

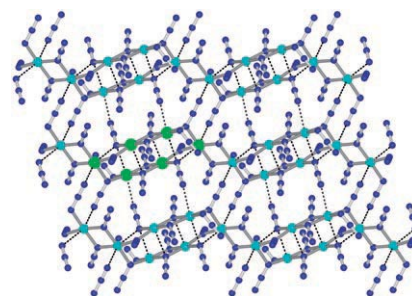
Magnetic Materials

T. Liu, Y.-F. Yang, Z.-M. Wang,*
S. Gao*

Two Diammonium Copper Azides with Similar Layerlike Magnetic Substructures Made of Chains of Serially Connected Cu₆ Rings Show Cation-Modulated Magnetism

Chem. Asian J.
DOI: 10.1002/asia.200800012

Tuning through spacing: Two compounds, each with a cationic diammonium ion and an anionic copper azide layer ([Cu₆(N₃)₁₄]²⁻) with serially connected elongated Cu₆ rings (see anionic structure; Cu green and cyan, N blue), display magnetic properties that are tuned by the nature of the diammonium counteranions located in the interlayer spacing.

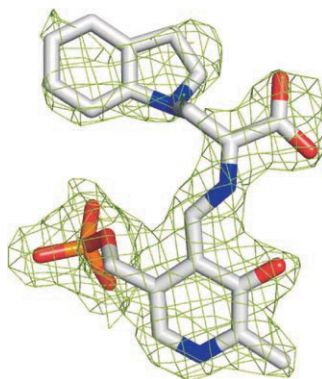


Enzyme Catalysis

T. R. M. Barends, T. Domratcheva,
V. Kulik, L. Blumenstein, D. Niks,
M. F. Dunn, I. Schlichting*

Structure and Mechanistic Implications of a Tryptophan Synthase Quinonoid Intermediate

ChemBioChem
DOI: 10.1002/cbic.200700703



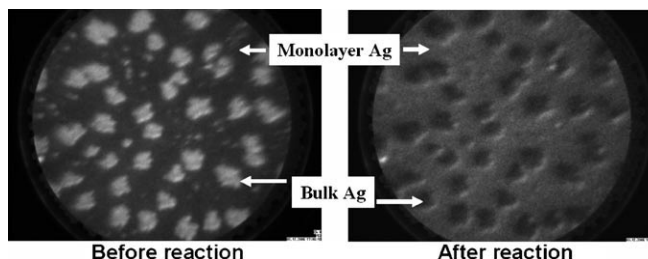
Way station. Quinonoid intermediates play a key role in the catalytic mechanism of pyridoxal 5'-phosphate (PLP)-dependent enzymes. Whereas structures of other PLP-bound reaction intermediates have been determined, a high-quality structure of a quinonoid species has not been reported. We present the crystal structure of the indoline quinonoid intermediate of tryptophan synthase (see figure) and discuss its implications for the enzymatic mechanism and allosteric regulation.

Metal Surfaces

Y.-X. Yao, X. Liu, Q. Fu,* W.-X. Li,*
D.-L. Tan, X.-H. Bao*

Unique Reactivity of Confined Metal Atoms on a Silicon Substrate

ChemPhysChem
DOI: 10.1002/cphc.200700840



Electronic effects: The unusual reactivity of the Ag monolayer compared to the bulk Ag surface on Si(111) (see figure) is due to the confinement of the 5s electrons. This change in reac-

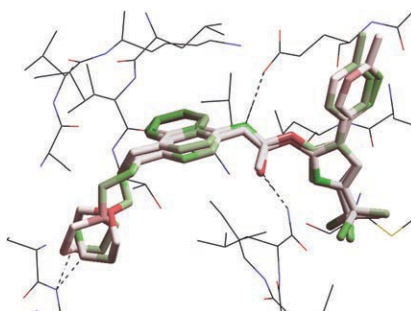
tivity has significant effects on the surface activity and selectivity, which could result in novel catalytic properties.

Molecular Design

I. Reulecke, G. Lange, J. Albrecht,
R. Klein, M. Rarey*

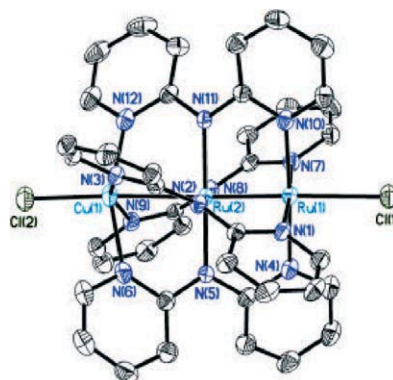
Towards an Integrated Description of Hydrogen Bonding and Dehydration: Decreasing False Positives in Virtual Screening with the HYDE Scoring Function

ChemMedChem
DOI: 10.1002/cmdc.200700319



HYDE is a new empirical scoring function for the evaluation of protein-ligand complexes that estimates binding free energy based on two terms for dehydration and hydrogen bonding only. In contrast to other scoring functions, HYDE accounts for destabilizing dehydration effects in a consistent manner, thereby decreasing the rate of false positive hits in virtual screening.

The nonplanar dipyriddyamide ligand is widely used to stabilize homonuclear strings made of three M^{II} atoms in $M_3-(dpa)_4L_2$ complexes. Heterometallic frameworks such as $Co^{II}Pd^{II}Co^{II}-(dpa)_4Cl_2$ can also be obtained. The metal strings presented here are heterometallic, nonsymmetric and mixed-valent, and a $[Ru_2]^{5+}$ moiety was coupled with a Cu^+ or Ni^+ atom.



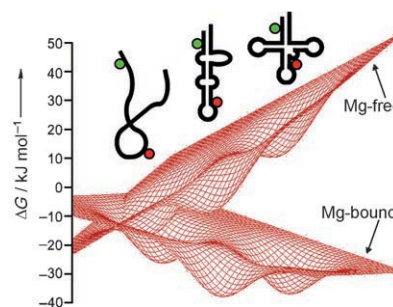
Heterometal String Complexes

G.-C. Huang, M. Bénard,*
M.-M. Rohmer, L.-A. Li, M.-J. Chiu,
C.-Y. Yeh, G.-H. Lee, S.-M. Peng*

**$Ru_2M(dpa)_4Cl_2$ ($M = Cu, Ni$):
Synthesis, Characterization, and
Theoretical Analysis of Asymmetric
Heterometal String Complexes of the
Dipyriddyamide Family**

Eur. J. Inorg. Chem.
DOI: 10.1002/ejic.200701159

With or without Me: The effect of a single methyl group on the free energies of functional and nonfunctional conformations of human mitochondrial lysine transfer RNA ($tRNA^{Lys}$) has been studied. The stabilization of the three distinct conformations by Mg^{2+} ion binding and free energy changes resulting from methylation were characterized and revealed a preference for the functionally relevant clover-leaf-shaped tRNA form (see picture).

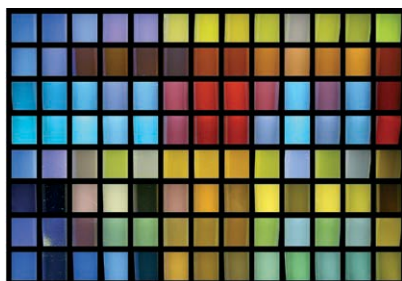


RNA Conformations

A. Y. Kobitski, M. Hengesbach,
M. Helm, G. U. Nienhaus*

**Sculpting an RNA Conformational
Energy Landscape by a Methyl Group
Modification—A Single-Molecule
FRET Study**

Angew. Chem. Int. Ed.
DOI: 10.1002/anie.200705675



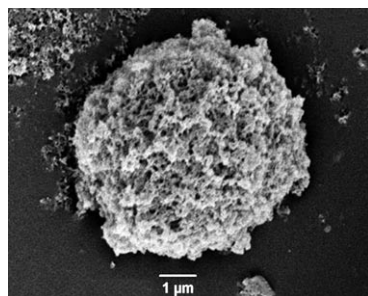
Amine detection made easy: One fluorophore in eight different solvents (columns in graphic) allows the unambiguous identification of twelve different amines (along one row) based on the distinct fluorescence changes through excited-state proton transfer.

Hydroxycruciforms

P. L. McGrier, K. M. Solntsev,
S. Miao, L. M. Tolbert,
O. R. Miranda, V. M. Rotello,
U. H. F. Bunz*

**Hydroxycruciforms:
Amine-Responsive Fluorophores**

Chem. Eur. J.
DOI: 10.1002/chem.200800296



Holey starch! Mesoporous materials with tuneable characteristics have been prepared from biomass-derived α -D-polysaccharides by a microwave-assisted strategy. Careful selection of the preparation temperature allows control of the crystallinity, particle morphology and textural properties of the resulting materials, leading to surface areas approaching $200 \text{ m}^2 \text{ g}^{-1}$ and mesopore volumes over $0.6 \text{ cm}^3 \text{ g}^{-1}$.

Mesoporous Materials

R. J. White, V. L. Budarin,
J. H. Clark*

**Tuneable Mesoporous Materials from
 α -D-Polysaccharides**

ChemSusChem
DOI: 10.1002/cssc.200800012



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