# SYMEORIM

People, Trends and Views in Synthetic Organic Chemistry

2008/08

#### SYNSTORIES

- Efficient, Selective, and **Green: Catalyst Tuning for Highly Enantioselective Reactions of Ethylene**
- Total Synthesis of (+)-Pinnatoxin A

- FOCUS on the Second **European Workshop in Drug** Synthesis (II EWDSy), May 25-30, 2008, Siena (Italy)
- **Thieme-Chemistry Journals Editorial Board Meetings 2008**

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SYNFORM **A84** 



#### Dear readers,

August is holiday time for many colleagues, particularly in the south of Europe, but **SYNFORM** is still fully operative and eager to inform you about the latest trends and the most exciting achievements in organic chemistry.

Certainly exciting is the total synthesis of the complex natural structure pinnatoxin A that was elegantly synthesized by the group of Professor A. Zakarian (USA). Another important work covered in this issue of SYNFORM is the use of an abundantly available building block like ethylene for catalytic enantioselective synthesis, as described by Professor T. V. RajanBabu (USA). For the first time, SYNFORM presents a report on a symposium, specifically the Second European Workshop in Drug Synthesis, recently held in Siena (Italy). The organizers of conferences who are interested in having their event covered by **SYNFORM** are invited to get in touch (synform@chem.polimi.it) well in advance. The issue closes with a brief report on the 2008 Thieme Chemistry journals editorial board meetings recently held in Thessaloniki (Greece).

Enjoy your reading!

Matteo Zanda

Editor of SYNFORM

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# FOCUS on the Second European Workshop in Drug Synthesis (II EWDSy), May 25–30, 2008, Siena (Italy)

■ From May 25th to 30th 2008, Siena (Ital y) hosted the Second European Workshop in Drug Synthesis (II EWDSy). The venue was the prestigious "Certosa di Pontignano", with its beautiful and char ming cloisters, the Cong ress Center of the University of Siena. The site has a rich and ancient history, dating back to 1314 w hen the order of Car thusian monks was expanding throughout Italy.

In such a culturally stimulating environment, the organizer Professor Maurizio Botta (University of Siena) and the International Scientif ic Committee, for med by Professor Botta, Professor Maurizio Taddei (Italy), Professor Stephen Hanessian (Canada and USA), Professor Gerhard F. Ecker (Austria), Dr. Hugo Kubinyi (Germany), and Dr. Chiara Ghiron (Italy), put together an impressive list of international speakers. The medieval atmosphere of the Certosa, together with the additional help of fantastic food and wine, combined to produce a magic blend of science and relaxation, to the g reat advantage of the about 40 "students" attending the Workshop, who were selected on the basis of their cur riculum vitae. Not to mention the benefits for the "teaching body"... Some of the lectures that were presented at the II EWDSy are highlighted here.

The first lecture w as held by Professor Trevor M. Jones, who gave a conceptually very stimulating and informative talk on the oppor tunities and challenges for medicinal chemistry in the search for better medicines. Professor Stephen Hanessian, affiliated both to the University of Montreal (Canada) and to the University of California, Irvine (USA), described the recent achievements of his group with proximity-assisted cycloaddition reactions, and provided as usual an inspiring and passionate lesson of or ganic chemistry. Professor Oli ver Kappe from the University of Graz (Austria) explained how to fully exploit the potential of micro waves in organic synthesis. His lecture was followed by Dr. Jieping Zhu from the Institut de Chimie des Substances Naturelles of the CNRS (France), who reported on the potential of multicomponent reactions.

Professor P eter Wipf from the Uni versity of Pittsb urgh (USA) reported on recent research by his group on the synthesis and medicinal chemistry of highly cytotoxic natural molecules that target the vinca-domain of tub ulin, such as disorazoles and tubulysins, which represent very promising potential drugs for cancer chemotherapy. Professor Wipf's talk was the first of a long series dedicated to the topic of cancer . Epothi-



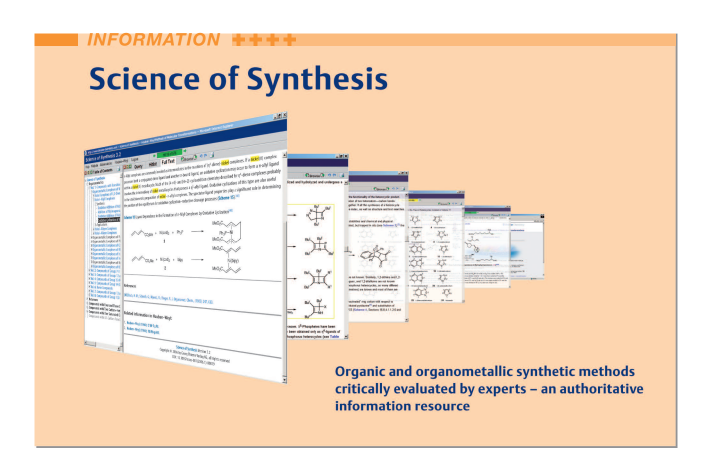
Group picture of the II EWDSy attendees

lones and their synthetic deri vatives are another class of extremely important natural molecules acting at the le vel of microtubules, interrupting cell mitosis and causing apoptosis; this was discussed by an expert in the field: Professor Karl-Heinz Altmann from the ETH Zürich (Switzerland). The targeting of cancer w as the topic presented b y Professor Lutz Tietze from the Georg-August-Universität of Göttingen (Germany), who described synthesis and bioactivity of the immunoconjugates recently synthesized by his research group. Professor Jürgen Borlak from the Medical School of Hanno ver (Germany) gave a highly appreciated and stimulating talk on cancer stem cells, and their possib le role in cancer re-g rowth and metastasis, while Professor Alan P. Kozikowski from the University of Illinois at Chicago (USA) g ave a lecture about new findings on histone deacetylase inhibitors, which are also of interest in the f ield of CNS disorders and malaria. Professor Stephen Neidle from the London School of Phar macy (UK) covered the use of G-Quadr uplex DNAs as therapeutic targets in human cancer, and Professor Carlo Melchiorre from the University of Bologna (Italy) gave a talk on new strategies for identifying high-af finity lig ands for the therap y of the Alzheimer disease.

Last b ut not least, a round-tab le chaired by Professor Antonio Giordano (University of Siena) addressed the increasingly important topic of "stem cells", with the participation of international experts from academia and industry. The rich program was completed by at least 10 further multidisciplinary lectures, and by a practical training program involving organic synthesis problems, chiral HPLC purifications and microwave-promoted reactions. A small poster session, from which a few communications were selected for an after-dinner oral presentation (with beer and wine served) was also included.

Overall, the II EWDSy w as a v ery useful and e xtremely well or ganized e vent that cer tainly can be considered as a small gem in the cro wded agenda of courses and w orkshops dedicated to organic chemistry.

Matteo Zanda



## Efficient, Selective, and Green: Catalyst Tuning for Highly Enantioselective Reactions of Ethylene

Org. Lett. 2008, 10, 1657-1659

According to Professor T. V. RajanBabu from the Chem istry Department of the Ohio State Uni versity in Columb us (USA) the "discovery of efficient catalytic processes for activation and selective incorporation of ab undantly a vailable feedstocks such as eth ylene, butadiene, styrene, HCN, CO, and H<sub>2</sub> could have an enor mous impact on ho w chemical intermediates are synthesized in the lab, and eventually manufactured on an industrial scale. Among these," he continued, "processes that yield practical levels of asymmetric induction will be especially attractive to synthetic chemists. Seldom has chemistry seen an area where the scientific goals are so challenging, the economic and environmental benefits so obvious, and the intellectual and ethical reasons for doing the research so compelling. Through an approach that relies primaril y on mechanistic insights and a systematic e xamination of lig and effects, our g roup has been looking for highly efficient and selective protocols for the use of feedstock materials for asymmetric synthesis."

Recently, the RajanBab u g roup described an impro ved methodology for the asymmetric hydrovinylation (HV) of styrene derivatives using ethylene as the vinyl group source.

"This paper is an impor tant milestone in our w ork on the nickel-catalyzed asymmetric hydrovinylation," said Professor RajanBabu, "a reaction that was initially discovered by the Wilke group at Mülheim almost 36 y ears ago (Review: T. V. RajanBabu Chem. Rev. 2003, 103, 2845). This reaction received only scant attention until Nobu Nomura in our group discovered a ne w protocol (J. Am. Chem. Soc. 1998, 120, 459). Our g roup also disco vered stringent demands with respect to ligands and counter ions (J. Am. Chem. Soc. 1999, 121, 9899) for this exacting reaction." Since then, a number of successful lig ands, among them phospholanes ( Org. Lett. 2004, 6, 1515; J. Am. Chem. Soc. 2006, 128, 54), phosphinites (J. Am. Chem. Soc. 2002, 124, 734) and phosphoramidites (the title paper and others cited under reference 6) ha ve been successfully used for asymmetric h ydrovinylation reactions of vin yl arenes, 1,3-dienes and strained bic yclic molecules. "The Feringa phosphoramidite, f irst used by Leitner's group (J. Am. Chem. Soc. 2002, 124, 736) under our protocols," continued Professor RajanBabu, "are especially valuable for the reaction of vinyl arenes. As this communication illustrates, this highly tunable ligand can be easily modified to

New approaches to the control of exocyclic stereochemistry

achieve the best selectivities observed to date for all classes of substrates, including vinylarenes that produce all-carbon quaternary centers ( *J. Am. Chem. Soc.* **2006**, *128*, 5620)." Especially noteworthy is the synthesis of 3-arylbutenes, which are precursors for widel y used 2-ar ylpropionic acids such as ibuprofen, naproxen and k etoprofen. "Enantiomeric excesses over 96 % can be achie ved for these v aluable intermediates," explained Professor RajanBabu. "In more recent w ork, Craig Smith has since achie ved substrate-to-catalyst ratios of o ver 7000 for some of these transfor mations, making this one of the most selective and efficient carbon—carbon bond-forming reactions known."

Dan Mans, a recent g raduate from the RajanBab u group, completed the total syntheses of v arious pseudopterosins using back-to-back h ydrovinylations of vin ylarenes and dienes, demonstrating the excellent functional group compatibility of these reactions. "The inter mediates he prepared are potentially useful for other highly sought after molecules such as elisabethin and colombiasin, " conf irmed Professor RajaBabu. "The hydrovinylation reaction has also been used to construct side chains of steroid D-rings car rying either 20(S)- or 20(R)- substituents (J. Am. Chem. Soc. 2008, 130, 9000). These discoveries provide a new approach to the control of exocyclic stereochemistry in several biologically relevant molecules like arylpropionic acids, pseudopterosins and steroids," he added. "Detailed procedures for the syntheses of ligands described in this paper and the h ydrovinylation reactions have recently appeared in the literature ( Org. Synth. **2008**, 85, 235; 245)."

"Our ongoing research encompasses reactions of bic yelic and acyclic dienes (best to date: 80% ee), use of other metals (Ru, Co) and ligands to expand the scope of the reaction, and further applications in natural product synthesis, "concluded Professor RajanBab u. "Exciting leads on ruthenium- and cobalt-catalyzed reactions have been reported by Yi and Vogt. A recent modification of the ruthenium-catalyzed HV reaction (*J. Org. Chem.* **2003**, *68*, 8431) should provide further impetus for work in this area."

Matteo Zanda



From left: Prof. T. V. RajanBabu, Craig R. Smith

#### About the corresponding authors

T. V. (Babu) RajanBabu received his undergraduate education in India (Kerala University and IIT, Madras). He obtained his PhD degree from The Ohio State University (USA) under the direction of Professor Harold Shechter, and was a post-doctoral fellow at Harvard University with the late Professor R. B. Woodward. He then joined the Research Staff of Dupont Central Research. He returned to Ohio State as a Professor of Chemistry in 1995. His research interests are in new practical methods for stereoselective synthesis focusing on enantioselective catalysis, free-radical chemistry, applications in natural product synthesis, and organic reactions in water.

#### Thieme-Chemistry Journals Editorial Board Meetings 2008



The White Tower of Thessaloniki

The 2008 Editorial Board Meetings of the Thieme-Chemistry journals SYNTHESIS, SYNLETT, and SYNFACTS, including the supplement SYNFORM, were held on June 20th and 21st 2008, in the vibrant city of Thessaloniki (Greece). The Editorial Boards of the three journals and the staf f of the Thieme-Chemistry editorial of fice met at the Mediterranean Palace Hotel

as shown in the group picture. Many topics were addressed in the general meeting session, as well as in the single jour nals meetings, including a brand new joint Editorial Advisory Board that will be full y operative from 2009 onwards, and a stimulating discussion on the strategies that will be pursued to further increase the quality of the jour nals. The program was completed by a highly appreciated cultural event—a guided tour of the archaeological site of Vergina, about 80 km from Thessaloniki, that hosts the magnificent palaces and tombs of ancient Macedonian kings, including Philip II, and queens, with perfectly preserved works of miniature art and original paintings. The appointment for the next Editorial Board Meetings is in 2009 in Granada (Spain).

Matteo Zanda



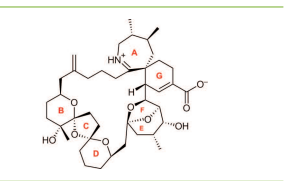
#### Total Synthesis of (+)-Pinnatoxin A

J. Am. Chem. Soc. 2008, 130, 3774-3776; Synfacts 2008, 782 (SYNFACT of the Month)

According to Professor Armen Zakarian from the Uni versity of California at Santa Barbara (USA), from a certain perspective the total synthesis of complex natural products can be occasionally regarded as a 'guilty pleasure' of Organic Chemistry. "For example," he said, "for many people outside the field, it is sometimes not easy to imagine that the chemical synthesis of 0.9 mg of the powerful marine toxin palytoxin A would be anything more than a high-le vel chemical chess game, which requires a very expensive chess board with no apparent practical benefits." Possibly, one of the reasons is that the benefits, although substantial, are not direct. "Unlike aspirin," continued Professor Zakarian, "which can be used to relieve a headache, the anesthetic potential of pal ytoxin A is unlik ely to attract consumers, not to mention its prohibitive cost." Another possible reason might be that the endea vor of complex molecule synthesis is largely unpredictable. "One has to have a rational plan," he e xplained, "b ut also anticipate that things are not going to go according to the plan. The important specific contributions resulting from the work culminating with the synthe sis of, for example, palytoxin A, although quite logical in retrospect, could hardly be predicted at the onset of the endea vor. They are, however, now employed in the preparation of pharmaceuticals and many other useful organic materials."

The interest of Professor Zakarian and his group in the synthesis of pinnato xin A began with a f ascination with its chemical structure, while keeping its potent bioacti vity in mind. The 27-membered all-carbon macroc yclic frame work of the marine to xin incor porates several medium-sized rings, most notably a unique 6,7-spirobic yclic imine, w hich is a critical constituent of the phar macophore. In addition, a 6,5,6-spirotricyclic bis-ketal and a dioxabicyclo[3.2.1]octane ring system are embedded in the macrocycle.

"It was pretty clear that our overall synthetic strate gy would have to accommodate for the small size of the new group," said Professor Zakarian. "It was one of the very first projects for me as an Assistant Professor, so efficient use of resources was a crucial factor." Thus, an early strategic decision was to develop a convergent synthesis plan. According to Professor Zakarian, another easily identified challenge in the synthesis, based on both the structural analysis and prior art, was going to be the assembly of the spiroimine with its quaternary chiral center at the core. "Thus," he confirmed, "we planned to explore this challenge at the onset of the total synthesis, and our synthesis plan would have to allow for that."

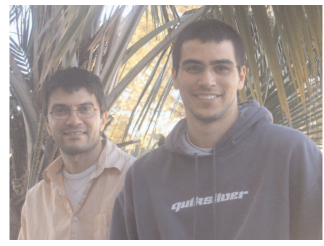


Pinnatoxin A

"Because several groups investigated a Diels-Alder approach to form ring G and the quater nary stereocenter re vealing its challenges," e xplained Professor Zakarian, "w e became in trigued by the potential of a [3,3]-sigmatropic process as a solution to this synthetic problem. Our initially conceived tandem sigmatropic Claise n-Mislow-Evans rear rangement, developed by graduate student Matthe w Pelc, was very successful in b uilding the quater nary stereocenter and ring G. " However, this strate gy w as ultimatel y abandoned because Zakarian and co-workers could not properly functionalize the product, in spite of e xtensive efforts. "Needless to sa y, it is times like these that put one's mental perseverance to the test," said Professor Zakarian. The group was able to redesign the synthesis almost completely, while keeping the Claisen rearrangement as a key concept behind the synthesis plan. Importantly, this exercise led to a solution for a long-standing limi tation of the classic Ireland-Claisen rearrangement, which has previously had poor diastereoselecti vity with α-branched esters. Highly stereoselective rearrangement could be achieved using readily available chiral Koga-type bases.

"The synthesis of the B ,C,D-dispirotricylic fragment w as achieved by Postdoc Chongdao Lu, who joined the group after obtaining his doctorate from the Chengdu Institute of Organic Chemistry, China," said Professor Zakarian. "The total synthesis was ultimately completed thanks to the skillful hands of another graduate student, Craig Stivala, whose previous athletic career in track and field fortunately [for me] ended with an injury in the pole vault," he concluded, "thus allowing him to pursue his true passion – Organic Synthesis!"

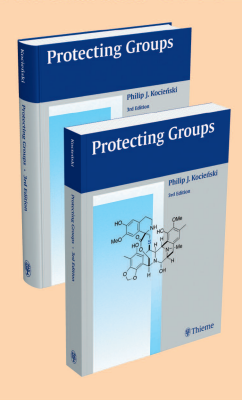
According to Dr. David Chen from the Chemical Synthesis Laboratory @ Biopolis, A\*Star Sing apore, an expert in the total synthesis of comple x natural molecules, "chemical synthesis of marine toxins continues to serve as the cradle for the chemical and biological investigations of these har mful substances. The recent total synthesis of pinnatoxin A reported by Zakarian and Sti vala illustrated, in a highly efficient and stereocontrolled manner, one of the most comple x examples of chirality transfer by means of an Ireland —Claisen rearrangement. In a conceptuall y different approach to the pioneering synthesis repor ted Kishi and co-w orkers, the macroc yclic scaffold was casted by the venerable RCM reaction. The work reported by the Zakarian g roup," concluded Dr. Chen, "is a beautiful demonstration of the state-of-the-art technologies in complex natural product synthesis."



Matteo Zanda

From left: Prof. A. Zakarian, C. Stivala

#### **INFORMATION**



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P. DeShong, University of Maryland, USA, J. AM. CHEM. SOC., 2006

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T. Laird, Scientific Update, UK, Organic Process Research & Development, 2005

#### COMING SOON ▶ ▶ COMING SOON ▶ ▶

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#### In the next issues:

#### SYNSTORIES . .

■ The Total Synthesis of (–)-Cyanthiwigin F by Means of **Double Catalytic Enantioselective Alkylation** 

(Focus on an article from the current literature)

■ Nickel-Catalyzed Cross-Coupling of Aryl Methyl Ethers with Aryl Boronic Esters

(Focus on an article from the current literature)

- Pd(II)-Catalyzed Cross-Coupling of sp³ C-H Bonds with sp<sup>2</sup> and sp<sup>3</sup> Boronic Acids Using Air as the Oxidant (Focus on an article from the current literature)
- Asymmetric Total Synthesis of (-)-Quinocarcin (Focus on an article from the current literature)
- Stereoselective Synthesis of β-L-Rhamnopyranosides (Focus on an article from the current literature)

#### ■ FURTHER HIGHLIGHTS

#### **SYNTHESIS**

Special Topic on "N-Heterocyclic Carbenes" in issue 17/2008

#### SYNLETT

Account on: Corrole: The Little Big Porphyrinoid (by R. Paolesse)

#### **SYNFACTS**

Synfact of the Month in category "Metal-Catalyzed Asymmetric Synthesis and Stereoselective Reactions":

Synthesis of Substituted Enol Ethers and Their Synthetic **Application** 

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