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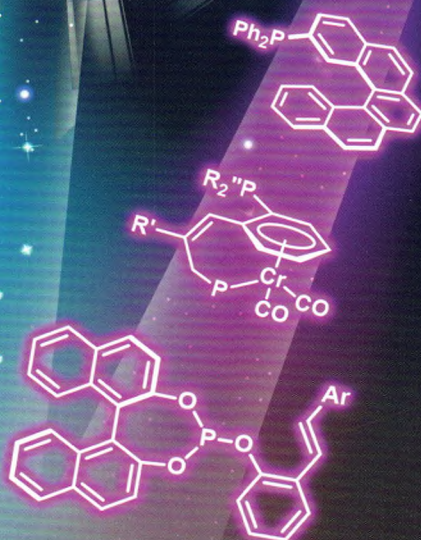
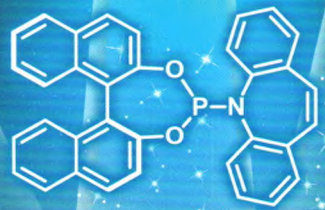
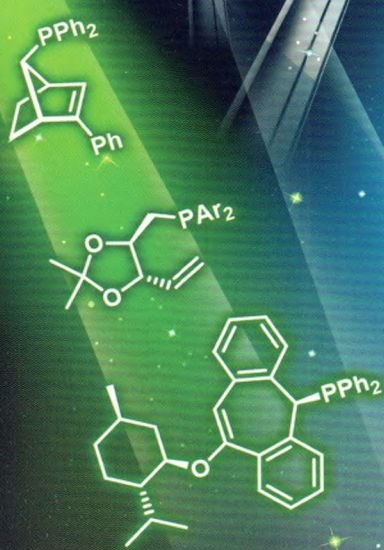


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# 化 学 报

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中国化学会  
中国科学院上海有机化学研究所

主办

# 化 学 学 报

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(Huaxue Xuebao)

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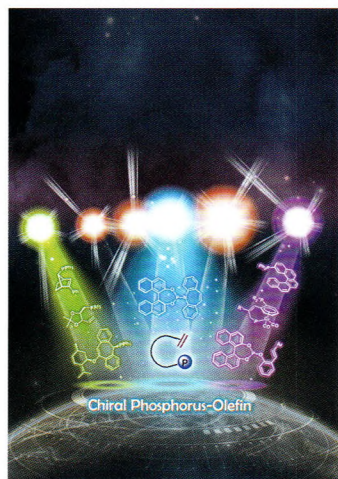
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\* 通信联系人。

Contents

**On the cover:** Since the first report of chiral phosphine-olefins for asymmetric hydrogenation by Grützmacher in 2004, phosphorus-based hybrid olefins have emerged as a conceptually new and particularly interesting class of chiral ligands for asymmetric catalysis. Over the past few years, a diverse range of structurally different phosphorus-olefin ligands have been designed and developed, and many of them have proved to be highly effective for various transition metal-catalyzed enantioselective transformations. This paper highlights the recent progress and remarkable advances in the use of various P-olefins in asymmetric catalysis. [Xu, Ming-Hua *et al.* on page 655-670.]



Review

Chiral Phosphorus-Olefin Ligands for Asymmetric Catalysis

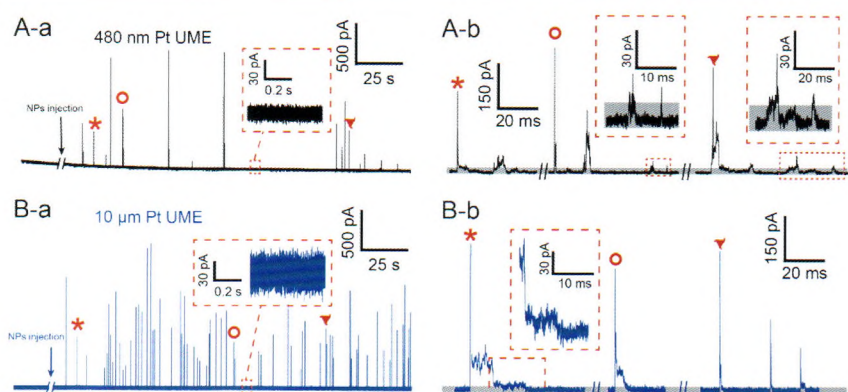


Yu, Yue-Na; Xu, Ming-Hua\*

Acta Chim. Sinica 2017, 75(7), 655-670

Communication

Study on the Resolution of Single Silver Nanoparticles Electrochemical Behavior at Nanoelectrode

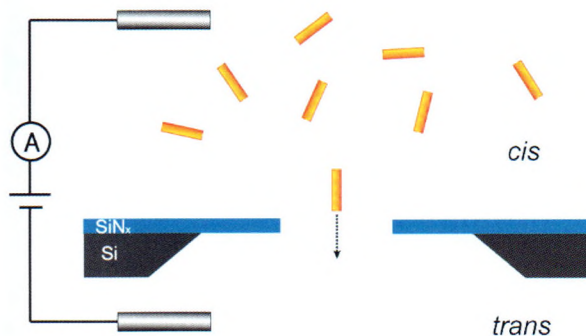


A 480 nm (diameter) nanoelectrode and a 10  $\mu\text{m}$  (diameter) microelectrode were employed as working electrode in the experiments of single AgNPs collision/oxidation. The data demonstrated that the collision frequency at varying size of electrode are in reasonable agreement with Fick's Law. In such cases, a single current peak was as a result of single collision. Result revealed that more detail information was detected by nanoelectrode with the reducing of the noise. Our findings explored that employing nanoelectrode as working electrode can enhance the current resolution of single entity electrochemistry.

Yang, Zheyao; Ma, Wei; Ying, Yilun; Long, Yitao\*

Acta Chim. Sinica 2017, 75(7), 671-674

**Analysis of Single-entity Anisotropy with a Solid-state Nanopore**

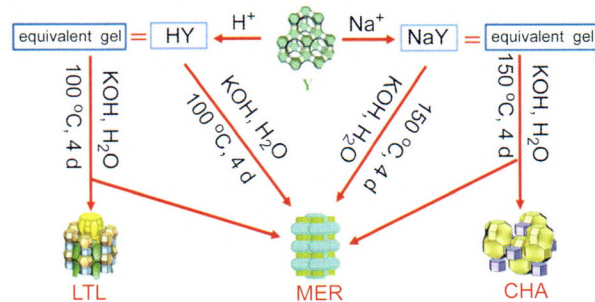


Lin, Yao; Ying, Yilun\*; Gao, Rui; Wang, Huifeng\*; Long, Yitao

*Acta Chim. Sinica* 2017, 75(7), 675-678

**Article**

**Phase Transition Behavior of Zeolite Y under Hydrothermal Conditions**

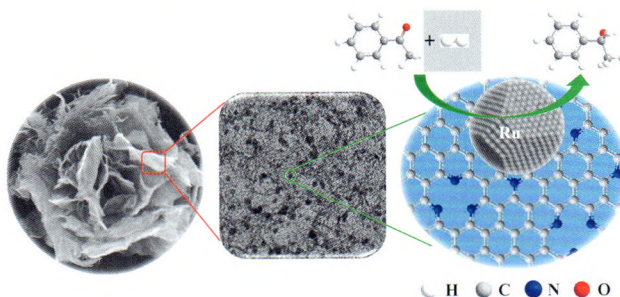


Hu, Chengyu; Yan, Wenfu\*; Xu, Ruren

*Acta Chim. Sinica* 2017, 75(7), 679-685

With phase transition of zeolite Y (HY and NaY) in the KOH solution under hydrothermal treatment, highly crystalline zeolite MER can be obtained within 2 d. Phase transition of HY can be conducted at either 100 or 150 °C, whereas that of NaY can only be conducted at 150 °C. KOH/SiO<sub>2</sub> and H<sub>2</sub>O/SiO<sub>2</sub> have significant influence on the phase transition behavior of zeolite Y.

**Ruthenium Nanoparticles Supported on Hierarchical Nitrogen-Doped Carbon Nanocages for Selective Hydrogenation of Acetophenone in Mild Conditions**

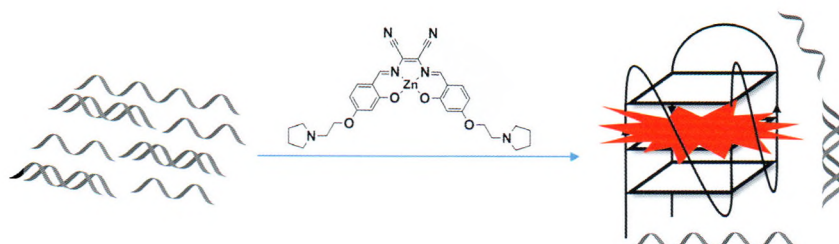


Cai, Yuejin; Liu, Chenxia; Zhuo, Ou; Wu, Qiang\*; Yang, Lijun; Chen, Qiang; Wang, Xizhang\*; Hu, Zheng

*Acta Chim. Sinica* 2017, 75(7), 686-691

Ruthenium nanoparticles ( $\approx 3$  nm) supported on hierarchical nitrogen-doped carbon nanocages (hNCNC) exhibit excellent catalytic performance for hydrogenation of acetophenone to 1-phenylethanol in mild conditions, which is attributed to the highly dispersed Ru nanoparticles owing to the high specific surface area and nitrogen dopants of hNCNC, the tuned electron structure of Ru by nitrogen incorporation, and the facile mass transportation throughout the porous hNCNC.

**Induction Stabilization and Fluorescence-based Switch-on Detection of G-Quadruplex by Zinc(II)-salen Complex**

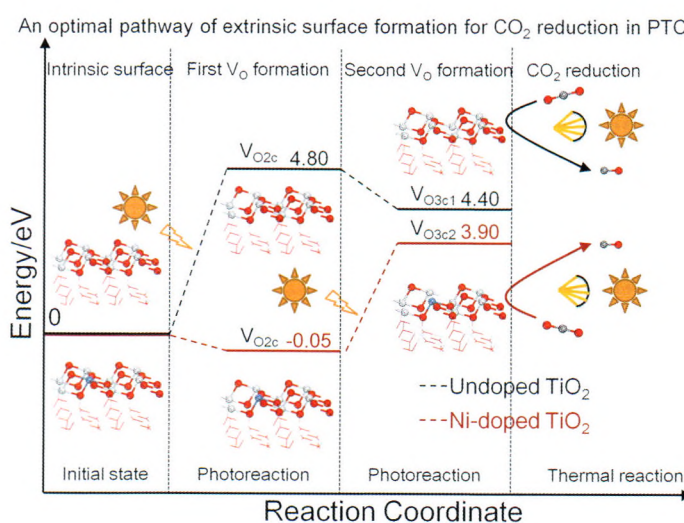


Wang, Yafen; Zhang, Xiong; Liu, Chaoxing;  
Zhou, Xiang\*

*Acta Chim. Sinica* 2017, 75(7), 692-698

Zinc(II)-salen Complex is excellent G-quadruplex DNA stabilizer, which can both selectively induce and fluorescence-based switch-on detect G-quadruplex DNA.

**Photo-thermochemical Cycle for CO<sub>2</sub> Reduction based on Effective Ni ion Substitute-doped TiO<sub>2</sub>**

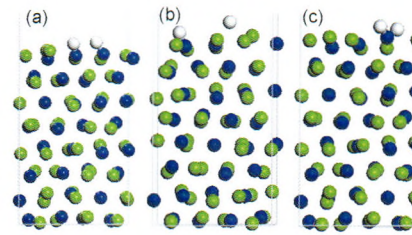


Xu, Chenyu; Lin, Jiayi; Pan, Fuqiang; Deng, Bowen; Wang, Zhihua; Zhou, Junhu; Chen, Yun; Ma, Jingcheng; Gu, Zhien; Zhang, Yanwei\*

*Acta Chim. Sinica* 2017, 75(7), 699-707

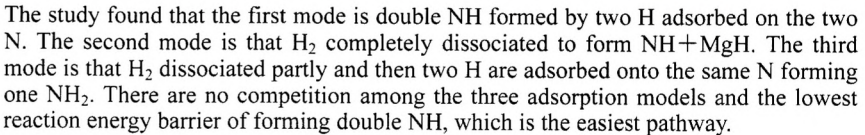
Photo-thermochemical cycle (PTC) is an innovation to dissociate carbon dioxide (CO<sub>2</sub>) utilizing both solar luminous energy and thermal energy, and the photo-induced oxygen vacancy is regarded as an important reaction core in the extrinsic surface formation step of PTC. An optimal pathway of extrinsic surface formation has been proposed by a DFT calculation. Ni substituted doping could reduce the V<sub>O</sub> formation energy in TiO<sub>2</sub>, and CO<sub>2</sub> reduction has been enhanced based on Ni doped TiO<sub>2</sub> in PTC.

**First Principles Study on the Adsorption of H<sub>2</sub> Molecules on Mg<sub>3</sub>N<sub>2</sub> Surface**



Chen, Yuhong\*; Liu, Tingting; Zhang, Meiling; Yuan, Lihua; Zhang, Cairong

*Acta Chim. Sinica* 2017, 75(7), 708-714

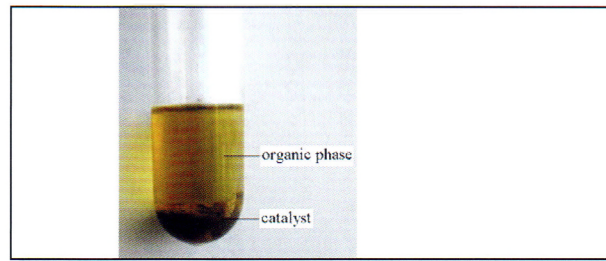


The study found that the first mode is double NH formed by two H adsorbed on the two N. The second mode is that H<sub>2</sub> completely dissociated to form NH + MgH. The third mode is that H<sub>2</sub> dissociated partly and then two H are adsorbed onto the same N forming one NH<sub>2</sub>. There are no competition among the three adsorption models and the lowest reaction energy barrier of forming double NH, which is the easiest pathway.

**Synthesis of Chiral Salen Mn(III) Complex Immobilized on Phenoxy-modified AlPS-PVPA as Catalysts for Epoxidation of Olefins**

Wang, Yifan; Fan, Yimei; Jian, Jun; Pan, Yuming; Zhao, Liang; Jing, Xueping; Zhou, Shenjia; Chen, Xiaohong\*; Du, Quan\*; Wang, Ling; Wu, Xiaoju; Fu, Xiangkai

*Acta Chim. Sinica* 2017, 75(7), 715-722



Chiral salen Mn(III) complex immobilized on phenoxy-modified AlPS-PVPA as catalysts display superior performance in the epoxidations of olefins.



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