reviews are surveys, with entire, systematic, and important information, of recent progress in important topics of catalysis. Rather than an assemblage of detailed information or a complete literature survey, a critically selected treatment of the material is desired. Unsolved problems and possible developments should also be discussed. Authors should have published articles in the field. Reviews should have more than 80 references.
- *Communications* rapidly report studies with significant innovation and major academic value. They are limited to four Journal pages. After publication, their full-text papers can also be submitted to this or other journals.
- Articles are original full-text reports on innovative, systematic and completed research on catalysis.
- Highlight Comments describe and comment on very important new results in the original research of a third person with a view to highlight their significance. The results should be presented clearly but concisely without the comprehensive details required of an original article. Highlight comment should not be more than 2–3 Journal pages (approximately 9000 characters) in length, and should be appropriately organized by the author. Chemical formulae, figures, and schemes should be restricted to important examples. The number of references should be restricted to about 15.
- Academic Arguments can discuss, express a different opinion or query the idea, concept, data, data processing method, characterization method, computational method, or the conclusion of published articles. The objective of an academic argument should be to enliven the academic atmosphere.

#### **Impact Factor**

2012 SCI Impact Factor: 1.304 2012 SCI 5-Year Impact Factor: 1.011 2011 ISTIC Impact Factor: 1.288

#### **Abstracting and Indexing**

Abstract Journals (VINITI)
Cambridge Scientific Abstracts (CIG)
Catalysts & Catalysed Reactions (RSC)
Current Contents/Engineering, Computing
and Technology (Thomson ISI)
Chemical Abstract Service/SciFinder
(CAS)

Chemistry Citation Index (Thomson ISI)

Japan Information Center of Science and Technology

Journal Citation Reports/Science Edition (Thomson ISI)

Science Citation Index Expanded (Thomson ISI)

SCOPUS (Elsevier)

Web of Science (Thomson ISI)



## 催化学报

(CUIHUA XUEBAO)

#### CHINESE JOURNAL OF CATALYSIS

月刊 SCI 收录 2013 年 7 月 第 34 卷 第 7 期



### 目 次

#### 

#### 1277 (英/中)

纳米银催化剂上CO氧化反应研究进展 张晓东, 曲振平, 于芳丽, 王奕

#### 研究快讯

#### 1291 (英)

通过调变碳纳米管大π体系与含氧官能团的共轭作用以改变 碳纳米管的催化性能

杜钰珏, 李振华, 范康年

#### 1297 (英)

表面修饰的钒氧化物纳米带上甲苯选择氧化反应 李轩,叶霜,赵建波,李磊,彭路明,丁维平

#### 研究论文

#### 1303 (英/中/封面文章)

石墨型氮化碳对氨硼烷放氢性能的影响 张静,何腾,刘彬,柳林,赵泽伦,胡大强,鞠晓花,吴国涛, 陈萍

#### 1312 (英/中)

铁基费托合成催化剂相变调控及反应性能 高芳芳, 王洪, 青明, 杨勇, 李永旺

#### 1326 (英)

镍(II)配合物官能化的MCM-41催化分子氧环氧化苯乙烯 杨刚, 陈星, 王小丽, 邢卫红, 徐南平

#### 1333 (英)

苯甲酰二茂铁修饰碳纳米管糊电极电催化测量蛋氨酸浓度 Hadi BEITOLLAHI, Alireza MOHADESI, Farzaneh GHORBANI, Hassan KARIMI MALEH, Mehdi BAGHAYERI, Rahman HOSSEINZADEH

#### 1339 (英/中)

有机溶剂/缓冲液两相体系中全细胞催化拆分环氧氯丙烷 邹树平, 颜海蔚, 胡忠策, 郑裕国

#### 1348 (英/中)

SAPO-34晶粒形貌对甲醇转化制低碳烯烃反应的影响 吴磊, 刘子玉, 夏林, 丘明煌, 刘旭, 朱浩佳, 孙予罕

#### 1357 (英)

Co取代的六铁酸盐在N<sub>2</sub>O催化分解反应中的应用 Barkat Ul-AIN, Safeer AHMED, 黄廷强

#### 1363 (英/中)

双功能钛硅分子筛的合成、表征及催化性能 李颜, 雷骞, 张小明, 索继栓

#### 1373 (英)

钌(II)-吡啶基NNN配合物催化酮的室温(不对称)氢转移反应 杜旺明, 王清福, 余正坤

#### 1378 (英)

可回收Fe<sub>3</sub>O<sub>4</sub>@SiO<sub>2</sub>-Ag磁性纳米微球对染料污染物的快速脱色处理

孙丽娟, 何疆, 安松松, 张军伟, 郑金敏, 任栋

#### 1386 (英)

 $Au/FeO_x$ -羟基磷灰石催化CO氧化反应中羟基磷灰石和 $FeO_x$ 的作用

赵昆峰, 乔波涛, 张彦杰, 王军虎

#### 1395 (英/中)

介孔炭的孔结构对其负载的Ru基氨合成催化剂性能的影响 周亚萍,蓝国钧,周斌,姜维,韩文锋,刘化章,李瑛

#### 1402 (英)

用Lewis酸或Brönsted酸催化剂和N<sub>2</sub>汽提法使木糖脱水为糠醛

Iker AGIRREZABAL-TELLERIA,

Cristina GARCÍA-SANCHO, Pedro MAIRELES-TORRES, Pedro Luis ARIAS

#### 1407 (英/中)

Ni基催化剂催化燃油重整耦合选择性催化还原NO<sub>x</sub>反应 赵娇娇, 余运波, 韩雪, 贺泓

#### 1418 (英/中)

钯氮共掺杂TiO<sub>2</sub>的制备与表征及其可见光催化活性 于新变, 王岩, 孟祥江, 杨建军

#### 1429 (英)

多级孔ZSM-5分子筛: 丰富的外表面酸中心和良好的二甲苯 异构化催化性能

周健, 刘志成, 李丽媛, 王仰东, 高焕新, 杨为民, 谢在库, 唐颐

#### 1434 (英/中)

氧气气氛下Pt<sub>3</sub>Ni(111)表面结构变化的从头算原子热力学研究

孙大鹏, 赵永慧, 苏海燕, 李微雪

#### 1443 (英)

MgO纳米晶负载镍催化剂用于甲烷蒸汽重整反应 Mahmood ANDACHE, Mehran REZAEI, Mansour KAZEMI MOGHADAM

#### 1449 (英/中)

Ni-Ti-O混合氧化物的制备、表征及其富氧选择性催化还原 NO

袁德玲, 李新勇, 肇启东

#### 1456 (英)

非对称N-(亚水杨基)二亚乙基三胺铜(II)配合物的合成、结构分析和催化活性评价

Hassan HOSSEINI-MONFARED, Sohaila ALAVI, Milosz SICZEK

#### 1462 (英)

镍基催化剂上稻草水蒸气重整制富氢合成气 李庆远,季生福,胡金勇,蒋赛

#### 相关信息

1469 作者索引

英文全文电子版(国际版)由Elsevier出版社在ScienceDirect上出版

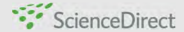
http://www.sciencedirect.com/science/journal/18722067

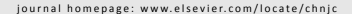
http://www.elsevier.com/locate/chnjc

http://www.chxb.cn



#### available at www.sciencedirect.com







#### Chinese Journal of Catalysis

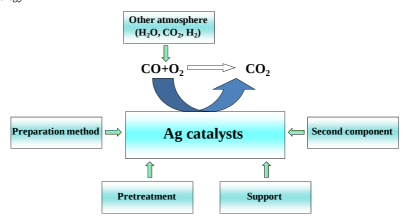
#### **Graphical Contents**

#### **Review**

Chin. J. Catal., 2013, 34: 1277-1290 doi: 10.1016/S1872-2067(12)60610-X

#### Progress in carbon monoxide oxidation over nanosized Ag catalysts

ZHANG Xiaodong, QU Zhenping\*, YU Fangli, WANG Yi Dalian University of Technology



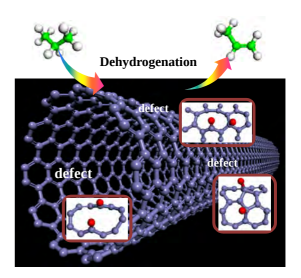
Recent developments of Ag catalysts for CO oxidation mainly focus on the structures of Ag catalysts (particle size, species states, etc.) and their catalytic activities.

#### **Communications**

Chin. J. Catal., 2013, 34: 1291-1296 doi: 10.1016/S1872-2067(12)60627-5

Tuning the catalytic performance of carbon nanotubes by tuning the conjugation between the  $\pi$  orbitals of carbon nanotubes and the active oxygenic functional groups

DU Yujue, LI Zhenhua\*, FAN Kangnian\* Fudan University

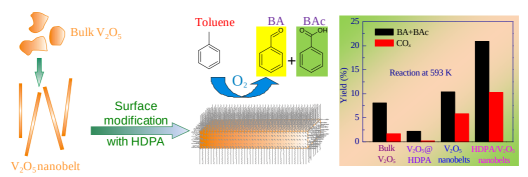


The conjugation between CNT  $\pi$  orbitals and the orbitals of functional groups plays a key role in balancing the activity of the two C–H activation steps in the propane dehydrogenation reaction.

Chin. J. Catal., 2013, 34: 1297–1302 doi: 10.1016/S1872-2067(12)60619-6

#### Selective oxidation of toluene by surface-modified vanadium oxide nanobelts

LI Xuan, YE Shuang, ZHAO Jianbo, LI Lei, PENG Luming, DING Weiping\* Nanjing University



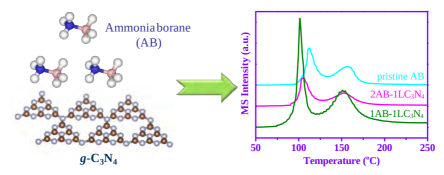
Vanadium oxide nanobelts are more active, but less selective, in toluene oxidation than bulk vanadium oxide is. Significant modification of the selective oxidation of toluene on vanadium oxide nanobelts is achieved by doping with hexadecylphosphonic acid.

#### **Articles**

Chin. J. Catal., 2013, 34: 1303-1311 doi: 10.1016/S1872-2067(12)60566-X

#### Effects of graphitic carbon nitride on the dehydrogenation of ammonia borane

ZHANG Jing, HE Teng\*, LIU Bin, LIU Lin, ZHAO Zelun, HU Daqiang, JU Xiaohua, WU Guotao, CHEN Ping\* Dalian Institute of Chemical Physics, Chinese Academy of Sciences

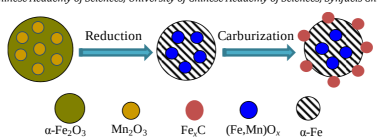


The dehydrogenation temperature of  $C_3N_4$ -modified ammonia borane was reduced to lower region with depression of byproduct and induction period compared with pristine ammonia borane.

Chin. J. Catal., 2013, 34: 1312-1325 doi: 10.1016/S1872-2067(12)60562-2

#### Controlling the phase transformations and performance of iron-based catalysts in the Fischer-Tropsch synthesis

GAO Fangfang, WANG Hong, QING Ming, YANG Yong\*, LI Yongwang
Institute of Coal Chemistry, Chinese Academy of Sciences; University of Chinese Academy of Sciences; Synfuels China Co. Ltd

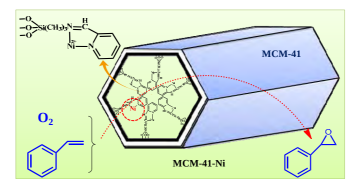


The control of the phase transformations of iron-based catalysts can be achieved by effectively controlling the reduction and carburization conditions.

Chin. J. Catal., 2013, 34: 1326–1332 doi: 10.1016/S1872-2067(12)60568-3

#### Nickel(II) complex anchored on MCM-41 for the epoxidation of styrene by oxygen

YANG Gang\*, CHEN Xing, WANG Xiaoli, XING Weihong, XU Nanping Nanjing University of Technology

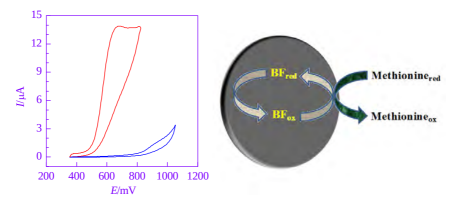


A nickel(II) Schiff base complex functionalized MCM-41 catalyst was used to catalyze epoxidation of styrene using green oxidant molecular oxygen as oxidant with styrene conversion of 95.2% and epoxide se-lectivity of 66.7%.

Chin. J. Catal., 2013, 34: 1333–1338 doi: 10.1016/S1872-2067(12)60582-8

### Electrocatalytic measurement of methionine concentration with a carbon nanotube paste electrode modified with benzoylferrocene

Hadi BEITOLLAHI\*, Alireza MOHADESI, Farzaneh GHORBANI, Hassan KARIMI MALEH, Mehdi BAGHAYERI, Rahman HOSSEINZADEH Graduate University of Advanced Technology, Iran; Payame Noor University, Iran; Hakim Sabzevari University, Iran; University of Mazanda-ran, Iran



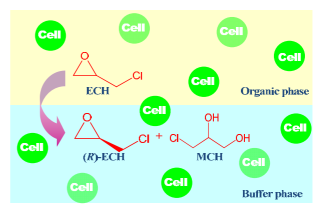
A sensor was fabricated for the measurement of methionine concentration that reduced the oxidation potential of methionine as compared to the unmodified electrode.

Chin. J. Catal., 2013, 34: 1339-1347 doi: 10.1016/S1872-2067(12)60576-2

## Enzymatic resolution of epichlorohydrin catalyzed by whole cells in an organic solvent/buffer biphasic system

ZOU Shuping, YAN Haiwei, HU Zhongce, ZHENG Yuguo\* Zhejiang University of Technology

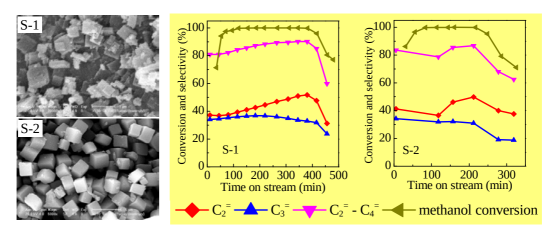
An isooctane/buffer biphasic system is useful for dissolving the substrate epichlorohydrin (ECH) and suppressing its non-enzymatic hydrolysis. This biphasic system could prove useful for improving the concentration of substrate and yield of (*R*)-ECH.



Chin. J. Catal., 2013, 34: 1348-1356 doi: 10.1016/S1872-2067(12)60575-0

#### Effect of SAPO-34 molecular sieve morphology on methanol to olefins performance

WU Lei, LIU Ziyu\*, XIA Lin, QIU Minghuang, LIU Xu, ZHU Haojia, SUN Yuhan\* Shanghai Advanced Research Institute, Chinese Academy of Sciences; University of Chinese Academy of Sciences; Institute of Coal Chemistry, Chinese Academy of Sciences



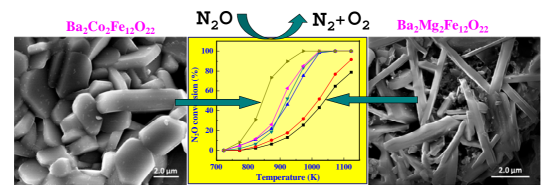
Methanol to olefins performance on SAPO-34 depends on the molecular sieve morphology. The sheet-like SAPO-34 (sample S-1) exhibits a longer lifetime and higher olefin selectivity than the cubic material (sample S-2) because of the shorter diffusion path of the former.

Chin. J. Catal., 2013, 34: 1357-1362 doi: 10.1016/S1872-2067(12)60587-7

#### Catalytic decomposition of N2O on cobalt substituted barium hexaferrites

Barkat UL-AIN, Safeer AHMED, HUANG Yanqiang \*

Dalian Institute of Chemical Physics, Chinese Acadamy of Sciences, China; Quaid-i-Azam University Islamabad, Pakistan



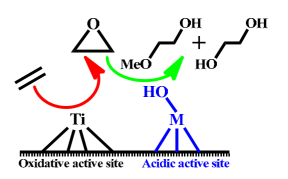
Co substituted barium hexaferrites displayed good activity for  $N_2O$  decomposition. Complete substitution of Mg for Co lowered the complete conversion temperature from 1123 K to 973 K.

Chin. J. Catal., 2013, 34: 1363-1372 doi: 10.1016/S1872-2067(12)60589-0

### Synthesis, characterization, and catalytic performance of bifunctional titanium silicalite-1

LI Hao, LEI Qian, ZHANG Xiaoming \*, SUO Jishuan Chengdu Institute of Organic Chemistry, Chinese Academy of Sciences; Yangtze University

The coincorporation of Ti and a trivalent ion into the MFI-type zeolite was studied. The acidity of TS-1 was enhanced by introducing trivalent ions. M-TS-1 showed higher activity in the selective oxidation of ethylene.



Chin. J. Catal., 2013, 34: 1373-1377 doi: 10.1016/S1872-2067(12)60583-X

## Ru(II) pyridyl-based NNN complex catalysts for (asymmetric) transfer hydrogenation of ketones at room temperature

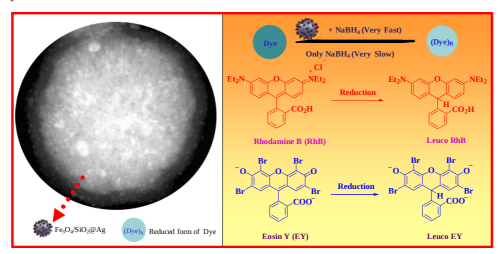
DU Wangming, WANG Qingfu, YU Zhengkun\*
Dalian Institute of Chemical Physics, Chinese Academy of Sciences

Ru(II) complexes bearing pyridyl-based benzimidazolyl-imidazolinyl tridentate NNN ligands exhibited excellent catalytic activity and selectivity in the asymmetric transfer hydrogenation of ketones at room temperature.

Chin. J. Catal., 2013, 34: 1378-1385 doi: 10.1016/S1872-2067(12)60605-6

#### Recyclable Fe<sub>3</sub>O<sub>4</sub>@SiO<sub>2</sub>-Ag magnetic nanospheres for the rapid decolorizing of dye pollutants

SUN Lijuan, HE Jiang\*, AN Songsong, ZHANG Junwei, ZHENG Jinmin, REN Dong Lanzhou University



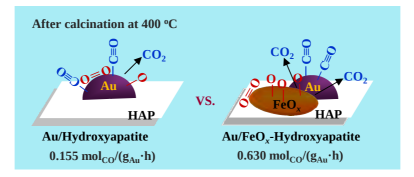
Show STEM image of  $Fe_3O_4@SiO_2$ -Ag nanospheres and their catalytic mechanism for reduction of rhodamine B and eosin Y in the presence of NaBH<sub>4</sub>. The catalytic activity depends on the Ag nanoparticles on the surface of  $Fe_3O_4@SiO_2$  nanoparticles.

Chin. J. Catal., 2013, 34: 1386–1394 doi: 10.1016/S1872-2067(12)60590-7

#### The roles of hydroxyapatite and FeO<sub>x</sub> in a Au/FeO<sub>x</sub>-hydroxyapatite catalyst for CO oxidation

ZHAO Kunfeng, QIAO Botao, ZHANG Yanjie, WANG Junhu\*

Dalian Institute of Chemical Physics, Chinese Academy of Sciences; University of Chinese Academy of Sciences



The hydroxyapatite stabilizes gold and  $FeO_x$  nanoparticles by a strong interaction with these, while  $FeO_x$  promotes the catalytic activity by changing the nature of the reaction mechanism and the intermediates.

Chin. J. Catal., 2013, 34: 1395-1401 doi: 10.1016/S1872-2067(12)60596-8

#### Effect of pore structure of mesoporous carbon on its supported Ru catalysts for ammonia synthesis

ZHOU Yaping, LAN Guojun, ZHOU Bin, JIANG Wei, HAN Wenfeng, LIU Huazhang, LI Ying \* Zhejiang University of Technology

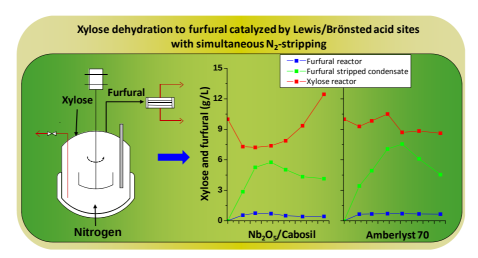


The pore structure of the Ba-Ru-K catalyst supported on mesoporous carbon was controlled by changing the particle size of  $SiO_2$  and the  $SiO_2/C$  ratio. The catalyst activity for ammonia synthesis increased with an increased mesoporous surface area.

Chin. J. Catal., 2013, 34: 1402-1406 doi: 10.1016/S1872-2067(12)60599-3

#### Dehydration of xylose to furfural using Lewis or Brönsted acid catalyst and N2 stripping

Iker AGIRREZABAL-TELLERIA\*, Cristina GARCÍA-SANCHO, Pedro MAIRELES-TORRES, Pedro Luis ARIAS University of the Basque Country, Spain; University of Málaga, Spain



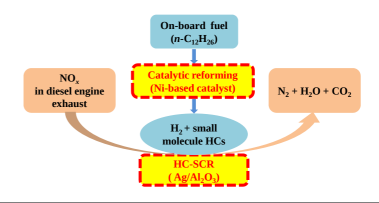
The present work studied the change of the xylose dehydration mechanism and furfural yield using Lewis ( $Nb_2O_5/Cabosil$ ) or Brönsted (Amberlyst 70) acid catalysts in combination with simultaneous discontinuous  $N_2$ -stripping.

Chin. J. Catal., 2013, 34: 1407-1417 doi: 10.1016/S1872-2067(12)60598-1

## Fuel reforming over Ni-based catalysts coupled with selective catalytic reduction of $NO_x$

ZHAO Jiaojiao, YU Yunbo \*, HAN Xue, HE Hong Research Center for Eco-Environmental Sciences, Chinese Academy of Sciences

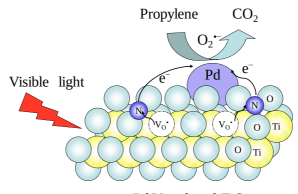
Small molecular weight hydrocarbons and  $H_2$  produced from on-board fuel reforming over Ni-based catalysts were used as reductants in the selective catalytic reduction of  $NO_x$  over  $Ag/Al_2O_3$  catalysts at typical diesel exhaust temperatures.



Chin. J. Catal., 2013, 34: 1418-1428 doi: 10.1016/S1872-2067(12)60597-X

#### Preparation and characterization of Pd/N codoped TiO2 photocatalysts with high visible light photocatalytic activity

YU Xinluan, WANG Yan, MENG Xiangjiang, YANG Jianjun\* Henan University



Pd/N codoped TiO<sub>2</sub>

Pd/N codoped  $TiO_2$  photocatalysts showed high efficiency for the visible light photocatalytic oxidation of propylene. The origin of visible light photocatalytic activity was attributed to both Pt and N dopant elements and the formation of single-electron-trapped oxygen vacancies.

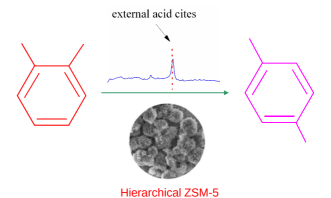
Chin. J. Catal., 2013, 34: 1429-1433 doi: 10.1016/S1872-2067(12)60602-0

## Hierarchical mesoporous ZSM-5 zeolite with increased external surface acid sites and high catalytic performance in o-xylene isomerization

ZHOU Jian, LIU Zhicheng\*, LI Liyuan, WANG Yangdong, GAO Huanxin, YANG Weimin, XIE Zaiku\*, TANG Yi

Shanghai Research Institute of Petrochemical Technology, SINOPEC; Fudan University

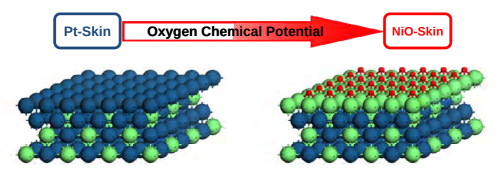
Hierarchical mesoporous ZSM-5 zeolite synthesized by the steam-assisted crystallization method exposes lots of acid sites on the external surface, thereby improving catalytic activity in the isomerization of *o*-xylene.



Chin. J. Catal., 2013, 34: 1434-1442 doi: 10.1016/S1872-2067(12)60604-4

#### An atomistic thermodynamics study of the structural evolution of the Pt<sub>3</sub>Ni(111) surface in an oxygen environment

SUN Dapeng, ZHAO Yonghui, SU Haiyan, LI Weixue\*
Dalian Institute of Chemical Physics, Chinese Academy of Sciences



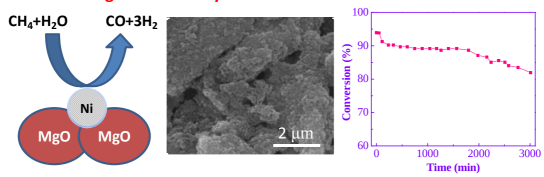
With a gradual increase of oxygen chemical potential, the clean Pt-skin surfaces of Pt<sub>3</sub>Ni(111) transit to the oxygen-chemisorbed Ni-skin surfaces without stable oxygen-chemisorbed PtNi surface alloys formed.

Chin. J. Catal., 2013, 34: 1443-1448 doi: 10.1016/S1872-2067(12)60606-8

#### A nanocrystalline MgO support for Ni catalysts for steam reforming of CH<sub>4</sub>

Mahmood ANDACHE, Mehran REZAEI\*, Mansour KAZEMI MOGHADAM
Islamic Azad University, Iran; University of Kashan, Iran; Iran University of Science and Technology, Iran

#### High stable catalyst with low carbon formation



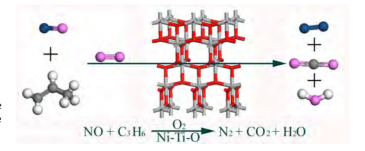
A 7% Ni catalyst supported on mesoporous nanocrystalline MgO showed high catalytic stability and low degree of carbon formation in steam reforming of methane for syngas production.

Chin. J. Catal., 2013, 34: 1449-1455 doi: 10.1016/S1872-2067(12)60614-7

## Preparation and characterization of Ni-Ti-O mixed oxide for selective catalytic reduction of NO under lean-burn conditions

YUAN Deling, LI Xinyong\*, ZHAO Qidong Dalian University of Technology

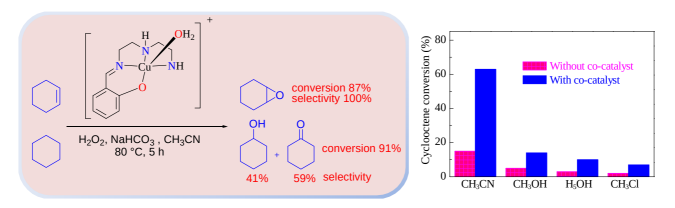
A novel Ni-Ti-O mixed-oxide catalyst was used in the selective catalytic reduction of NO by  $C_3H_6$  under lean-burn conditions; the catalyst demonstrated good activity at 430 °C.



Chin. J. Catal., 2013, 34: 1456-1461 doi: 10.1016/S1872-2067(12)60616-0

## Synthesis, structural analysis and evaluation of the catalytic activity of a non-symmetric N-(salicylidene)diethylenetriamine complex of copper(II)

Hassan HOSSEINI-MONFARED\*, Sohaila ALAVI, Milosz SICZEK University of Zanjan, Iran; University of Wroclaw, Poland

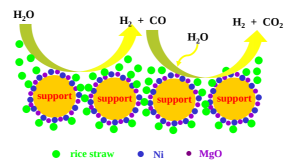


Synthesis and crystal structural analysis of a new copper(II) complex of a non-symmetric Schiff base have been reported. The complex showed high levels of catalytic activity and selectivity towards the oxidation of cycloalkenes and cyclohexane by  $H_2O_2$  and  $NaHCO_3$ .

Chin. J. Catal., 2013, 34: 1462–1468 doi: 10.1016/S1872-2067(12)60618-4

#### Catalytic steam reforming of rice straw biomass to hydrogen-rich syngas over Ni-based catalysts

LI Qingyuan, JI Shengfu\*, HU Jinyong, JIANG Sai Beijing University of Chemical Technology

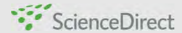


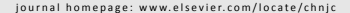
Supported Ni-based catalysts

A series of supported Ni-based catalysts were prepared and used for steam reforming of rice straw biomass to hydrogen-rich syngas. A 1.0% MgO-7.5% Ni/ $\gamma$ -Al<sub>2</sub>O<sub>3</sub> catalyst exhibited the highest catalytic activity of the series.



#### available at www.sciencedirect.com







#### Article

## An atomistic thermodynamics study of the structural evolution of the Pt<sub>3</sub>Ni(111) surface in an oxygen environment

SUN Dapeng, ZHAO Yonghui, SU Haiyan, LI Weixue\*

State Key Laboratory of Catalysis, Dalian Institute of Chemical Physics, Chinese Academy of Sciences, Dalian 116023, Liaoning, China

#### ARTICLE INFO

Article history:
Received 8 January 2013
Accepted 28 April 2013
Published 20 July 2013

Keywords:
Pt<sub>3</sub>Ni
Oxygen
Segregation
Phase diagram
Ab initio calculation

#### ABSTRACT

The structural evolution of the  $Pt_3Ni(111)$  surface under oxidizing conditions was studied by ab initio atomistic thermodynamics. The thermodynamic phase diagram from Ni-rich to Pt-rich conditions with oxygen coverages up to one monolayer was constructed from their 560 possible surface structures. With an increase in the oxygen chemical potential, there were only two types of thermodynamically stable structures, which were a clean Pt-skin surface and a Ni-skin surface with chemisorbed oxygen, regardless of the underlying Pt-rich or Ni-rich conditions. Bimetallic surfaces with chemisorbed oxygen were only metastable. The detail analysis revealed that the structural evolution is determined by the factors of segregation cost, difference between oxygen-metal (Pt and Ni) bonding strength, and oxygen chemical potential.

© 2013, Dalian Institute of Chemical Physics, Chinese Academy of Sciences.

Published by Elsevier B.V. All rights reserved.

#### 1. Introduction

Alloys, especially bimetallic alloy systems comprising Pt and a non-noble transition metal (TM), are widely used as catalysts [1–10] because they have unique activities in various catalytic reactions, such as water-gas-shift reaction [11], oxygen reduction reaction [12–15], and hydrogenation [16,17]. The Pt-Ni alloy has received special attention because it was found that the Pt<sub>3</sub>Ni(111) surface with a Pt-skin surface exhibited a much higher activity for oxygen reduction that was ten times that of Pt(111) and 90 times higher than a commercialized Pt/C catalyst [18]. Also, a "sandwich-like" structure comprising NiO<sub>1-x</sub>/Pt/Ni/Pt(111) has shown the highest activity for CO oxidation [19]. It was further found that the surface structures of the Pt-Ni bimetallic system changed reversibly under the different conditions of an oxidizing or a reducing atmosphere [20,21].

These results indicated that the chemical properties of an alloy surface such as its reactivity, selectivity, and stability

strongly depend on its composition, structure, and chemical state. Under a reactive environment, the adsorption of reactants on the alloy surface can result in surface reconstruction, segregation, and oxidation or reduction, which affect the catalytic mechanism and activity [22]. A systematic and quantitative study of the structural evolution of the alloy surface under a catalytic reaction environment can reveal the important factors that should be considered in the design and improvement of the stability and activity of the catalyst.

This work is a comprehensive study of surface segregation and oxidation under an oxygen environment of the  $Pt_3Ni(111)$  surface by an ab initio atomistic thermodynamic method [23–25]. We discuss the role of the bulk alloy reservoir by considering the effect of a small amount of non-stoichiometry in a  $Pt_3Ni$  bulk alloy. The thermodynamic surface phase diagram of  $Pt_3Ni(111)$  under an oxygen environment was constructed. In addition, we analyzed the influence of the segregation energy, difference in the oxygen binding energy to Pt and Ni, and the

<sup>\*</sup>Corresponding author. Tel: +86-411-84379996; Fax: +86-411-84694447; E-mail: wxli@dicp.ac.cn
This work was supported by the National Natural Science Foundation of China (21173210, 21225315, 21273224, 21103164).
DOI: 10.1016/S1872-2067(12)60604-4 | http://www.sciencedirect.com/science/journal/18722067 | Chin. J. Catal., Vol. 34, No. 7, July 2013

oxygen chemical potential on the structural evolution.

#### 2. Methods

## 2.1. Gibbs free energy of formation of the alloy surface under an oxygen environment

For a given temperature T and oxygen partial pressure  $p_{02}$ , the stability of the alloy surface is described by the Gibbs free energy of formation, which is defined as follows for the Pt-Ni bimetallic system:

$$\gamma = (G_{\rm s} - N_{\rm s} x_{\rm Pt} \mu_{\rm Pt} - N_{\rm s} x_{\rm Ni} \mu_{\rm Ni} - N_{\rm 0} \mu_{\rm 0})/A \tag{1}$$

where  $N_{\rm s}$  and  $N_{\rm 0}$  are the numbers of metal atoms and oxygen atoms in the system, and  $x_{\rm Pt}$  and  $x_{\rm Ni}$  are the mole factions of Pt and Ni ( $x_{\rm Pt}+x_{\rm Ni}=1$ ). A is the normalized surface area, which is equal to the twice of one side surface area of the center-symmetric model as below.  $G_{\rm s}$  is the Gibbs free energy of the alloy system, which is usually approximately equal to the total energy  $E_{\rm s}$  [24].  $\mu_{\rm 0}$  is the oxygen chemical potential at a specified temperature and pressure condition:

$$\mu_0 = 0.5 \times (E_{02} + \Delta \mu_{02}(T, p^{\circ}) + k_{\rm B}T \ln(p_{02}/p^{\circ})) = 0.5 \times E_{02} + \Delta \mu_{0}(T, p_{02})$$
 (2)

where  $k_{\rm B}$  is the Boltzmann constant, and  $E_{\rm O2}$  is the total energy of an isolated O<sub>2</sub> molecule.  $\Delta\mu_{\rm O2}(T,\,p^{\rm o})$  can be obtained from tabulated enthalpy and entropy values at the standard pressure  $p^{\rm o}=0.1$  MPa.  $\mu_{\rm Pt}$  and  $\mu_{\rm Ni}$  are the chemical potentials of Pt and Ni, respectively. For bulk Pt<sub>3</sub>Ni alloy in equilibrium,  $4\mu_{\rm Pt3Ni}=3\mu_{\rm Pt}+\mu_{\rm Ni}$ , where  $\mu_{\rm Pt3Ni}$  is the chemical potential of the Pt<sub>3</sub>Ni alloy, which is approximately equal to the total energy per atom ( $E_{\rm Pt3Ni}$ ) of the bulk Pt<sub>3</sub>Ni alloy. Eq. (1) can be rewritten as

$$\gamma \approx (E_{\rm s} - N_{\rm s} E_{\rm Pt3Ni} - N_{\rm s} \Delta \mu_{\rm Pt-Ni} (x_{\rm Pt} - 0.75) - 0.5 \times N_0 E_{\rm 02} - N_0 \Delta \mu_0 (T, p_{\rm 02}))/A$$
(3)

where  $\Delta\mu_{\text{Pt-Ni}} = \mu_{\text{Pt}} - \mu_{\text{Ni}}$  is the difference between the chemical potentials of Pt and Ni in the alloy system, and 0.75 is the mole faction of Pt in stoichiometric bulk Pt<sub>3</sub>Ni.

With a Pt-rich or Ni-rich condition, the corresponding chemical potential  $\mu_{\text{Pt}}$  or  $\mu_{\text{Ni}}$  in the alloy is higher. However, their values are lower than that of pure bulk Pt or Ni, namely,  $\mu_{\text{Pt}} \leq \mu_{\text{Pt(fcc)}}$  and  $\mu_{\text{Ni}} \leq \mu_{\text{Ni(fcc)}}$ , because otherwise the Pt-Ni alloy will not be stable. Because  $4\mu_{\text{Pt3Ni}} = 3\mu_{\text{Pt}} + \mu_{\text{Ni}}$ ,  $\Delta\mu_{\text{Pt-Ni}}$  can only vary in range

$$\Delta\mu_{\text{Pt-Ni}}$$
 (Ni-rich)  $\leq \Delta\mu_{\text{Pt-Ni}} \leq \Delta\mu_{\text{Pt-Ni}}$  (Pt-rich) (4) where  $\Delta\mu_{\text{Pt-Ni}}$  (Ni-rich) =  $(\mu_{\text{Pt3Ni}} - \mu_{\text{Ni}(\text{fcc})})/0.75$ , and  $\Delta\mu_{\text{Pt-Ni}}$  (Pt-rich) =  $4(\mu_{\text{Pt}(\text{fcc})} - \mu_{\text{Pt3Ni}})$ . By approximating  $\mu_{\text{Pt}(\text{fcc})}$  and  $\mu_{\text{Ni}(\text{fcc})}$  by the total energy of bulk face centered cubic (fcc) Pt and Ni,  $E_{\text{Pt}(\text{fcc})}$  and  $E_{\text{Ni}(\text{fcc})}$ , the lower and upper limits of  $\Delta\mu_{\text{Pt-Ni}}$  are  $\Delta\mu_{\text{Pt-Ni}}$  (Ni-rich) =  $-0.67$  eV and  $\Delta\mu_{\text{Pt-Ni}}$  (Pt-rich) =  $-0.32$  eV, respectively.

Equation (3) indicates that the Gibbs free energy of formation of the alloy surface depends on the relative contents of Pt and Ni, and also on the chemical potential of oxygen. We note that under a high  $\Delta\mu_0$ , the alloy surface can be oxidized to surface oxides and even bulk oxides. For simplicity, we limit the present work to the chemisorption regime with oxygen coverages up to one monolayer (ML) only. These results from oxygen chemisorption as a precursor, however, provide valuable insights for the subsequent oxidation.

#### 2.2. Computational details

Spin-polarized density functional theory (DFT) calculations were performed using the Vienna Ab initio Simulation Package (VASP) [26,27] using the all-electron projected augmented wave (PAW) potentials [28,29] and the Perdew-Wang 91 (PW91) exchange-correlation functional [30]. The wave function was expanded in plane waves with a kinetic cutoff of 400 eV. The Monkhorst-Pack method (containing the  $\Gamma$  point) was used for k-point sampling in the reduced Brillouin zone. For the bulk fcc structures of Pt, Ni, and Pt<sub>3</sub>Ni, a grid of (12×12×12) was used. The optimized lattice constants were  $a_0(Pt) = 0.399$ nm,  $a_0(Ni) = 0.352$  nm, and  $a_0(Pt_3Ni) = 0.389$  nm, which agreed well with a previous study [31]. The alloy surfaces were simulated by using the (1×1) structures of center-symmetric slab models with 7 layers, and the two identical surfaces were separated by a vacuum of 1.5 nm. The middle 3 layers were fixed as the bulk while the other layers were fully relaxed during structure optimization. For the oxygen covered surfaces, oxygen atoms were allowed to adsorb symmetrically on both sides of the unit cell. (6×6×1) Monkhorst-Pack k-point grids were used in the surface structure calculations. For the computation of gaseous oxygen, a (1.45 nm  $\times$  1.50 nm  $\times$  1.56 nm) cell was used. The force convergence for all structural relaxation was set to less than 0.1 eV/nm.

#### 3. Results and discussion

The structure of the bulk Pt<sub>3</sub>Ni alloy is L1<sub>2</sub>, with a stoichoimetric ratio of 3:1 for Pt:Ni in the bulk. For the Pt3Ni(111)-(1×1) surface, each layer contains three Pt atoms and one Ni atom. In order to study the effect of nonstoichiometry, the ratios of Pt:Ni in the outer two layers (for each side of the symmetric cell) were allowed to change gradually from 8:0 (Pt-rich condition) to 0:8 (Ni-rich condition), while the bulk phase (the middle 3 layers) was fixed at 3:1. Taking into account the relative positions of the bulk atoms, there are 35 possible substrate configurations. Based on these clean surface structures, oxygen adsorption was considered by placing from zero up to four 0 atoms on each side of the unit cell onto the fcc sites, which corresponded to oxygen coverages  $\theta_0$  = 0.00, 0.25, 0.50, 0.75, 1.00 ML, respectively. A combination of  $C_4^0 + C_4^1 + C_4^2 + C_4^3 + C_4^4 = 16$ possible 0 adsorption configurations for the surface grid was obtained ( $\theta_0 = 0$  ML was considered one type of adsorption). From these, we constructed 560 (35×16) possible structures. They were some equivalent surface structures, but to make it easier to manipulate in the batch program, we kept all the configurations. We calculated the total energy of each configuration after structure optimization by DFT and used this in Eq. (3) to get the relative stability of the configurations for different oxygen chemical potentials under Ni-rich (Fig. 1) and Pt-rich (Fig. 2) conditions. A higher coverage of oxygen gave a steeper slope of the lines of Gibbs free energy of formation, and a higher sensitivity to the oxygen environment. For a given oxygen chemical potential, the surface configuration with the lowest Gibbs free energy of formation (the lowest line in Fig. 1 and Fig. 2) is the thermodynamically most favorable structure.

#### 3.1. The phase diagram of the Pt<sub>3</sub>Ni(111) surface

Figure 1 shows the Gibbs free energy of formation of the Pt<sub>3</sub>Ni(111) surface under Ni-rich conditions versus the oxygen chemical potential  $\Delta\mu_0$ . At the very low oxygen chemical potential conditions of  $\Delta\mu_0$  < -1.68 eV that pertain to a high temperature and low pressure (see the upper axes of Fig. 1), the most stable surface structure is a clean Pt-skin structure with 100% Pt in the first layer and 100% Ni in the second layer. As illustrated in Fig. 1(a), the outer two layers have a Pt-Ni ratio of 4:4, which is lower than the bulk value (3:1), manifesting a Ni-rich condition. With increasing oxygen chemical potential, the dissociative adsorption of oxygen on the surface becomes exothermic, and the coverage of oxygen increases gradually. When  $\Delta\mu_0 > -1.68$  eV, a structure of  $\theta_0 = 0.50$  ML with 100% Ni in the first layer (Ni-skin) and Pt:Ni = 3:1 in the second layer is the most stable (Fig. 1(b)). In this structure, the Pt-Ni ratio in the outer two layers is 3:5, which is also a Ni-rich condition. Under the condition of  $\Delta\mu_0 > -1.58$  eV, the most stable structure is shown in Fig. 1(c), with an oxygen coverage of  $\theta_0$  = 0.75 ML. At even higher oxygen chemical potentials of  $\Delta\mu_0 > -0.62$  eV, the coverage of oxygen reaches 1 ML, as shown in Fig. 1(d). During the evolution of the surface from Fig. 1(b) to 1(d), the configurations in the outer two layers are the same as in Fig. 1(b), and the Pt-Ni ratio is always 3:5 under the Ni-rich condition.

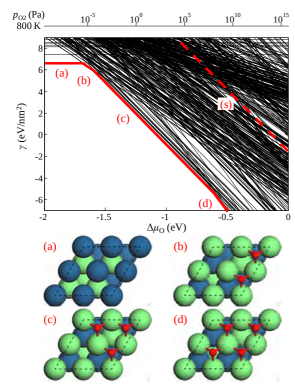


Fig. 1. Surface free energy of  $Pt_3Ni(111)$  versus oxygen chemical potential  $\Delta\mu_0$  under Ni-rich condition. Each line corresponds to one possible surface configuration. The red segments (a, b, c, and d) represent the most stable structure for the range of  $\Delta\mu_0$ . The corresponding structures (top view) are shown at the bottom. The red dashed line (s) corresponds to the stoichiometric  $Pt_3Ni(111)$  surface with  $\theta_0=0.75$  ML, as shown in Fig. 2(s). The blue, green, and red balls are, respectively, Pt, Ni, and O atoms. The correlation between  $\Delta\mu_0$  and oxygen partial pressure at 800 K is indicated at the top.

It should be emphasized that we considered all possible compositions and configurations with different oxygen coverages. With increasing oxygen chemical potential, there are only two types of stable structures, namely, the clean Pt-skin surface and the Ni-skin surfaces with  $\theta_0 \geq 0.50$  ML. There were no Pt-Ni bimetallic surfaces (e.g. the stoichiometric Pt<sub>3</sub>Ni surface in Fig. 1(s), with the schematic structure shown in Fig. 2(s)) nor structures with oxygen coverages less than 0.25 ML that exists as stable structures, that is, these are only metastable. The reason is the interplay between the segregation energies and oxygen-metal binding energies of Pt and Ni, which will be discussed later.

The phase diagram of  $Pt_3Ni(111)$  under a Pt-rich condition is presented in Fig. 2. When  $\Delta\mu_0 < -1.48$  eV, the most stable surface is composed of 100% Pt in the first layer (Pt-skin) and Pt:Ni = 3:1 in the second layer without any adsorbed 0 on the surface (Fig. 2(a)). With increasing oxygen chemical potential, the dissociative adsorption of oxygen on the surface starts to be exothermic, and the coverage of oxygen increases correspondingly. Our calculations show that when  $\Delta\mu_0 > -1.48$  eV, a structure of  $\theta_0 = 0.75$  ML with 100% Ni in the first layer (Ni-skin) and 100% Pt in the second layer is the most stable (Fig. 2(b)). At the even higher oxygen chemical potentials of  $\Delta\mu_0 > -0.68$  eV, the coverage of oxygen reaches 1 ML, as shown in Fig. 2(c), where the configurations in the outer two layers of this Ni-skin surface are the same as Fig. 2(b).

Similarly to the Ni-rich condition, with increasing oxygen chemical potential, there are only two types of stable surface structures, namely, the Pt-skin surface without adsorbed O and the Ni-skin surfaces with  $\theta_0 \ge 0.50$  ML. Pt-Ni bimetallic surfaces (e.g. the stoichiometric Pt<sub>3</sub>Ni surface alloy in Fig. 2(s)) and

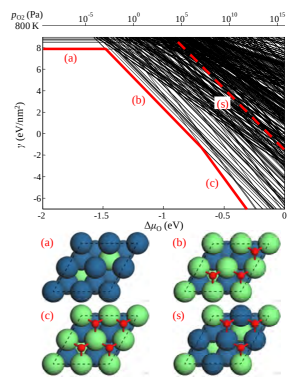


Fig. 2. Surface free energy of  $Pt_3Ni(111)$  versus oxygen chemical potential  $\Delta\mu_0$  under a Pt-rich condition.

structures with oxygen coverage of 0.25, and 0.50 ML are only metastable.

#### 3.2. Segregation energy and oxygen-metal binding

To understand the above results, we define three quantities as follows:

$$E_{\text{Seg}} = E_{\text{s}}^{\text{dean}} - E_{\text{s}|\text{Pg:NN}|}^{\text{dean}} - N_{\text{S}} \Delta \mu_{\text{Pt-Ni}} (x_{\text{Pt}} - 0.75)$$
(5)  
$$E_{\text{b}} = (E_{\text{s}} - E_{\text{s}}^{\text{dean}} - 0.5 \times N_0 E_{02}) / N_0$$
(6)

$$E_{\rm b} = (E_{\rm s} - E_{\rm s}^{\rm clean} - 0.5 \times N_0 E_{02})/N_0 \tag{6}$$

$$\gamma_{\text{Pt3Ni}} = (E_{\text{s(Pt3Ni)}}^{\text{clean}} - N_{\text{s}}E_{\text{Pt3Ni}})/A \tag{7}$$

where  $E_{\text{seg}}$  is the segregation energy, which is the energy cost of the particular surface configuration (without adsorbates) with respect to the stoichiometric surface, and  $E_{
m s}^{
m clean}$  and  $E_{
m s}^{
m clean}$  are the corresponding total energies.  $E_b$  is the average adsorption energy per O atom on the particular surface. YPt3Ni is the surface energy of the stoichiometric Pt<sub>3</sub>Ni(111) surface. Equation (3) can be rewritten as

$$\gamma \approx \gamma_{\text{Pt3Ni}} + (E_{\text{seg}} + N_0 E_b - N_0 \Delta \mu_0(T, p_{02}))/A$$
 (8)

Equation (8) indicates that the Gibbs free energy of formation of a particular surface is determined by the segregation energy  $E_{\text{seg}}$ , average O adsorption energy  $E_{\text{b}}$ , and oxygen chemical potential in addition to the surface energy of stoichiometric Pt<sub>3</sub>Ni(111). The dependence of  $E_{seg}$  and  $E_b$  on composition and configuration will be discussed below.

We first calculated  $E_{\text{seg}}$  for all the structures considered. Figure 3 is the calculated  $E_{\text{seg}}$  (per unit surface area) as a function of the Pt concentration in the topmost layer ( $x_{Pt}$ ). At a given  $x_{Pt}$ , only the result with the least segregation energy is plotted. Regardless of the structures present under Pt-rich or Ni-rich conditions, the segregation energies decrease linearly with xPt. When  $x_{Pt}$  is more than 75%,  $E_{seg}$  even becomes negative (exothermic). This implies that in the absence of oxygen adsorption, Pt atoms tend to segregate to the surface. This is understandable because the surface energy of Pt(111) is lower than that of Ni(111) [32]. As seen in Fig. 3, when the surface configuration changes from the Ni-skin ( $x_{Pt}$  = 0%, containing 4 Ni atoms in the top layer of the unit surface area) to the Pt-skin ( $x_{Pt} = 100\%$ , containing 4 Pt atoms in the top layer of the unit surface area), the segregation energy decreases by 2.54 (Pt-rich) and 1.97 (Ni-rich) eV. That is, there is an average energy cost of 0.56 eV/atom when a surface Pt atom is substituted by a bulk Ni

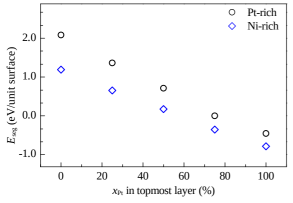


Fig. 3. Segregation energy per (1×1) surface as a function of Pt concentration in the topmost layer  $(x_{Pt})$  for all the clean surface configurations. Only the lowest value at each  $x_{Pt}$  is shown.

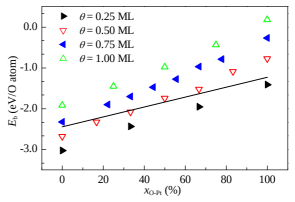


Fig. 4. Average adsorption energy of oxygen as a function of the percentage of O-Pt bonds over all O-metal bonds,  $x_{0-Pt} = N(O-Pt)/(N(O-Pt)$ + N(O-Ni)), for different coverages of oxygen. The line is a linear interpolation between the average adsorption energy of oxygen (0.25 ML) on Ni(111) and Pt(111) surfaces and is given as a guide to the eye.

atom.

Using Eq. (6), we calculated the average oxygen dissociative adsorption energy  $(E_b)$  and studied its dependence on surface composition and oxygen coverage. For a given oxygen coverage, the calculated E<sub>b</sub> was very sensitive to the percentage of O-Pt bonds of all the O-metal bonds ( $x_{O-Pt}$ ). As shown in Fig. 4,  $E_{\rm b}$  increases with  $x_{\rm 0-Pt}$ . This means that the oxygen-metal bond strength decreases with more O-Pt bond, which is understandable because the O-Ni bond strength is higher than that of the O-Pt bond, as seen from the calculated oxygen average adsorption energy of -2.44 and -1.23 eV on Ni(111) and Pt(111) at  $\theta_0$  = 0.25 ML. It was found further that at a given  $x_{0-Pt}$ , a higher coverage of oxygen gave a higher average adsorption energy, which was due to the increase of lateral repulsion between adsorbed oxygen. From the above results, the difference in the O-metal bond strength between O-Ni and O-Pt on Pt<sub>3</sub>Ni and their dependence on oxygen coverage were extracted. For  $\theta_0$  = 0.25 ML, when  $x_{0-Pt}$  increases from 0% (three 0-Ni bonds) to 100% (three O-Pt bonds), Eb decreases by 1.62 eV. Thus an O-Ni bond strength is on average 0.54 eV stronger than that of the O-Pt bond. With the increase of oxygen coverage to 0.50, 0.75, and 1.00 ML, the difference in the bond strength increases to 0.64, 0.69, and 0.70 eV, respectively.

#### The trends of oxygen-induced segregation

The above analysis indicates that once a Ni atom segregates onto the surface by substituting for a Pt atom, the energy of the system rises about 0.56 eV. As a consequence, at a low oxygen chemical potential without dissociative oxygen adsorption, the segregation of the more noble metal is preferred in order to minimize the total Gibbs free energy, which finally gives a clean Pt-skin surface. While increasing oxygen chemical potential in the range of  $\Delta\mu_0 > -1.68$  eV (Ni-rich, Fig. 1) or  $\Delta\mu_0 > -1.48$  eV (Pt-rich, Fig. 2), oxygen from the gas environment starts to adsorb dissociatively on the surface. From Fig. 4, we note that when  $\theta_0$  = 0.25 ML, the difference between the O-Ni bond strength and O-Pt bond strength is 0.54 eV, which would not compensate for the cost of the segregation energy of Ni atom

from the bulk to the surface (about 0.56 eV). However, when  $\theta_0 \geq 0.50$  ML, the difference between the 0–Ni bond strength and 0–Pt bond strength increases to 0.64, 0.69, and 0.70 eV respectively, which are all large enough to compensate the cost of the segregation energy. On the other hand, when  $\theta_0 \geq 0.50$  ML, all metals on the top layer will bind to at least one 0 atom.

Thus, once the thermodynamic condition for oxygen dissociative adsorption is reached, oxygen-induced Ni segregation will occur to maximize the O–Ni bonds, so that a Ni-skin surface with  $\theta_0 \geq 0.50$  ML chemisorbed O will be formed, if there is no kinetic restrictions. To better see why the Pt-Ni bimetallic surfaces are only metastable, we consider a stoichiometric Pt<sub>3</sub>Ni(111) surface with a oxygen coverage of 0.75 ML. Although its segregation energy is lower than that of the Ni-skin surface, the weaker O–Pt bond leads to a higher Gibbs free energy of formation (shown by the red dashed lines (s) in Figs. 1 and 2) compared to the Ni-skin surface with the same oxygen coverage.

For oxygen-induced segregation to cause the change from the Pt-skin to Ni-skin, the corresponding oxygen chemical potential should be  $\Delta\mu_0$  = –1.68 eV (Ni-rich) and  $\Delta\mu_0$  = –1.48 eV (Pt-rich). As indicated in the top panel of Figs. 1 and 2, these correspond to oxygen partial pressures of 3.5  $\times$  10<sup>-6</sup> Pa (Ni-rich) and 1.2  $\times$  10<sup>-3</sup> Pa (Pt-rich) at 800 K, respectively. At 300 K, they are equivalent to 5.6  $\times$  10<sup>-43</sup> Pa (Ni-rich) and 2.9  $\times$  10<sup>-36</sup> Pa (Pt-rich), which are far less than possible at ultra-high vacuum (UHV) conditions. This means that oxygen-induced Ni segregation to form the Ni-skin surface occurs easily at elevated temperatures. However, with possible kinetic hindrance at low temperatures, metastable structures can however be formed.

There is no systematic experimental study of  $Pt_3Ni(111)$  under oxidizing conditions so far. However, there was an experimental study of the Pt(111) surface deposited with 1.3 ML Ni [21]. It was found that at UHV and 800 K (corresponding to  $\Delta\mu_0 < -1.8$  eV), all the Ni atoms diffused into the subsurface region to leave a Pt-skin surface. On increasing the pressure to

 $p_{02}=5.2\times10^{-5}$  Pa ( $\Delta\mu_0=-1.6$  eV), most Ni atoms segregated onto the surface and a NiO film was formed. Considering the errors in the DFT calculations from the exchange-correlation functional and the difference between the Pt<sub>3</sub>Ni(111) alloy and the 1.3 ML Ni/Pt(111) system, the agreement between theory and experiment is satisfactory. The experimental study also found that when the temperature was decreased to 700 K but with the same oxygen pressure of  $p_{02}=5.2\times10^{-5}$  Pa, there was less Ni segregation and oxidation. Because the corresponding oxygen chemical potential (equivalent to  $\Delta\mu_0=-1.4$  eV) was even higher than that at the higher temperature, this means that kinetic hindrance to Ni segregation on Pt<sub>3</sub>Ni(111) was present at 700 K.

#### 4. Conclusions

A systematic ab initio atomistic thermodynamics study on the structural evolution of the  $Pt_3Ni(111)$  surface with oxygen adsorption up to 1 ML coverage was performed. With increasing oxygen chemical potential, there are only two types of thermodynamically stable structures, namely, the clean Pt-skin surface and Ni-skin surfaces with oxygen coverage higher than 0.50 ML. The interplay between the segregation energy and the difference in oxygen bonding strength between Pt and Ni plays an essential role in the structural evolution in the presence of oxygen. Surface alloys are metastable, but they can be formed when there is kinetic hindrance to structural change.

#### References

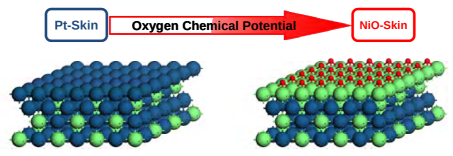
- [1] Shao Y Y, Yin G P, Gao Y Z. J Power Sources, 2007, 171: 558
- [2] Ferrando R, Jellinek J, Johnston R L. Chem Rev, 2008, 108: 845
- [3] Antolini E, Lopes T, Gonzalez E R. J Alloys Compd, 2008, 461: 253
- [4] Peng Z M, Yang H. Nano Today, 2009, 4: 143
- [5] Wang D S, Li Y D. Adv Mater, 2011, 23: 1044
- [6] Fu Q, Li W X, Yao Y X, Liu H Y, Su H Y, Ma D, Gu X K, Chen L M, Wang Z, Zhang H, Wang B, Bao X H. Science, 2010, 328: 1141

#### **Graphical Abstract**

Chin. J. Catal., 2013, 34: 1434–1442 doi: 10.1016/S1872-2067(12)60604-4

An atomistic thermodynamics study of the structural evolution of the  $Pt_3Ni(111)$  surface in an oxygen environment

SUN Dapeng, ZHAO Yonghui, SU Haiyan, LI Weixue\* Dalian Institute of Chemical Physics, Chinese Academy of Sciences



With a gradual increase of oxygen chemical potential, the clean Pt-skin surfaces of  $Pt_3Ni(111)$  transit to the oxygen-chemisorbed Ni-skin surfaces without stable oxygen-chemisorbed PtNi surface alloys formed.

- [7] Sun D P, Gu X K, Ouyang R H, Su H Y, Fu Q, Bao X H, Li W X. J Phys Chem C, 2012, 116: 7491
- [8] Su H Y, Bao X H, Li W X. J Chem Phys, 2008, 128: 194707
- [9] Ma X F, Deng H Q, Yang M M, Li W X. J Chem Phys, 2008, 129: 244711
- [10] Ma X F, Su H Y, Deng H Q, Li W X. Catal Today, 2011, 160: 228
- [11] Knudsen J, Nilekar A U, Vang R T, Schnadt J, Kunkes E L, Dumesic J A, Mavrikakis M, Besenbacher F. J Am Chem Soc, 2007, 129: 6485
- [12] Marković N M, Ross P N Jr. Surf Sci Rep, 2002, 45: 117
- [13] Stamenkovic V, Mun B S, Mayrhofer K J J, Ross P N, Markovic N M, Rossmeisl J, Greeley J, Norskov J K. Angew Chem, Int Ed, 2006, 45: 2897
- [14] Greeley J, Jaramillo T F, Bonde J, Chorkendorff I, Norskov J K. Nat Mater, 2006, 5: 909
- [15] Chen S, Ferreira P J, Sheng W C, Yabuuchi N, Allard L F, Shao-Horn Y. J Am Chem Soc, 2008, 130: 13818
- [16] Skoplyak O, Menning C A, Barteau M A, Chen J G. Top Catal, 2008, 51: 49
- [17] Chen J G, Menning C A, Zellner M B. Surf Sci Rep, 2008, 63: 201
- [18] Stamenkovic V R, Fowler B, Mun B S, Wang G, Ross P N, Lucas C A,

- Markovic N M. Science, 2007, 315: 493
- [19] Mu R T, Fu Q, Xu H, Zhang H, Huang Y Y, Jiang Z, Zhang S, Tan D L, Bao X H. J Am Chem Soc, 2011, 133: 1978
- [20] Mu R T, Fu Q, Liu H Y, Tan D L, Zhai R S, Bao X H. Appl Surf Sci, 2009, 255: 7296
- [21] Mu R T, Guo X G, Fu Q, Bao X H. J Phys Chem C, 2011, 115: 20590
- [22] Zafeiratos S, Piccinin S, Teschner D. Catal Sci Technol, 2012, 2: 1787
- [23] Kitchin J R, Reuter K, Scheffler M. Phys Rev B, 2008, 77: 075437
- [24] Reuter K, Scheffler M. Phys Rev B, 2003, 68: 045407
- [25] Li W-X, Stampfl C, Scheffler M. Phys Rev Lett, 2003, 90: 256102
- [26] Kresse G, Hafner J. Phys Rev B, 1993, 48: 13115
- [27] Kresse G, Furthmuller J. Phys Rev B, 1996, 54: 11169
- [28] Bloechl P E. Phys Rev B, 1994, 50: 17953
- [29] Kresse G, Joubert D. *Phys Rev B*, 1999, 59: 1758
- [30] Perdew J P, Chevary J A, Vosko S H, Jackson K A, Pederson M R, Singh D J, Fiolhais C. Phys Rev B, 1992, 46: 6671
- [31] Ma Y, Balbuena P B. Surf Sci, 2008, 602: 107
- [32] Su H Y, Gu X K, Ma X F, Zhao Y H, Bao X H, Li W X. Catal Today, 2011, 165: 89

#### 氧气气氛下Pt<sub>3</sub>Ni(111)表面结构变化的从头算原子热力学研究

孙大鹏,赵永慧,苏海燕,李微雪\*

中国科学院大连化学物理研究所催化基础国家重点实验室, 辽宁大连116023

摘要:采用从头算原子热力学方法系统研究了Ni-rich和Pt-rich条件下Pt<sub>3</sub>Ni(111)在不同偏析、表面化学吸附氧覆盖度下560个可能结构的相对稳定性,构建了氧气气氛下Pt<sub>3</sub>Ni(111)表面结构演化、直至满覆盖化学吸附氧的热力学相图.结果表明,随着氧的化学势的升高,在热力学上仅出现两类稳定的结构,主要包括没有化学吸附氧的干净Pt-skin表面,以及在很低氧的化学势下就形成的含有化学吸附氧的Ni-skin表面,而有化学吸附氧的PtNi表面合金化的中间结构则处于亚稳态.仔细分析发现,这些结构的形成主要由金属的偏析能、氧与两种金属成键强弱的差别、氧的化学势的高低三个因素共同决定. 关键词: Pt<sub>3</sub>Ni; 氧气; 偏析; 相图; 从头算

收稿日期: 2013-01-08. 接受日期: 2013-04-28. 出版日期: 2013-07-20.

\*通讯联系人. 电话: (0411)84379996; 传真: (0411)84694447; 电子信箱: wxli@dicp.ac.cn

基金来源: 国家自然科学基金(21173210, 21225315, 21273224, 21103164).

本文的英文电子版由Elsevier出版社在ScienceDirect上出版(http://www.sciencedirect.com/science/journal/18722067).

#### 1. 前言

合金催化剂广泛应用于多相催化反应中, 近年来尤以Pt基贵金属与活泼金属形成的二元合金体系的研究居多[1-10]. 在水煤气变换[11]、氧还原[12-15]和加氢反应[16,17]中, 合金表面因其双功能特点而往往成为催化反应的活性中心. 作为燃料电池工业的一种电极催化剂, Pt-Ni合金因具有比纯Pt更高的活性和更强的抗CO中毒能力, 而引起广泛的关注. 研究发现, Pt<sub>3</sub>Ni(111)表面对氧还原催化活性比Pt(111)表面提高了一个数量级, 而比工业中使用的Pt/C催化剂提高了近两个数量级<sup>[18]</sup>. 通过调变Pt-Ni体系的制备条件以及切换氧化或还原气氛, 能够可逆地改变表面结构, 发现通过控制温度和反应气氛得到的"三明治"结构NiO<sub>1-x</sub>/Pt/Ni/Pt(111), 相比Pt-Ni体

系其它结构,显示出最高的CO氧化反应活性[19~21].

综上可见,合金催化剂的活性、选择性及稳定性与 其表面的组分、化学价态和结构紧密相关.合金表面的 几何结构和电子结构等性质既不同于各单独组分,又不 同于合金的体相.在实际反应气氛下,反应物分子在合 金表面吸附或解离,由于分子与合金的各组分间的相互 作用存在差异,导致合金表面的化学组分和化学价态发 生变化,从而直接影响催化剂的机理和性能<sup>[22]</sup>.因此, 系统定量地研究制备条件和真实反应条件下Pt-Ni合金 表面组分结构的变化规律,并揭示其内在的作用机制, 对于正确揭示反应机理、优化催化剂的设计、提高性能 和稳定性具有重要的意义.

基于从头算的原子热力学方法<sup>[23-25]</sup>,本文系统研究 了在不同氧气气氛下Pt<sub>3</sub>Ni(111)面的结构演化,并考虑 到合成Pt<sub>3</sub>Ni合金过程中Pt和Ni含量可能的变化,定量地研究了该合金表面偏析和表面氧化的行为,从而构造了表面化学吸附氧直至一个单层时的热力学相图;此外,从偏析能、吸附能和氧气化学势等因素出发,揭示氧化诱导表面偏析的热力学本质及其相互制约的关系,以期对类似体系的制备和反应条件下结构稳定性的研究有所启发.

#### 2. 理论研究方法

#### 2.1. 氧气气氛下合金表面热力学

在任意给定的温度T和氧气压力 $p_{02}$ 情况下,合金表面的稳定性由其表面形成自由能来描述,相应的自由能越低,表面越稳定.以Pt-Ni双金属合金为例,其表面形成自由能为:

 $\gamma = (G_s - N_s x_{P_t} \mu_{P_t} - N_s x_{N_i} \mu_{N_i} - N_o \mu_o)/A$  (1) 其中,  $N_s \pi N_o \gamma$ 别是体系金属原子和氧原子的总数;  $x_{P_t}$  和 $x_{N_i} \gamma$ 别是体系中Pt和Ni的摩尔分数, 且 $x_{P_t} + x_{N_i} = 1$ ; A为表面面积,在下文所述中心对称的单胞模型下,是一侧表面单胞面积的2倍;  $G_s$ 是扩展体系的吉布斯自由能,通常情况下可近似为体系总能 $E_s^{[24]}$ . 给定温度和压力下氧的化学势 $\mu_o$ 为:

$$\mu_{O} = 0.5 \times (E_{O2} + \Delta \mu_{O2}(T, p^{\circ}) + k_{B}T \ln(p_{O2}/p^{\circ})) = 0.5 \times E_{O2} + \Delta \mu_{O}(T, p_{O2})$$
(2)

其中 $k_B$ 为波尔兹曼常数,  $E_{O2}$ 为孤立氧分子的总能,  $\Delta\mu_{O2}(T, p^\circ)$ 可在热力学手册中查得( $p^\circ$  = 0.1 MPa).  $\mu_{Pt}$ 和 $\mu_{Ni}$ 分别是Pt和Ni的化学势. 对于本文所研究的Pt<sub>3</sub>Ni有序体相合金,在热力学平衡时4 $\mu_{Pt3Ni}$  =  $3\mu_{Pt} + \mu_{Ni}$ , 其中 $\mu_{Pt3Ni}$  是Pt<sub>3</sub>Ni体相合金的化学势,可近似为Pt<sub>3</sub>Ni体相合金每个原子的平均总能 $E_{Pt3Ni}$ . 考虑到实际的材料合成过程中所用Pt和Ni源不同,使得相应的化学势有所变化,所生成的合金中Pt和Ni的含量会与理论计量比发生一定的偏差,进而影响合金在氧气气氛下的结构演化. 为了很好地描述这一情况,将方程(1)改写为:

$$\gamma \approx (E_{\rm s} - N_{\rm s} E_{\rm Pt3Ni} - N_{\rm s} \Delta \mu_{\rm Pt-Ni} (x_{\rm Pt} - 0.75) - 0.5 \times N_{\rm O} E_{\rm O2} - N_{\rm O} \Delta \mu_{\rm O} (T, p_{\rm O2})) / A$$
(3)

其中,  $\Delta\mu_{\text{Pt-Ni}} = \mu_{\text{Pt}} - \mu_{\text{Ni}}$ . 为了保证Pt<sub>3</sub>Ni合金相的稳定而不发生两相分离, Pt和Ni的化学势不能任意变化, 必须满足 $\mu_{\text{Pt}} \leq \mu_{\text{Pt(fcc)}}$ 和 $\mu_{\text{Ni}} \leq \mu_{\text{Ni(fcc)}}$ . 其中 $\mu_{\text{Pt(fcc)}}$ 和 $\mu_{\text{Ni(fcc)}}$ 分别是体相fcc Pt和fcc Ni的化学势, 分别对应于富铂(Pt-rich)和富镍(Ni-rich)的极端情况. 根据4 $\mu_{\text{Pt3Ni}} = 3\mu_{\text{Pt}} + \mu_{\text{Ni}}$ , 可以推导出 $\Delta\mu_{\text{Pt-Ni}}$ 允许变化的范围:

$$\Delta\mu_{\text{Pt-Ni}}(\text{Ni-rich}) \leq \Delta\mu_{\text{Pt-Ni}} \leq \Delta\mu_{\text{Pt-Ni}}(\text{Pt-rich})$$
 (4)

其中, $\Delta\mu_{\text{Pt-Ni}}$  (Ni-rich) =  $(\mu_{\text{Pt3Ni}} - \mu_{\text{Ni(fcc)}})/0.75$ ,  $\Delta\mu_{\text{Pt-Ni}}$  (Pt-rich) =  $4(\mu_{\text{Pt(fcc)}} - \mu_{\text{Pt3Ni}})$ . 以体相fcc Pt和fcc Ni的总能  $E_{\text{Pt(fcc)}} \pi E_{\text{Ni(fcc)}}$ 近似 $\mu_{\text{Pt(fcc)}} \pi \mu_{\text{Ni(fcc)}}$ ,可算得 $\Delta\mu_{\text{Pt-Ni}}$ 的边界条件为-0.67 eV (Ni-rich)和-0.32 eV (Pt-rich).

由式(3)可见,表面形成自由能不仅与Pt和Ni的含量有关,还取决于氧的化学势.在较高的氧的化学势下,合金表面可能会生成表面氧化物,并最终完全氧化形成稳定的体相氧化物,从而出现两相分离.本文主要研究合金表面氧化的情况,即表面化学吸附氧的覆盖度仅达到1个单层(1 ML)时的结构,而有关深度氧化的研究不在此列.

#### 2.2. 模型选择和计算方法

自旋极化总能计算和结构优化采用基于密度泛函理论(DFT)和周期性模型的VASP程序包 $^{[26,27]}$ ,原子赝势采用PAW势 $^{[28,29]}$ ,使用广义梯度近似(GGA)的PW91泛函 $^{[30]}$ ,平面波截断能为400 eV. 用包含 $\Gamma$ 点的Monkhorst-Pack格点方法对k空间简约Brillouin区分布取点,对于体相fcc结构Pt,Ni,以及Pt<sub>3</sub>Ni,k点选取( $12\times12\times12$ ). 优化的晶格常数分别为 $a_0$ (Pt) = 0.399 nm,  $a_0$ (Ni) = 0.352 nm和 $a_0$ (Pt<sub>3</sub>Ni) = 0.389 nm,与文献[31]一致. 合金表面采用7层( $1\times1$ )中心对称的原子超晶胞模型,中间三层固定为体相结构,其余部分完全弛豫,所有的氧吸附构型同样保持中心对称,真空层厚度为1.5 nm,k点选取( $6\times6\times1$ ).  $E_{02}$ 的计算采用(1.45 nm  $\times$  1.50 nm  $\times$  1.56 nm)的正交晶胞.所有结构优化的力收敛条件为小于0.1 eV/nm.

#### 3. 结果与讨论

体相Pt<sub>3</sub>Ni合金的晶体结构为L1<sub>2</sub>, Pt-Ni化学计量比为3:1. Pt<sub>3</sub>Ni(111)-(1×1)晶胞内每层包括3个Pt原子和1个Ni原子. 为了研究在Pt-rich和Ni-rich两种情况下Pt和Ni含量的变化,本文考察了表面两层共计8个金属原子相对含量发生变化的情况,相应的Pt:Ni比例从8:0逐渐变化到0:8,而体相(中间三层)Pt:Ni仍然维持在3:1. 考虑到相对位置的变化,共得到35个不等价的表面结构. 对于氧原子的解离化学吸附,为简单起见,我们仅考察了氧原子在稳定的fcc位吸附,且吸附氧原子的个数从0逐渐增至4,即对应于氧覆盖度 $\theta_0$ =0.00,0.25,0.50,0.75,1.00 ML. 在不考虑吸附衬底构型影响下,共有 $C_4^0$ + $C_4^1$ + $C_4^2$ 

相同的氧化学势下,氧原子的覆盖度越大,相应结构的表面形成自由能斜率越大,对气氛的变化也越敏感,而表面形成自由能最低的结构对应于热力学上最稳定的结构.

#### 3.1. Pt<sub>3</sub>Ni(111)表面相图

Ni-rich条件下的表面相图见图1. 在氧的化学势极 低条件下,  $\Delta\mu_0$ <-1.68 eV(对应于高温、低压的情况, 参 见图1上与化学势相对应的温度和压力值), 最稳定的结 构为表层全部由Pt原子组成而次表层全部为Ni原子组 成的没有氧化学吸附的干净Pt-skin表面(见图1(a)). 此 时,表面两层的Pt-Ni的计量比为4:4,低于相应的体相值 (3:1), 即对应于Ni-rich的情况. 随着氧的化学势的升高, 氧开始在表面上发生解离吸附,并且表面氧的覆盖度也 逐渐变大. 当 $\Delta\mu_{\rm O}$ >-1.68eV时, 形成 $\theta_{\rm O}$  = 0.50 ML表面暴 露原子全部是Ni原子而次表层Pt:Ni = 3:1的稳定结构(见 图1(b)). 此时,表面两层所有金属原子的计量比为Pt:Ni = 3:5, 仍然是对应于Ni-rich的情况. 当 $\Delta\mu_0$ >-1.58 eV时, 表面氧的覆盖度达到 $\theta_0 = 0.75$  ML, 形成的稳定结构见 图1(c). 当 $\Delta\mu_0 > -0.62 \text{ eV时}$ ,  $\theta_0 = 1.00 \text{ ML}$ , 形成的结构 见图1(d). 在这些结构变化过程中, 表面两层所有金属 原子的Pt:Ni计量比和构型与 $\theta_0 = 0.50$  ML的相同,没有 发生变化.

需要强调的是,上述计算涵盖了在不同的Pt和Ni含量变化、相对位置以及氧覆盖度情况下所有可能形成的表面结构. 随着 $\Delta\mu_0$ 的增加,所得最稳定的表面结构中没有出现 $\theta_0=0.25$  ML的中间结构;同时最稳定的表面结构表面组分从没有氧化学吸附的Pt-skin的结构直接过渡到Ni-skin的结构,没有PtNi表面合金化的中间结构出现. 这是由于偏析能与氧-金属键能之间的差异和竞争所致,具体原因将在下文讨论.

Pt-rich条件下的表面相图见图2. 当 $\Delta\mu_0$ < -1.48 eV时,氧分子未在表面上发生解离吸附. 这时形成表面全部由Pt原子组成而次表层Pt:Ni = 3:1的最稳定的Pt-skin表面(见图2(a)). 随着氧化学势的升高,氧开始在表面上发生解离吸附,并且表面氧的覆盖度也逐渐增加. 当发生解离吸附,形成 $\theta_0$  = 0.75 ML,此时表面暴露的全是Ni原子(Ni-skin),而次表层全为Pt原子(见图2(b)). 当 $\Delta\mu_0$ > -0.68 eV时,Ni-skin表面上的氧覆盖度将会达到1 ML,次表层仍然全是Pt原子(图2(c)).

同样地,随着 $\Delta\mu_0$ 由低到高变化,所形成的表面最稳定的结构并未出现 $\theta_0$ =0.25, 0.50 ML的中间结构;与Ni-rich相同,外表面的金属组分从没有氧化学吸附的

Pt-skin结构直接过渡到有氧化学吸附的Ni-skin结构,并未出现稳定的有氧化学吸附的表面PtNi合金中间结构.

#### 3.2. 吸附诱导表面偏析的贡献因素

为了理解上述计算结果, 揭示控制相应结构变化的物理本质, 定义了如下三个物理量:

$$E_{\text{seg}} = E_{\text{s}}^{\text{clean}} - E_{\text{s(Pt3Ni)}}^{\text{clean}} - N_{\text{s}} \Delta \mu_{\text{Pt-Ni}} (x_{\text{Pt}} - 0.75)$$
 (5)

$$E_{\rm b} = (E_{\rm s} - E_{\rm s}^{\rm clean} - 0.5 \times N_{\rm O} E_{\rm O2}) / N_{\rm O}$$
 (6)

$$\gamma_{\text{Pt3Ni}} = (E_{\text{s(Pt3Ni)}}^{\text{clean}} - N_{\text{s}} E_{\text{Pt3Ni}}) / A \tag{7}$$

其中 $E_{\text{seg}}$ 为偏析能,对应于所考察的不同Pt和Ni含量时各种可能的合金表面与按照体相结构中断的理想  $Pt_3Ni(111)$ 表面的能差;  $E_s$  和 $E_{\text{sep}}$  和 $E_{\text{sep}}$  为相应表面的总能.  $E_b$  为在相应的合金表面上的解离吸附氧原子的平均吸附能,  $\gamma_{Pt3Ni}$  是理想 $Pt_3Ni(111)$ 表面的表面能. 相应地,方程(3)可改写为:

 $\gamma \approx \gamma_{\text{Pt3Ni}} + (E_{\text{seg}} + N_0 E_b - N_0 \Delta \mu_0 (T, p_{02}))/A$  (8) 方程(8)表示不同Pt和Ni含量情况下合金表面的形成自由能,由相应的表面上的氧的平均解离吸附能、偏析能和理想Pt<sub>3</sub>Ni(111)表面的表面能共同决定,并且显著地依赖于氧的化学势.

图3是根据方程(5)算得不同Pt和Ni含量时干净表面最稳定结构的偏析能与相应的 $x_{Pt}$ 之间的关系. 结果显示,无论是在Pt-rich或Ni-rich条件下,随着 $x_{Pt}$ 增加,偏析能近似于线性减小;当 $x_{Pt}$ 大于75%时,偏析能甚至变为负值. 这说明在没有氧吸附的情况下,Pt倾向于在表面偏析. 这是是由于Pt的原子半径较大,表面能相对较低所致<sup>[32]</sup>. 还可以发现,当表面由Ni-skin ( $x_{Pt}$ =0%,单位(1×1)表面上包含有四个Ni原子)变为Pt-skin ( $x_{Pt}$ =100%,单位(1×1)表面上包含四个Pt原子)时,偏析能降低了2.54 eV (Pt-rich)和1.97 (Ni-rich) eV. 这意味着将一个表面Pt原子替换为一个Ni原子需平均耗能0.56 eV.

根据方程(6)计算得到氧原子在不同表面上的平均解离吸附能,结果见图4. 可以看出,在给定氧覆盖度时,随着O-Pt键在所有的氧-金属键中百分比( $x_{O-Pt}$ )增加,平均解离吸附能逐渐升高,化学键强度逐渐削弱,这是由于O-Ni键强于O-Pt键所致. 以Ni(111)和Pt(111)为例, $\theta_O$  = 0.25 ML时氧的平均吸附能分别为-2.44 eV/O和-1.23 eV/O; 另一方面,相同 $x_{O-Pt}$ 时,随着氧覆盖度的增加,平均解离吸附能逐渐升高,对应于氧-金属化学键的强度逐渐削弱. 这是由于表面上带负电的氧原子之间存在静电偶极排斥所致: 氧覆盖度越高,静电排斥越强. 由图4可得到不同Pt和Ni含量时合金表面上O-Ni和O-Pt键强度差别的定量结果以及随着氧覆盖度的变化关系. 在 $\theta_O$ 

= 0.25 ML时,当 $x_{O-Pt}$ 的比例从0%增至100%,即表面完全由O-Ni键变为O-Pt键时,氧的解离平均吸附能减小了1.62 eV. 因为所考察的氧吸附在fcc位,每一个氧原子和三个金属原子成键,这意味着所考察的合金表面上平均每个O-Ni键比O-Pt键要强0.54 eV. 当 $\theta_0$  = 0.50, 0.75, 1.00 ML时,相应的每个O-Ni键要比O-Pt键强0.64, 0.69和0.70 eV. 换言之,随着氧覆盖度增加,O-Ni键与O-Pt键能差越来越大.

#### 3.3. 吸附诱导表面偏析的总体趋势

上述分析表明,每个Ni原子偏析到表面上将导致体系总能升高约0.56 eV. 因此,在氧的化学势比较低、不发生化学吸附时,热力学上为达到自由能最低,Pt尽可能在表面上偏析,从而形成Pt-skin结构. 由图1和2可知,当分别在 $\Delta\mu_0>-1.68$  eV (Ni-rich)和 $\Delta\mu_0>-1.48$  eV (Pt-rich)时,氧开始在Pt-skin表面上化学吸附. 由图4可知,当 $\theta_0=0.50$ 和0.75 ML时,每一个O-Ni键至少比O-Pt键分别强0.64和0.69 eV,均大于偏析一个Ni原子到表面上的偏析能(~0.56 eV). 同时,当 $\theta_0 \geq 0.50$  ML时,所有暴露在表面上的金属原子都将至少和一个氧原子成键.换言之,一旦氧可以在Pt-skin表面上发生解离吸附,如果没有氧诱导偏析所引起的扩散动力学上可能的限制,解离吸附的氧原子将会诱导Ni完全偏析到表面上,直接形成稳定的 $\theta_0 \geq 0.50$  ML化学氧吸附的Ni-skin结构(见图1(b)和图2(b)).

随着氧化学势的增加, 从没有氧化学吸附的Pt-skin 结构, 直接形成有氧化学吸附( $\theta_0 \ge 0.50$  ML)的Ni-skin 结构, 而中间没有其它的稳定结构, 如PtNi表面合金的结构出现. 为了更好地理解这一点, 用红色虚线标出图1和2中理想中断的计量比Pt<sub>3</sub>Ni(111)表面上有化学吸附0.75 ML氧的表面所对应结构(见图2(s))的结果. 该表面上同时暴露有Pt和Ni原子, 且比例为3:1. 很明显, 虽然该表面的偏析能为零, 在能量上较Ni-skin更为有利, 但是由于表面上出现大量的Pt原子, O—Pt成键较O—Ni成键相对较弱, 其相应的表面形成自由能远高于同样氧覆盖度条件下Ni-skin所对应的结构(见图1(c)和图2(b)).

综上, 从Pt-skin变化到Ni-skin结构的条件为 $\Delta\mu_0$  = -1.68 eV (Ni-rich)和 $\Delta\mu_0$  = -1.48 eV (Pt-rich). 如图1和2 所示, 在800 K时, 相对应氧的压力为 $3.5 \times 10^{-6}$  Pa和 $1.2 \times 10^{-3}$  Pa. 当温度降低到300 K时, 由方程(2)算得所需氧压降为 $5.6 \times 10^{-43}$  Pa和 $2.9 \times 10^{-36}$  Pa (已超出当前实验所

能达到的真空条件);此时尽管所需氧的压力非常低,但是较低的温度导致结构变化可能会受到扩散动力学的限制,并不一定出现热力学上最稳定的结构,相反可能只生成一些亚稳态的中间结构.

迄今为止,尚未见有关氧化气氛下 $Pt_3N$ i结构变化的系统研究,但Pt(111)单晶上沉积Ni (= 1.3 ML)的实验发现<sup>[21]</sup>,在800 K超高真空条件下( $\Delta\mu_0<-1.8$ eV),沉积的Ni 全部扩散至次表层,形成Pt-skin结构;而当 $p_{02}=5.2\times10^{-5}$  Pa时( $\Delta\mu_0=-1.6$  eV),大部分Ni被偏析出表面形成NiO薄膜.考虑到理论计算交换关联泛函近似以及本文计算对象的不同,可以认为理论和实验结果较为一致.实验还发现,当温度降至700 K时,维持氧压不变,尽管此时氧的化学势( $\Delta\mu_0=-1.4$ eV)更高,Ni可被氧化得更为彻底,但实验发现,被氧化诱导偏析出来的Ni含量有所降低.这说明在700 K时,Ni由次表层偏析扩散到表面上来已经开始受动力学上限制.值得注意的是,当表面氧达到满覆盖度并进一步形成NiO膜时,氧气的活化和扩散动力学的受限将变得较为显著,从而现象有效抑制合金表面的深度氧化.

#### 4. 结论

采用从头算原子热力学方法和大量的理论计算系 统研究了氧化气氛下Pt<sub>3</sub>Ni(111)表面化学吸附氧直至满 覆盖度情况下的热力学相图. 研究发现, 随着氧化学势 的升高,在热力学上仅出现没有氧化学吸附的干净 Pt-skin表面和有氧化学吸附的Ni-skin表面两种特征结 构, 而没有形成PtNi表面合金的中间结构. 这主要是由 于O-Ni键与O-Pt键强度差刚好能够补偿Ni和Pt的偏析 能差. 另外, 当氧的化学势较低时, 合金表面倾向于从干 净的Pt-skin表面转变为有化学吸附氧的Ni-skin表面. 形 成较低的氧从Pt-skin表面由不同的Pt、Ni源(Ni-rich和 Pt-rich)所制备的Pt<sub>3</sub>Ni合金在氧化气氛下结构演化差别 不大. 该研究主要集中在热力学上, 而未考虑动力学限 制以及表面深度氧化; 而在实际条件下, 可能达不到热 力学平衡, 因此也可能出现含有氧化学吸附的、PtNi表 面合金的亚稳态中间结构; 而表面氧化膜的形成和钝化 则可能会在动力学上抑制合金表面的深度氧化. 本文所 采用的理论和分析方法具有一般性,对于研究 Pt<sub>3</sub>Ni(111)双金属催化剂在实际氧化条件下的活性相结 构及反应机理具有重要的参考价值.