

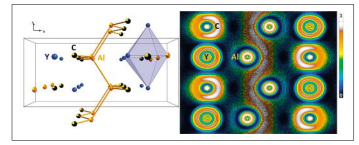
Swiss Science Concentrates

CHIMIA Column

Short Abstracts of Interesting Recent Publications of Swiss Origin

YAIC: A Bonding Chameleon with Heteropolyacetylene Features

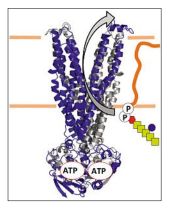
M. R. Kotyrba*, E. Cuervo-Reyes*, and R. Nesper*, Angew. Chem. Int. Ed. 2015, 54, 9606. ETH Zürich and EMPA Dübendorf Silicon carbide and aluminium carbide are extremely stable refractory compounds demonstrating the strength of the Si-C and the Al-C bonds, respectively. Kotyrba, Cuervo-Reyes and Nesper describe a novel ternary aluminide-carbide, YAIC, which was prepared by a flux method. It crystallizes as a partially filledup TII structure, showing remarkable structural aspects at the border between Zintl phases and intermetallics. The compound represents a fascinating compromise between the polarized metal (stuffed intermetallic) and the semicovalent Y³⁺[AlC]³⁻ model. A key feature is the coexistence of two-centre and multi-center bonds. The latter appear as tapes meandering wherever there would be a direct Al-Al or Y-Al contact, and contain the residual electron density after the formation of the Al-C semi-covalent chains.



Structure and Mechanism of an Active Lipid-linked Oligosaccharide Flippase

C. Perez, S. Gerber, J. Boilevin, M. Bucher, T. Darbre, M. Aebi, J.-L. Reymond, and K. P. Locher*, *Nature*, **2015**, *524*, 433. ETH Zürich

The flipping of membrane-embedded lipids containing large, polar head groups is slow and energetically unfavourable, and is therefore catalysed by flippases, the mechanisms of which are unknown. A prominent example of a flipping reaction is the translocation of lipid-linked oligosaccharides that serve as donors



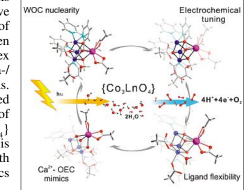
in *N*-linked protein glycosylation. In *Campylobacter jejuni*, this process is catalysed by the ABC transporter PglK. Locher and coauthors present a mechanism of PglK-catalysed, lipid-linked oligosaccharide flipping based on crystal structures in distinct states, a newly devised *in vitro* flipping assay, and *in vivo* studies. The proposed mechanism is distinct from the classical alternatingaccess model observed in other transporters.

3d–4f {Co^{II}₃Ln(OR)₄} Cubanes as Bio-Inspired Water Oxidation Catalysts

F. Evangelisti, R. Moré, F. Hodel, S. Luber*, and G. R. Patzke*, *J. Am. Chem. Soc.* **2015**, *137*, 11076. University of Zürich The { $CaMn_4O_5$ } oxygen evolving complex of photosystem II represents a major paradigm for the development of water oxidation catalysts (WOCs). However, translating its key features into active molecular WOCs remains a major challenge.

Patzke, Luber and co-workers now introduce $[Co^{II}_{3}Ln(hmp)_{4}(OAc)_{5}H_{2}O](\{Co^{II}_{3}Ln(OR)_{4}\};Ln=Ho-Yb,hmp=2-(hydroxymethyl)pyridine) cubanes as new functional and stable model systems to explore a range of crucial design parameters, including core nuclearity, redox-inactive promoters, and ligand exchange properties. The highly active <math>\{Co^{II}_{3}Ln(OR)_{4}\}$ cubanes advance promote bio-inspired design through the combination

of Ln³⁺ core cations as redox-inactive Ca2+ analogues of nature's oxygen evolving complex with flexible aqua-/ ligands. acetate The Ln³⁺-enhanced ligand exchange of the $\{Co^{II}, Ln(OR)_{4}\}$ WOCs is substantiated with molecular dynamics studies.

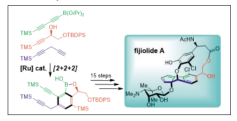


Synthesis of Fijiolide A *via* an Atropselective Paracyclophane Formation

C. Heinz and N. Cramer*, *J. Am. Chem. Soc.* **2015**, *137*, 11278. EPF Lausanne

Fijiolide A is a secondary metabolite isolated from a marinederived actinomycete displaying inhibitory activity against TNF- α -induced activation of NF κ B, an important transcription factor and a potential target for the treatment of different cancers and inflammation related diseases. Fijiolide A is a glycosylated complex paracyclophane, which is structurally closely related to the Bergman-aromatization product of enediyne C-1027. Cramer and Heinz report an enantioselective synthesis of fijiolide A demonstrating the power of fully intermolecular rutheniumcatalysed [2+2+2]-cyclotrimerizations with three different alkynes to assemble the heavily substituted central arene core.

The characteristic strained [2.6]-paracyclophane structure is accessed by a templated a t r o p s e l e c t i v emacroetherification reaction.



Prepared by Caroline D. Bösch, Markus Probst, Yuliia Vyborna, Mykhailo Vybornyi, Simon M. Langenegger and Robert Häner* **Do you want your article to appear in the SWISS SCIENCE CONCENTRATES highlight?** Please contact robert.haener@dcb.unibe.ch