

Heterocycles in the Service of Humankind

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Abstract

Alleviating the scourge of biting insects; Bioconjugates for new directions in pharmaceutical research; Sharing the benefits of research

Keywords: Malaria, insect repellents, bioconjugates, chemistry publishing

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4. Attempts to help Chemistry in Developing Countries by Innovations in the Publishing and Dissemination of Organic Chemistry Research

1. Overcoming the Problems of Biting Insects (Research with the participation of U. Bernier, D. Dobchev, G. Clark, C. D. Hall, K. Linthicum, S. Slavov, M. Tsikolia, Z. Wang)

Mosquito-borne diseases such as malaria, arboviral encephalitis, dengue fever, Rift Valley fever, and yellow fever, still result in significant morbidity and mortality in humans. Insect repellents often serve as a first line of personal protection. Disadvantages of current repellents include limited duration of protection from insect bites due to evaporative loss, absorption into the skin, removal by dissolution in water, skin irritation and a stinging sensation when in contact with eyelids or lips. Some repellents are only efficient when used in large quantities on the skin or

clothing. The repellent most often used is DEET but a major drawback is that DEET can lead to adverse side effects in some individuals, especially from its systemic uptake via dermal absorption.

New, improved long lasting repellents that are safe and efficacious against a wide range of insects are needed. Thus, there is a long-standing interest in the design of compounds that will be effective repellents against mosquitoes, sand flies, stable flies, black flies, tsetse flies, biting gnats and tabanids, all of which spread human and animal diseases.

Although the design of new insect repellents by computer-aided molecular modeling was discussed in a recent book, there have been few previous attempts to correlate the repellency of compounds towards mosquitoes with their chemical structure.² Suryanarayana *et al.*¹ used a small set of 31 repellants to propose the model (eqn 1) with the rather low R^2 of 0.304: where PT = Protection Time; $\log P$ = lipophilicity; V_p = Vapor pressure; ML = Molecular Length.

$$PT = \mathbf{a} \log P + \mathbf{b} \log V_p + \mathbf{c} \log ML + d \quad (1)$$

With the same data-set, and using descriptors derived solely from the chemical structures of the repellants we found an improved model with an R^2 of 0.79. This encouraged us to go further. Over the last few years we have been collaborating with the U.S. Department of Agriculture at the University of Florida, which over the past 60 years have recorded tests covering a wide range of insect species and some 30,000 different compounds. Properties such as protection time, effective dose and lethal dose towards a large numbers of insects including various species of mosquitoes, houseflies and other pests were amassed. We were given access to these records and in agreement with the Department of Agriculture embarked on a program to try and extract from them some information that would allow us to understand the relationship between the biological activity of compounds and their chemical structures. This lecture concerns the work that we have carried out with *N*-acylpiperidines, and which is described in detail in *Proc. Nat. Acad. Sci. USA*, 2008.

The USDA records included more than 150 different *N*-acylpiperidines which had been tested for their effectiveness as mosquito repellants. These compounds were classified according to their effectiveness into five classes, 1 being the least active and class 5 being the most active. The most active compounds retained effectiveness for 21 days. By building a neural network model, we were able to correlate protection times from the old USDA data with the chemical structures as is shown in Figure 1.

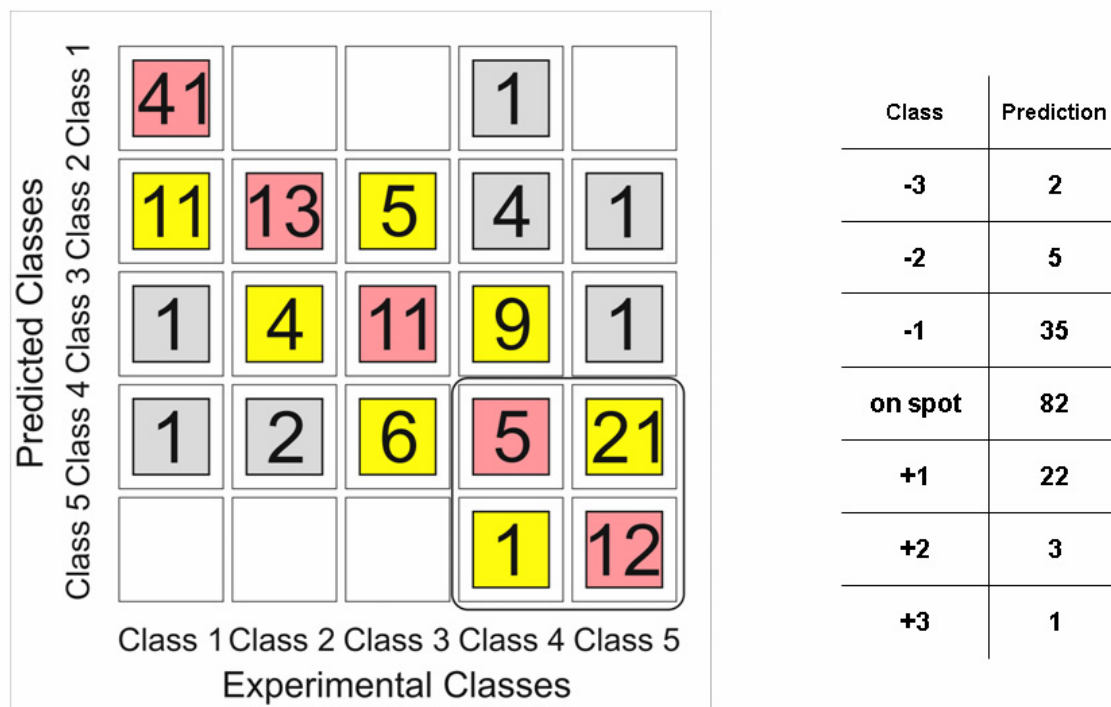


Figure 1. Correlation of protection times (old USDA data) with chemical structures by ANN.

We re-synthesized 11 of the *N*-acylpiperidines that were previously tested in order to anchor the envisaged biological testing into the existing data. Piperidines were synthesized utilizing acylbenzotriazoles prepared from carboxylic acids by treatment with thionyl chloride and benzotriazole. In addition to the 11 previously tested *N*-acylpiperidines, we synthesized 23 novel *N*-acylpiperidines as potential repellants. These 23 compounds were selected by first using our neural network model to predict the likely activity of many hundreds of compounds. Most of the 23 compounds selected were expected to be highly active but a few were also chosen that were predicted to have lower activity.

Biological testing was carried out for all 23+11=34 compounds synthesized, together with DEET. Bioassays were conducted by covering the hand of a volunteer with a soft-embossed long cuff poly glove and powder-free latex glove. To cover the arm a stocking is pulled over. A sleeve with an opening (3 cm x 8 cm) was fastened around the arm. Each cloth patch assembly was affixed over the open window with masking tape to hold it in place on the sleeve. The arm is then inserted into the cage of mosquitoes and held stationary for 1 minute. The number of feeding mosquitoes was counted prior to removal with a quick, brisk shake of the arm. Feeding mosquitoes that remained in the window were considered to have been biting.

The failure threshold for repellency for these experiments was set at 1% biting (5 bites) confirmed by achievement of two consecutive days of 5 or more bites. The results are shown in Table 1 and graphically in Figure 2. Gratifyingly, several of the compounds prepared showed considerably improved protection times compared to DEET.

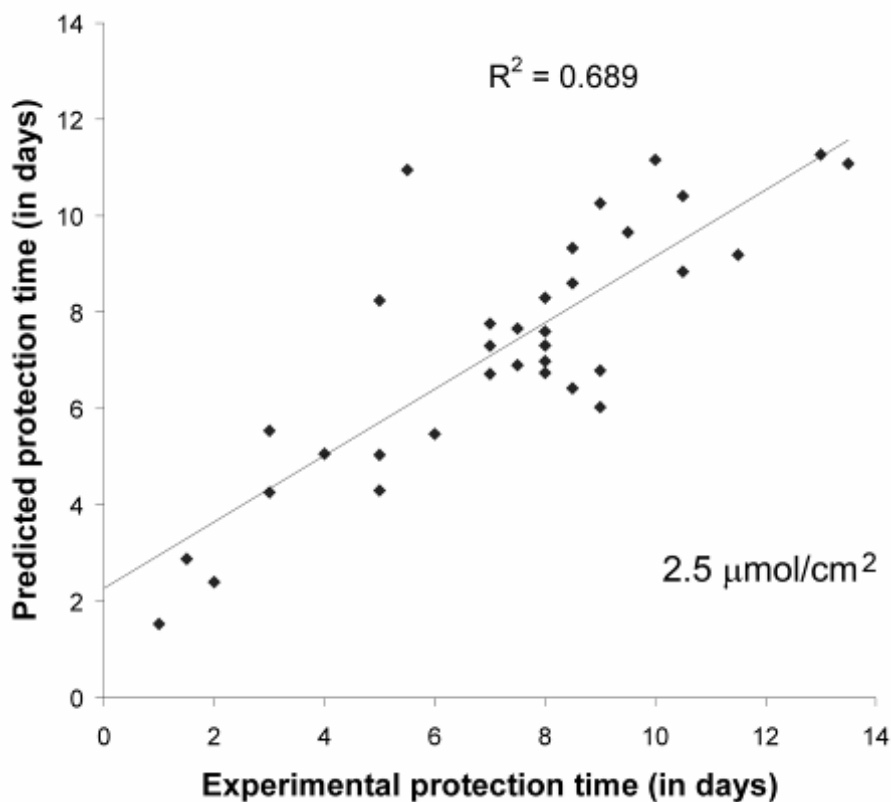
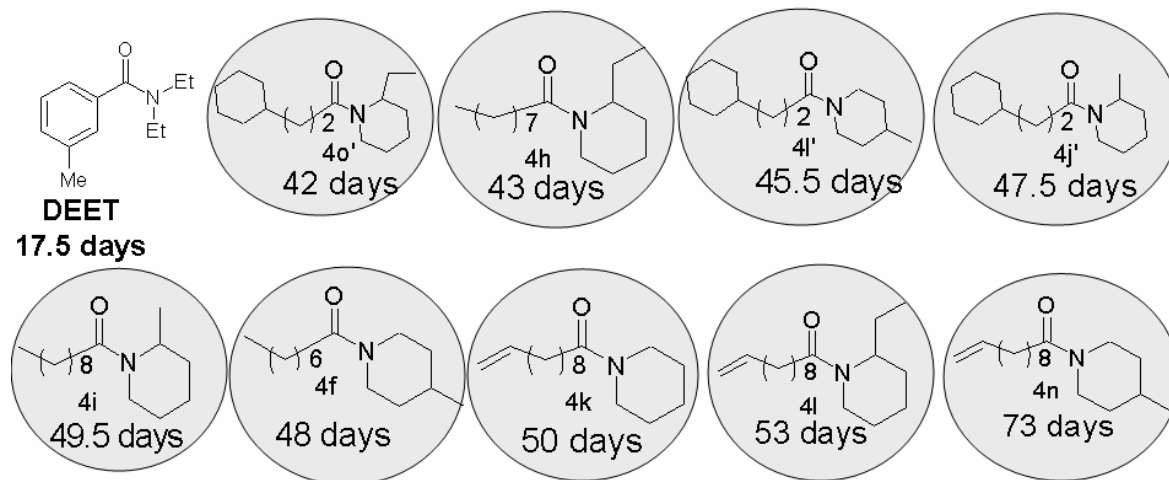


Figure 2. Correlation of protection time with chemical structure for the newly synthesized N-acylpiperidines.

The structures of the most effective compounds and their protection times are compared with DEET in Figure 3. These compounds are now being subjected to toxicity and other tests to see if they can be applied. We have particularly not asked for any patent protection for any of this work in order to make the results available to the whole world as quickly as possible.

Table 1. Average protection time in days

| Compound | | | Average PT | | Compound | | | Average PT | |
|-------------|---|------|--------------------------------|---------------------------------|-------------|--|------|--------------------------------|---------------------------------|
| ID # | R | R' | 25 $\mu\text{mol}/\text{cm}^2$ | 2.5 $\mu\text{mol}/\text{cm}^2$ | ID # | R | R' | 25 $\mu\text{mol}/\text{cm}^2$ | 2.5 $\mu\text{mol}/\text{cm}^2$ |
| <i>DEET</i> | | | 17.5 | 2.5 | | | | | |
| 4a | Me | 2-Me | 2 | 2 | 4a** | 1- <i>c</i> -C ₆ H ₉ | H | 17 | 5 |
| 4b | Et | H | 5 | 4 | 4b** | <i>c</i> -C ₆ H ₁₁ | H | 14 | 8 |
| 4c | Et | 2-Et | 5 | 3 | 4c** | <i>c</i> -C ₆ H ₁₁ | 3-Me | 17 | 6 |
| 4d | <i>n</i> -C ₄ H ₁₃ | 2-Me | 17 | 5 | 4d** | <i>c</i> -C ₆ H ₁₁ | 4-Me | 24.5 | 8.5 |
| 4e | <i>n</i> -C ₄ H ₁₃ | 3-Me | 15.5 | 7.5 | 4e** | <i>c</i> -C ₇ H ₉ (CH ₂) ₂ | H | 35 | 9 |
| 4f | <i>n</i> -C ₇ H ₁₅ | 4-Me | 48 | 8 | 4f** | 1-Me- <i>c</i> -C ₆ H ₁₀ | 3-Me | 12 | 7 |
| 4g | <i>n</i> -C ₇ H ₁₅ | 4-Bn | 13 | 7 | 4g' | 4-Me- <i>c</i> -C ₆ H ₁₀ | 2-Me | 33 | 8.5 |
| 4h | <i>n</i> -C ₈ H ₁₇ | 2-Et | 43 | 9.5 | 4h** | <i>c</i> -C ₆ H ₁₁ | 2-Et | 21.5 | 7 |
| 4i | <i>n</i> -C ₉ H ₁₉ | 2-Me | 49.5 | 8 | 4i** | <i>c</i> -C ₆ H ₁₁ CH ₂ | 2-Me | 29.5 | 7.5 |
| 4j | <i>n</i> -C ₉ H ₁₉ | 4-Me | 41 | 11.5 | 4j** | <i>c</i> -C ₆ H ₁₁ (CH ₂) ₂ | 2-Me | 47.5 | 10 |
| 4k* | CH ₂ =CH(CH ₂) | H | 50 | 13.5 | 4k** | <i>c</i> -C ₆ H ₁₁ (CH ₂) ₂ | 3-Me | 35 | 9 |
| 4l | CH ₂ =CH(CH ₂) | 2-Et | 53 | 9 | 4l' | <i>c</i> -C ₆ H ₁₁ (CH ₂) ₂ | 4-Me | 45.5 | 8 |
| 4m | CH ₂ =CH(CH ₂) | 4-Bn | 8.5 | 8 | 4m' | <i>c</i> -C ₆ H ₁₁ (CH ₂) ₃ | 4-Me | 33 | 3 |
| 4n | CH ₂ =CH(CH ₂) | 4-Me | 73 | 10.5 | 4n' | <i>c</i> -C ₇ H ₉ (CH ₂) ₂ | 2-Et | 40.5 | 8.5 |
| 4o | <i>n</i> -C ₁₀ H ₂₁ | H | 39.5 | 13 | 4o' | <i>c</i> -C ₆ H ₁₁ (CH ₂) ₂ | 2-Et | 42 | 10.5 |
| 4p | <i>n</i> -C ₁₁ H ₂₃ | 2-Me | 14.5 | 5 | 4p' | <i>c</i> -C ₆ H ₁₁ CH ₂ | 4-Bn | 3 | 1.5 |
| 4q | <i>n</i> -C ₁₁ H ₂₃ | 3-Me | 19.5 | 5.5 | 4q' | <i>c</i> -C ₆ H ₁₁ (CH ₂) ₂ | 4-Bn | 12 | 1 |

**Figure 3.** Protection time (25 $\mu\text{mol}/\text{cm}^2$) and structures.

2. Bioconjugates for New Directions in Pharma Research: Applications of Acylbenzotriazoles.

(Research with the participation of P. Angrish, B. E.-D. M. El-Gendy, D. Haase, L. Khelashvili, T. Narindoshvili, S. Tala)

The pharmaceutical industry is in crisis. It is becoming increasingly difficult to find new major drugs. Many of the time-honored strategies for drug research seem to be failing. One direction that is opening up is that of bioconjugates. Bioconjugates can be defined as compounds in which at least two fundamentally different types of organic structure are linked together. Table 2 shows the possibilities considering two out of six compound classes to be linked to each other or to another class. Some of the different types of bioconjugates displayed in Table 2 are very well known, but others have hardly been studied.

Table 2. Bioconjugates for new directions in pharma research

| | Amino Acids | Mono Saccharides | Lipids | Steroids Terpenes | Porphyrins | Biomarkers |
|-------------------|--|-----------------------------------|--|----------------------|----------------------------------|-------------------------|
| Amino Acids | Proteins | Glycoproteins Glycopeptides | Lipoprotein Lipoamino acids Lipopeptides | Peptidyl steroids | Porphyrin-amino acid derivatives | Biomarked amino acids |
| Mono Saccharides | Glycoproteins Glycopeptides | Sugars Starches | Glycolipids Lipopolysaccharide | Steroidal glycosides | Glycoporphyrins | Biomarked carbohydrates |
| Lipids | Lipoprotein Lipoamino acids Lipopeptides | Glycolipids Lipopolysaccharide | Fats | Lipoproteins | Lipoporphyrins | Biomarked lipids |
| Steroids Terpenes | Peptidyl steroids | Steroidal glycosides | Lipoproteins | Higher Terpenoids | — | Biomarked steroids |
| Porphyrins | Porphyrin-amino acid derivatives | Glycoporphyrins | Lipoporphyrins | — | Polyporphyrins | Biomarked porphyrins |
| Biomarkers | Amino acid biomarkers | Carbohydrate biomarkers | Lipid biomarkers | Steroid biomarkers | Porphyrin biomarkers | — |

An easy and often convenient way of linking two structural units together in organic chemistry is acylation. The classical way to affect acylation uses acid chlorides, but there are disadvantages. Recently³ we introduced *N*-acylbenzotriazoles as substitutes for acid chlorides. Several advantages are associated with these reagents:

1. Preparation: rapid one-pot procedures with mild reaction conditions (THF; 20 °C; 2 hr)

$$\text{BtH} + \text{SOCl}_2 \rightarrow [\text{BtSOBt}] + \text{RCO}_2\text{H} \rightarrow \text{RCOBt}; \quad \text{RCO}_2\text{Na} + \text{BtTs} \rightarrow \text{RCOBt} + \text{TsONa}.$$
2. Isolation: easily in crystalline form; high yields and purity without chromatography.
3. Stability: can be weighed out and handled in air, and stored at 20°C for many weeks.

4. Insensitive to water and thus can be used in partly aqueous solution: this allows peptide-coupling using amino acids with free carboxyl groups.
5. Protection not usually required for aliphatic or aromatic -OH, heterocyclic -NH, -SH, or -CONH₂.
6. They are more reactive and more crystalline than acyl imidazoles.
7. The Bt group is readily replaced by N-, S-, O-, and C- nucleophiles.

Acylbenzotriazoles can be applied to classical problems such as the synthesis of “difficult” peptide sequences, one of which is shown in Figure 4.⁴

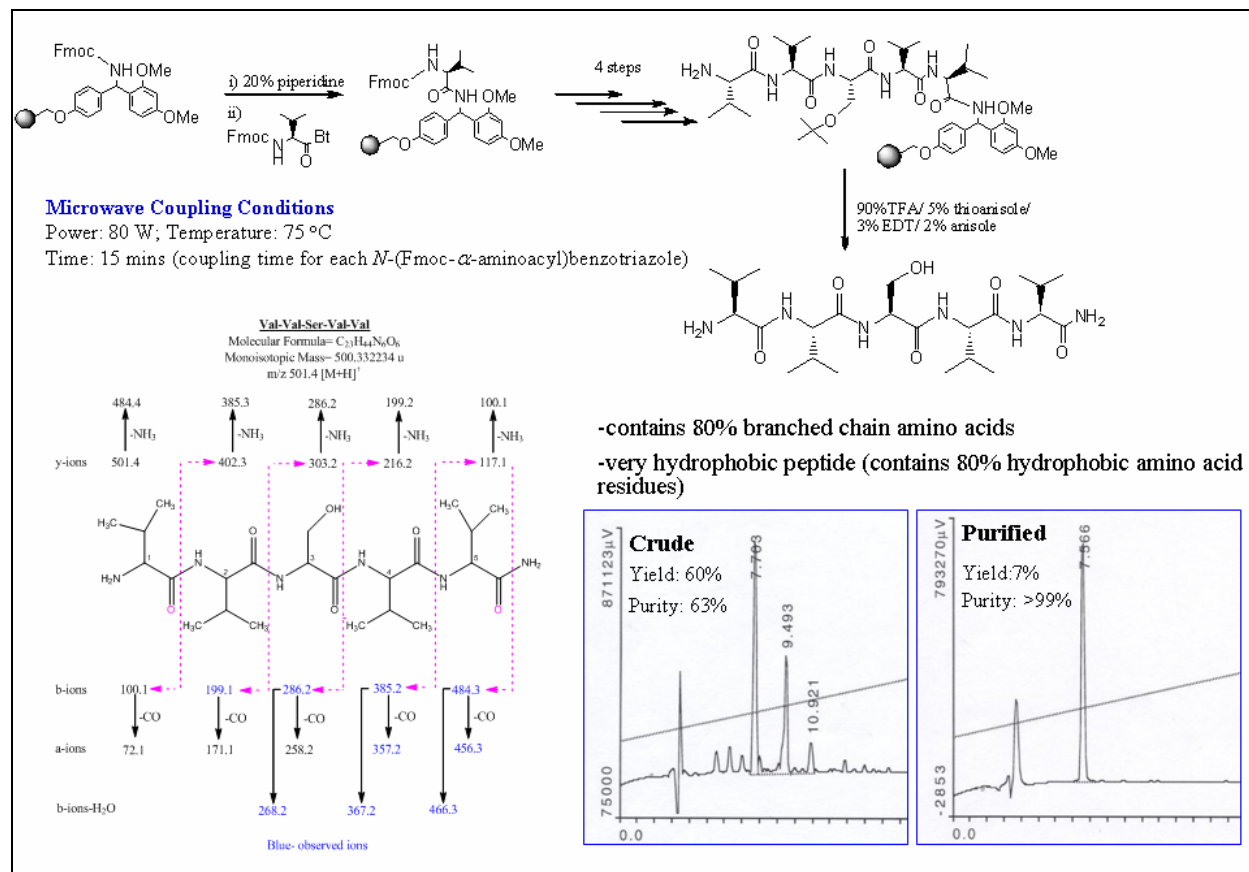


Figure 4. Microwave-assisted solid phase synthesis of a short difficult pentapeptide.⁴

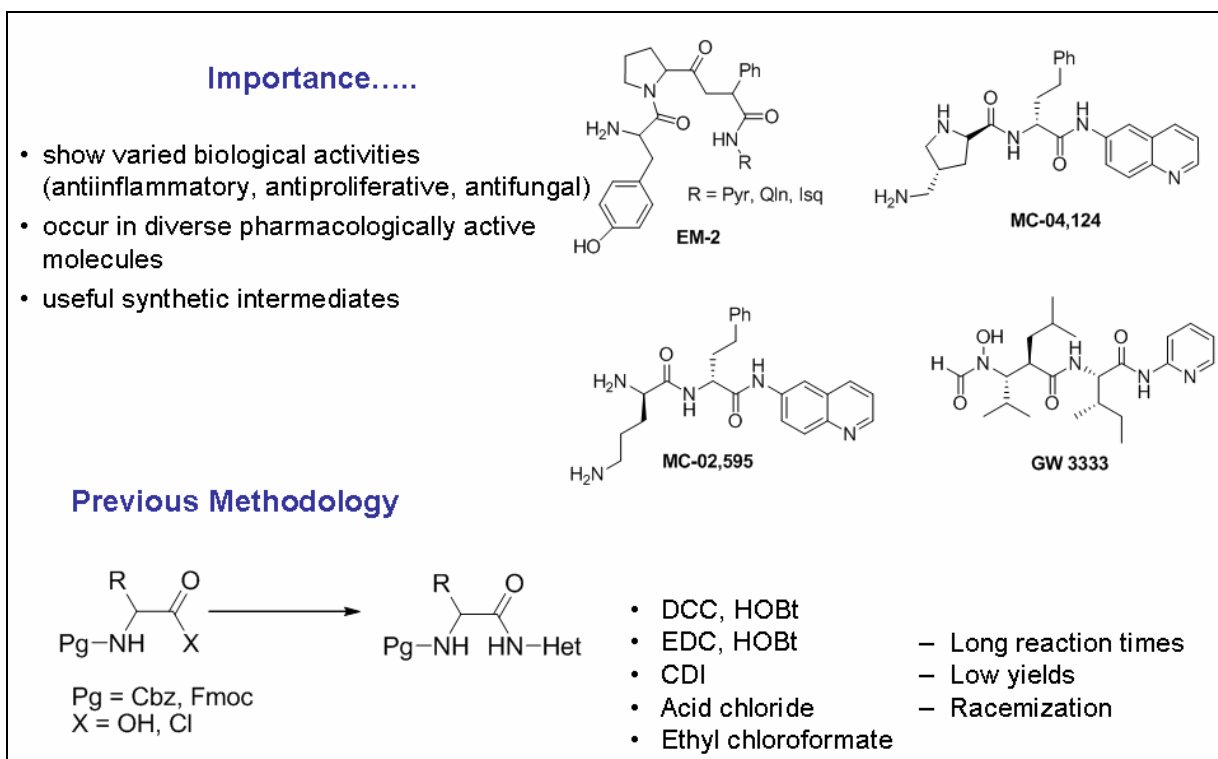


Figure 5. Importance and background for α -aminoacylamino-substituted heterocycles.

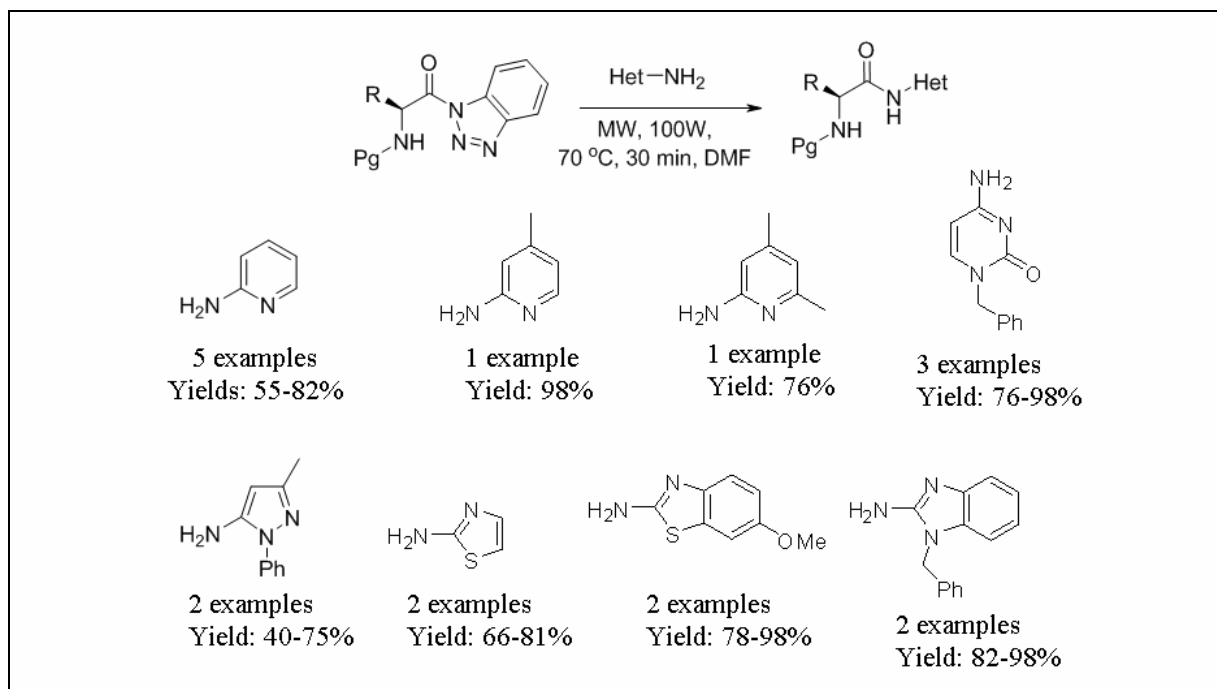


Figure 6. Novel α -aminoacylamino-substituted heterocycles prepared.⁵

Another application⁵ is in the preparation of α -aminoacylamino- substituted heterocycles which show many diverse biological activities. Figure 5 shows some examples of this, comparing the new methodology with the old, and resulting in the preparation of a diverse range of such compounds — additional examples of which are shown in Figure 6.

We have linked amino acids with terpenes through hydroxyl groups, as summarized in Figure 7.⁶ The data in Figure 7 also demonstrate the fact that chirality is preserved in the preparation and reactions of acylbenzotriazoles. Thus *DL*-Phe-*O*-citronellol shows retention times corresponding to each of the two diastereoisomers, whereas the analogous *L*-phenylalanine product shows only one peak.

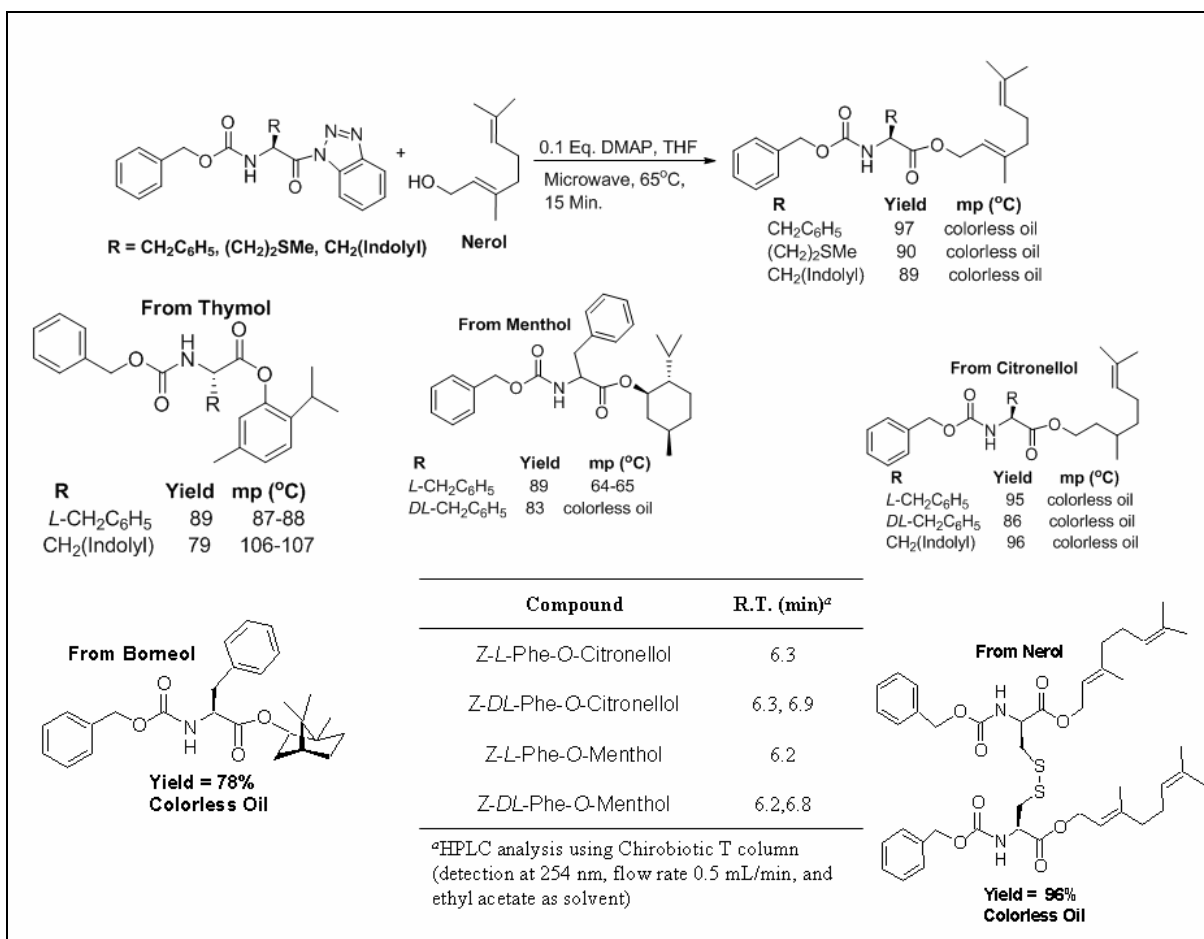


Figure 7. Preparation of *N*-protected (α -aminoacyl)oxy-substituted terpenes.⁶

Similarly amino- acid residues can be linked to steroids: Figure 8 shows some bioconjugates derived from cholesterol linked to a variety of amino acids.⁷

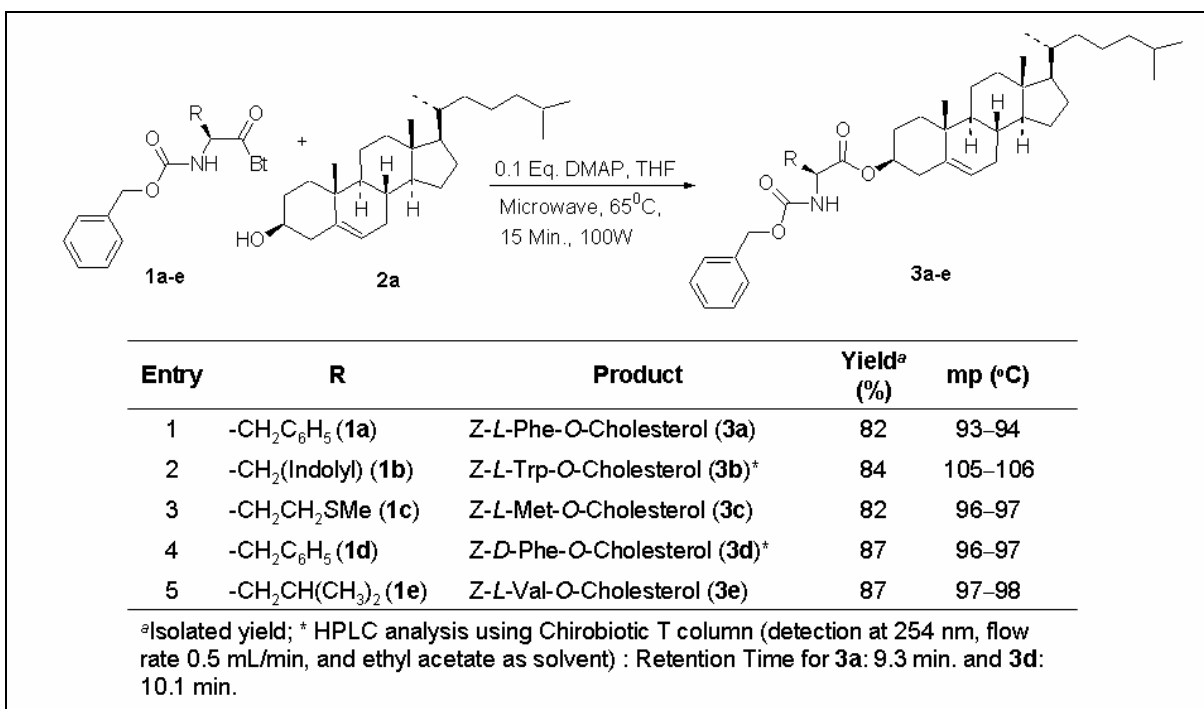


Figure 8. Microwave- assisted synthesis of α -(protected-aminoacyl)oxysteroids.⁷

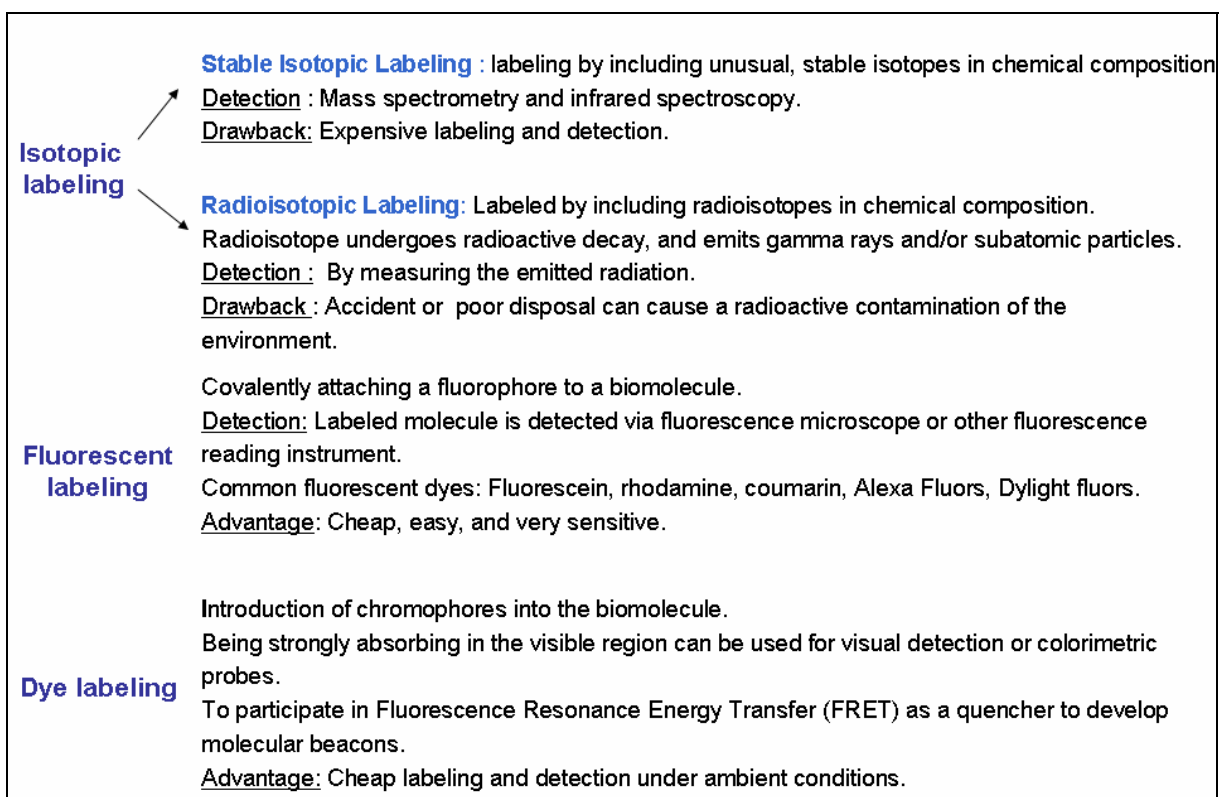


Figure 9. Tagging techniques for molecules.

Tagging has become extremely important for biomolecules (Figure 9).

Azo-dye- labeled *O*-aminoacyl terpenes, -sugars, and -steroids⁸ have been prepared by the new technique, as illustrated in Figure 10.

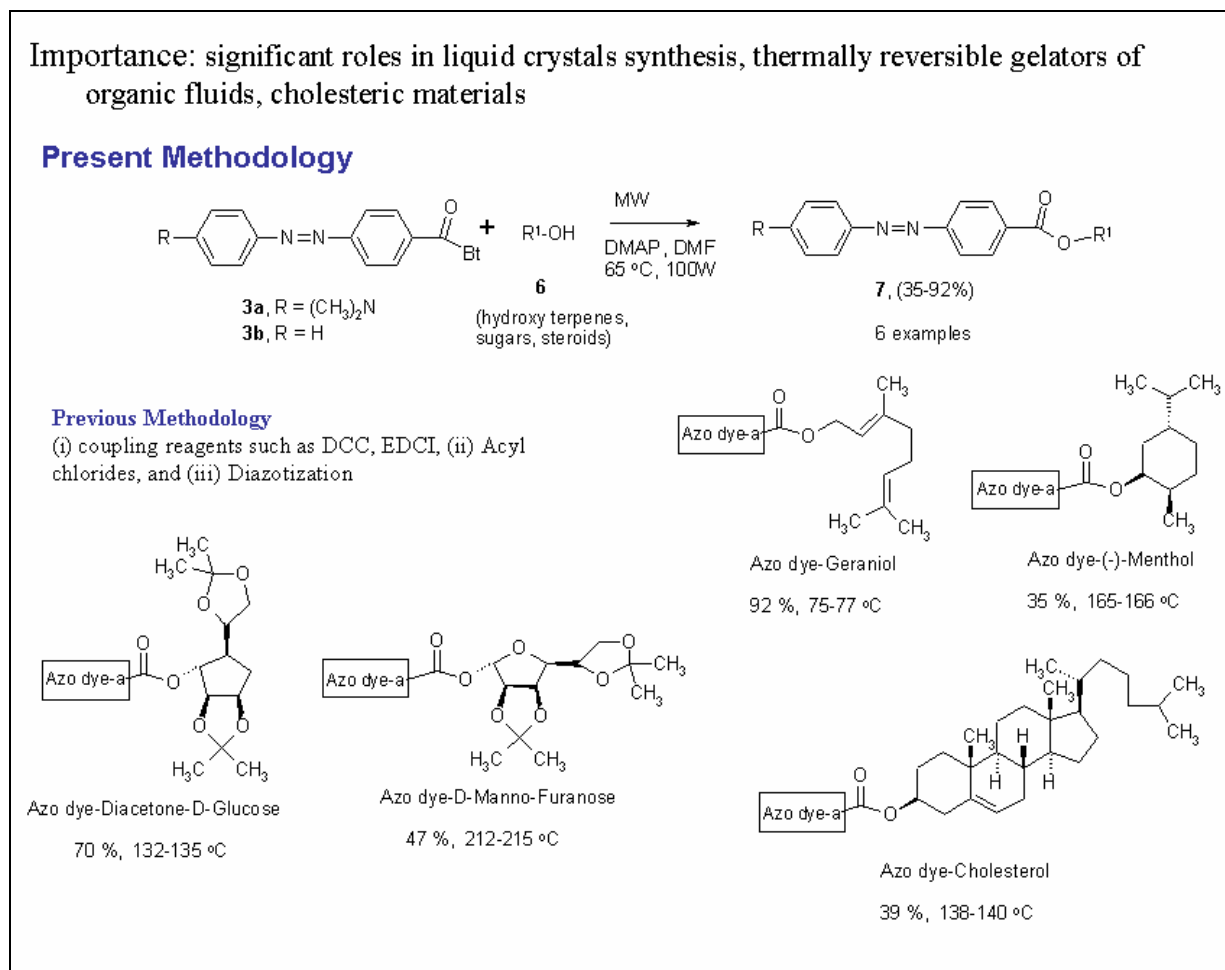


Figure 10. Preparation of azodye labeled terpenes, sugars and steroids.⁸

Fluorescent markers are often even more effective than dye markers. We have shown that coumarin units can be utilized in combination with lysine chemistry to tag a variety of amino acids and peptides⁹ (Figure 11).

Incorporation of a sugar unit confers solubility on such compounds and some examples of this technique¹⁰ are shown in Figure 12.

A series of dye- labeled nucleosides¹¹ is shown in Figure 13.

Similar techniques have been applied to dye- label amino acids and dipeptide amino alcohols,¹² as shown in Figure 14.

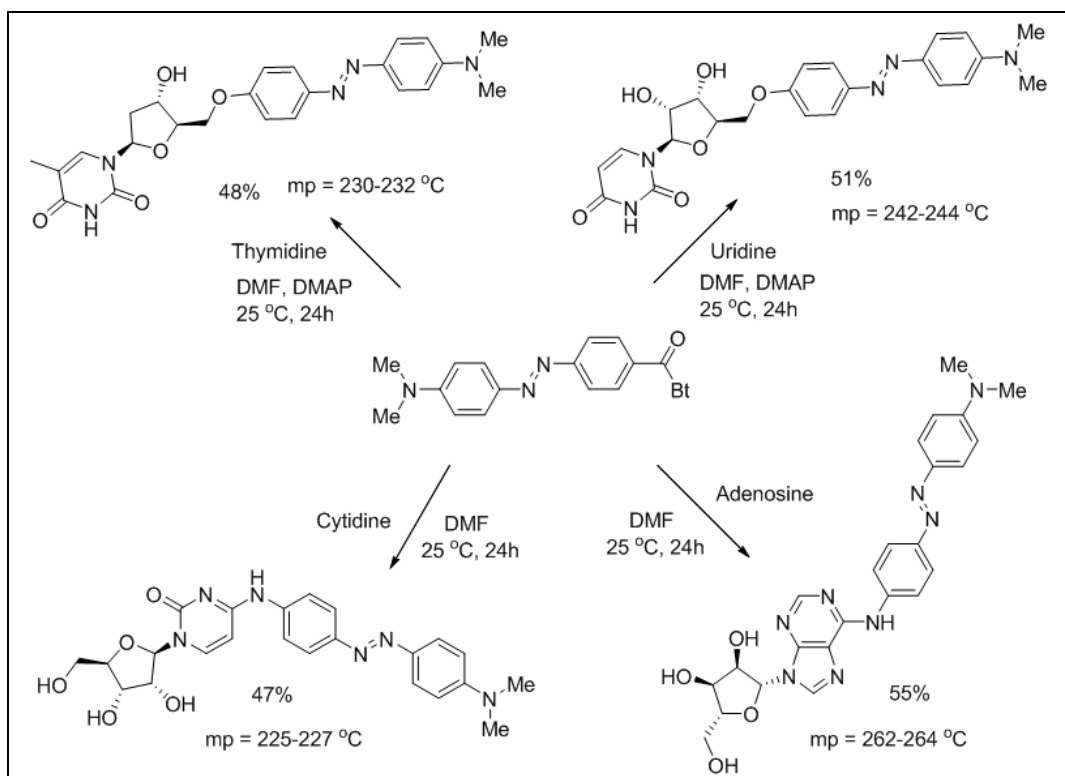


Figure 13. Preparation of dye- labeled nucleosides.¹¹

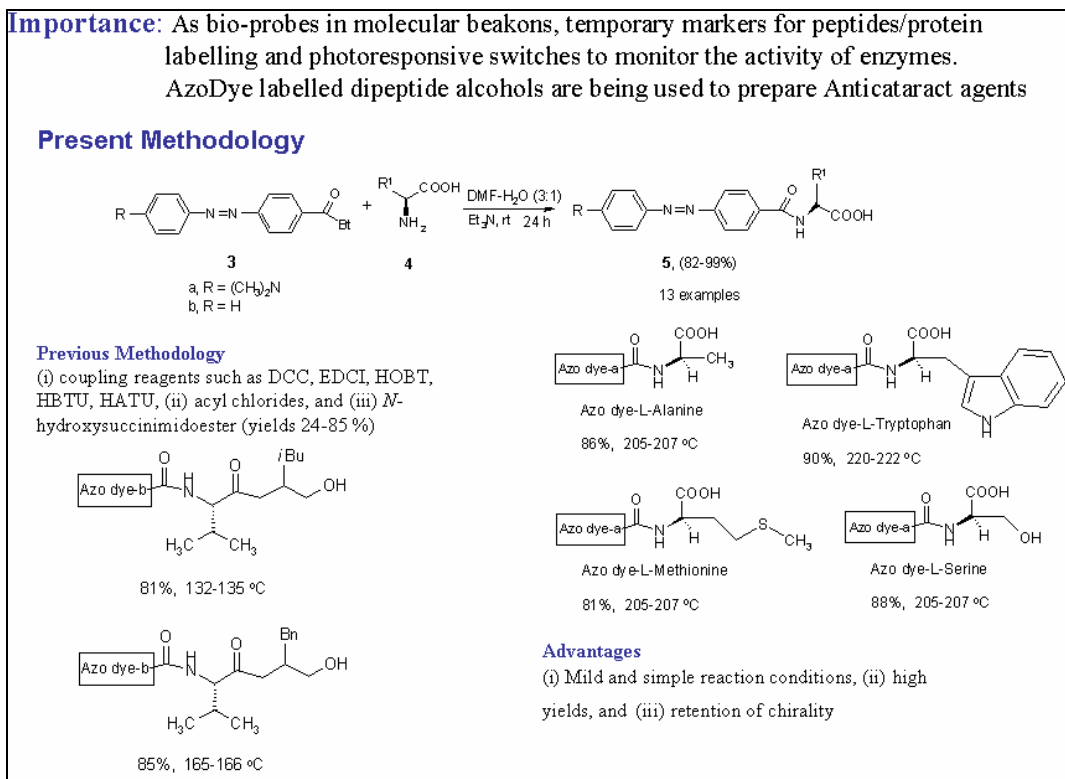


Figure 14. Synthesis of azo-dye- labeled amino acids and dipeptide alcohols.¹²

3. Attempts to help Chemistry in Developing Countries by Innovations in the Publishing and Dissemination of Organic Chemistry Research

The problems faced by libraries in meeting the high subscriptions required by commercial publishers are well known. The subscriptions set by learned societies for access to their publications are also high. This has given rise to the so-called “open access journals” where there is no charge to the reader, but page charges are levied on authors. In these circumstances it was decided eight years ago to launch a new journal with a very different philosophy: it would be free to authors, with no page charges or other fees, and also free to readers with no access or downloading charges. Thus Arkivoc was designed for universal on-demand distribution at no cost to authors or subscribers (see Figure 15).

| Subscription Journals (no charge to authors / Library pays) | Open Access Journals (no charge to reader; pages charged to authors) | Objectives for Arkivoc |
|--|--|---|
| European J. Org. Chem. (\$4933) | Bio. and Med. Chem. | Free to authors No page charges or other fees |
| J. Org. Chem. (\$2500) | Beilstein J. Org. Chem. | Free to readers No access or downloading fees |
| Tetrahedron (\$16,756) | Molecules | |
| Tetrahedron Letters (\$12,204) | | |
| Synthesis (\$1900) | | |
| Synlett (\$1300) | | |
| Designed for universal on-demand distribution at no cost to authors or subscribers | | |
| <ol style="list-style-type: none"> 1. On WWW electronically - free for all to access 2. Web edition formatted in pages to allow downloading and binding into hard copy - for free 3. Hard copy edition also available to send to deposit library and by subscription - for payment 4. Is kept in perpetuity on several servers worldwide | | |

Figure 15. Publishing of chemistry in 2008.

Many of the standard publishing procedures are used in Arkivoc, as detailed in Figure 16, but a major difference is that the “Control Board” which runs Arkivoc is unpaid. The composition of the Control Board as of spring 2008 is shown in Figure 17.

1. Instructions for authors including template available at www.arkat-usa.org
2. Authors contact Coordinating Editor by e-mail with MS title, authors, brief abstract
3. Coordinating Editor designates MS reference number, Scientific Editor, and referees
4. MS sent to Referees by e-mail
5. Referees send reports to Scientific Editor by e-mail
6. Scientific Editor makes decision as to accept the paper as it is, to ask author to amend manuscript, to seek additional Referee comments, or to reject paper
7. Author revises MS to satisfaction of Scientific Editor
8. Scientific Editor corresponds with authors until manuscript is accepted or rejected
9. Accepted manuscript sent to Publishing Editor, who arranges for technical editing and sending in for posting on the web

Figure 16. Submitting, refereeing and editing of manuscripts in ARKIVOC.

Referee Assigners: E. Anders (U. Jena, Germany), A. J. Boulton (also Coordinating Ed, UEA, UK), W. Dolbier (U. Florida, USA), B. Wakefield (U. Leeds, UK)

Scientific Editors: M. Begtrup (Roy. Sch. Pharmacy, Denmark), G. Cirrincione (U. Palermo, Italy), M. A. Iglesias- Arteaga (Nat U Mexico), H. Ila (Indian Inst Sci Kanpur), J. Joule (U. Manchester, UK), A. Kotali (U. Thessaloniki, Greece), P. Krapcho (U. Vermont, USA), B. Maes (U. Antwerp, Belgium), A. Marchand (Coral En, Inc.), R. Muthyala (U. Minnesota, USA), C. Ramsden (Keele U., UK), J. Schantl (U. Innsbruck, Austria), C. Stevens (Ghent U., Belgium), A. Waring (U. Birmingham, UK), V. Zhdankin (U. Minnesota, USA)

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Publishing Editor: E. Scriven (U. Florida, USA)

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Figure 17. ARKIVOC organization: Control Board as of March 2008.

We also have a very large Editorial Board of Referees. This has recently been extended significantly, and we now have close to 1,000 members. One difference from most editorial boards of referees is that we have about half our members from outside Western Europe, North America or Japan (see Figures 18 and 19) for representative lists.

During the eight years of existence, Arkivoc has progressed significantly in the number of manuscripts received and published. The number of visitors to its website has also increased dramatically; we now have about 100,000 visitors per month who make more than a million hits per month on the website.

| | | |
|---|---|--|
| Austria: U. Brinker, J. Froehlich, P. Gaertner, N. Haider, C. O. Kappe, M. D. Mihovilovic, T. Rosenau, M. Schnürch, P. Stanetty | G. Doddi, G. Favaro, S. Florio, L. Forlani, P. Fornasiero, F. Fringuelli, A. Gasco, G. La Manna, P. Linda, A. Maia, G. Musumarra, F. Naso, R. Noto, M. Peruzzini, M. Pulici, G. Scorrano, G. Sindona, D. Spinelli, L. Troisi, E. Valentin, P. Zanirato | Sandford, J. Shorter, K. Smith, E. Thomas, J. Walton |
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| Finland: K. Klika, M. Oivanen, K. Pihlaja, E. Sievänen | Portugal: P. Almeida, A. Lobo, C. Oliveira, A. Oliveira-Campos, A. Silva | |
| France: C. Amatore, G. Balme, G. Bashiardes, J. Bazureau, T. Besson, P. Bonnet, P. Dauban, J. Fehrentz, J. Finet, C. Gerardin, G. Kirsch, A. Lattes, C. Laurence, C. Le Roux, J. Majoral, M. Malacria, J. Mattalia, M. Medebielle, G. Queguiner, C. Roussel, D. Sinou, F. Terrier, G. Vo-Thanh | Spain: B. Abarca, M. Alvarez, J. Alvarez-Builla, M. Amat, J. Aurrecochea, F. Aznar, J. Barluenga, J. Fernández Bolaños, J. Bosch, L. Castedo, C. Cativiela, R. Claramunt, D. Diez, E. Díez-Barra, J. Elguero, B. Fontaniella, V. Gotor, C. Lopez, J. Marquet Cortes, J. Menendez, P. Merino, P. Molina, M. Montiel, F. Palacios, C. Palomo, M. Parra, J. Plumet, E. Roman, R. Suau, A. Tárraga, T. Torroba, M. Yus | |
| Germany: E. Anders, M. Arend, T. Balaban, R. Beckert, P. Czerney, U. Dietrich, D. Edelmann, W. Huebsch, H. Ihmels, K. Jug, W. Kantlehner, K. Krohn, T. Kurz, H. Martin, P. Metz, A. Mishra, F. Montforts, S. Nerdinger, P. Rademacher, N. Risch, M. Vogel, H. Wamhoff, E. Wuerthwein | Sweden: J. Bergman, S. Grivas, T. Janosik, C. Linde, J. Stawinski | |
| Ireland: R. Butler, M. Casey, D. Gilheany, A. Maguire, R. O'Ferrall, H. Moynihan | Switzerland: H. Heimgartner, C. Jefford | |
| Italy: A. Almerico, A. Andreani, L. Angiolini, O. Attanasi, M. Aversa, U. Azzena, E. Baciocchi, R. Ballini, M. Catellani, L. Ceraulo, F. Cermola, G. Cevasco, U. Chiacchio, S. Chimichi, G. Cirrincione, M. Costi, G. Daidone, R. Dalpozzo, M. D'Auria, P. Maria, A. Degli'Innocenti, G. Desimoni, | United Kingdom: K. Afarinkia, A. Aitken, D. Allen, S. Allin, R. Bedford, S. Bew, R. Chambers, R. Coombes, M. Cronin, A. de Silva, I. Eggleston, C. Frost, L. Harwood, H. Heaney, W. Horspool, H. Hudson, K. Jones, R. Jones, J. Joule, N. Karodia, D. Kelly, W. Kerr, P. Knowles, B. Linclau, S. Marsden, H. Maskill, T. McKervey, C. Moody, P. Murphy, P. Page, G. Pattenden, M. Plate, C. Ramsden, D. Rees, C. Reese, G. | |

Figure 18. ARKIVOC Editorial Board of Referees: members from W. Europe, N. America, Japan.

| | |
|---|---|
| Argentina: Alicia Baldessari, Maria T. Baumgartner, Graciela Yolanda Buldain, Rita Hoyos de Rossi, Teodoro S. Kaufman, Graciela Y. Moltrasio, Norma Sbarbati Nudelman, Alicia Penenory, Adriana B. Pierini, Julio C. Podesta, Javier Ramirez, Roberto A. Rossi, Juana J. Silber, Patricia Vazquez | Valentina K. Yu |
| Australia: Roger Bishop, David Black, Robert T. C. Brownlee, Darren Cundy, Chris J. Easton, Robin G. F. Giles, Stephen Glover, Andrew Hughes, Bill Kitching, Patrick Perlmutter, Jack Ryan, Paul Savage, Jamie Simpson, Bela Ternai, Curt Wentrup, Jonathan M. White | Korea: SouthHyun-Joon Ha, Yogesh R. Jorapur, Byeang Hyeon Kim, Jae Nyoung Kim, Seokjoon Lee |
| Bangladesh: M. Giasuddin Ahmed | Kuwait: Nouria Al-Awadi |
| Belarus: Alexander V. Baranovsky | Latvia: Pavel Arsenyan, Gunars Duburs, Edmunds Lukevics |
| Brazil: Wilhelm Josef Baader, Ivan P. de A. Campos, Paulo Cesar de Jesus, Luiz Carlos Dias, Ronaldo Pilli, Jose Augusto Rosario Rodrigues, Marcus M. Sa, Dulce Helena Siqueira Silva, Marcos Souza, Blanka Wladislaw | Lebanon: M. Haddadin |
| Bulgaria: Ivan G. Binev, Vladimir Dimitrov, Ivan G. Pojarlieff | Libya: George Y. Sarkis |
| Chile: Julio Belmar Mellado, Bruce Cassels, Marcos Caroli Rezende | Malaysia: Zuriati Zakaria |
| China: Shujiang Tu, Mei-Xiang Wang, Z. F. Xi, Deqing Zhang, Yongmin Zhang, Zhan-Hui Zhang, Guisen Zhao, Jian-Ping Zou | Mexico: Armando Ariza-Castolo, Carlos M. Cerda, Gabriel Cuevas, Francisco Delgado, Guillermo Delgado, Norberto Farfan, Felipe J. González, Barbara Gordillo, Eusebio Juaristi, Roberto Martinez, Rachel Mata, René Miranda, |
| Croatia: Mirjana Eckert-Maksic, Bono Lucic, Kata Mlinaric-Majerski, Marin Roje, Nenad TrinajsticHrvoj Vancik | Moldova: Fliur Macaev |
| Cuba: Daniel Garcia-Rivera, Carlos Perez, Margarita Suarez Navarro | New Zealand: Jim Coxon, Brian Halton, Gordon Rewcastle, Peter J. Steel |
| Cyprus: Panayiotis A. Koutentis | Poland: Ryszard Bodalski, Dariusz Bogdal, Jacek Brzezinski, Marek Chmielewski, Zofia Dega-Szafran, Tomasz Janecki, Janusz Jurczak, Andrzej Kapturkiewicz, T. Marek Krygowski, Jaroslaw Lewkowski, Barbara Malawska, Roman Mazurkiewicz, Jan Michalski, Jacek W. Morzycki, Wieslaw Oleszek, Jerzy Suwinski, Miroslaw Szafran, Piotr Tomasik |
| Czech Republic: Stanislav Radl | Puerto Rico: John A. Soderquist |
| Egypt: Ashraf A. Aly, El Sayed H. El Ashry, Ayman El-Faham, A. S. Hamad Elgazwy, Mohamed Helmy Elnagdi, A. Heshmat Moustafa, | Romania: Vasile N. Bercean, Florea Dumitrascu, Petru Filip, Florentina Georgescu, Ionel I. Mangalagiu, Vasile I. Parvulescu, Alexandru C. Razu,s |
| Estonia: Mati Karelson, Uno Mäeorg, Uko Maran | Russia: Vadim Annenkov, Eugene Babaev, Shainyan Bagrat, Vasilij Bakulev, Leonid Belen'kii, Irina Beletskaya, Konstantin Bryliakov, Andrei Ershov, Nina Gusarova, Nadezhda Itsikson, Andrey Ivanov, Rem Kostyanovsky, Mikhail Kuznetsov, Alevtina Medvedeva, Nina Nedolya, Vladimir Ostrovskii, Anatolii Pomogailo, Alexander Pozharskii, Valery Traven, Alexey Trofimov, Sergei Vasilevsky, Natalia Vlasova, Mikhail Voronkov, Nikolai Zefirov |
| Greece: John K. Gallos, Michael Orfanopoulos, Anastasios Varvoglis, GeorgeVarvounis | Saudi Arabia: Ashraf Ghanem |
| Hong Kong: Wei-Min Dai | Serbia: Ivan Juranic, Rade Markovic |
| Hungary: Ferenc Fülöp, Gyorgy Hajos, Kalman Hideg, Albert Lévai, Peter Molnar, József Nyitrai | Slovakia: Lubor Fišera, Alzbeta Krutosikova, Viktor Milata, Ladislav Petrus |
| India: Rajkumar Bansal, Deevi Basavaiah, Chinnappan Baskar, D. Subhas Bose, Ramesh Chandra, S. Chandrasekhar, Biswanath Das, Asish De, Rina Ghosh, Subrata Ghosh, Hiriyakkanavar Ila, Javed Iqbal, Sanjay Jain, N. N. Joshi, H. Junjappa, Srinivas Kantevari, Sambasivarao R. Kotha, Devinder Kumar, G. Kumaraswamy, Alka Mital, D. K. Mohapatra, Satbir Mor, Ganesh Pandey, Virinder S. Parmar, Mariappan Periasamy, S. Perumal, Om Prakash, Thingalur K. Raja, A. V. Rama Rao, B. C. Ranu, Diwan S. Rawat, Anil Kumar Saxena, G. V. M. Sharma, M. S. Shashidhar, Vishwakarma Singh, Neelima Sinha, A. Srikrishna, G. S. R. Subba Rao, G. Sundararajan, Yashwant D. Vankar, Shirodkar Prabhakar Yeshawant, IranMohammad Reza Islami, Kazem Saidi, Hassan Sheibani, Issa Yavari, Mohammad Ali Zolfigol | Slovenia: Jernej Iskra, Danijel Kikelj, Slovenko Polanc, Branko Stanovnik, Jurij Svete |
| Israel: Eli Breuer, Alfred Hassner, Shmaryahu Hoz | South Africa: Jaco Breytenbach, Ivan Green, Roger Hunter, Perry T. Kaye, Joseph P. Michael |
| Jamaica: Yvette Jackson | Taiwan: Long-Yong Chiang, Reuben Jih-Ru Hwu, Tien-Yau Luh, Chin-Kang Sha |
| Kazakhstan: Svetlana Achmetshakirovna Vizer, | Trinidad and Tobago: Gurdial Singh |
| | Turkey: Esin Aki-Sener, K. Husnu Can Baser, Sevim Bilgic, Nihat Celebi, Necdet Coskun, Mustafa Güllü, Cemil Ibis, Cemil Ogretir, Ismail Ozdemir, Turan Ozturk, Bilge Sener, Süleyman Servi, Birsen Tozkoparan, Lemi Turker |
| | Ukraine: Dmitriy Volochnyuk |
| | Uruguay: Hugo Cerecetto, Horacio Heinzen |
| | Uzbekistan: Andrey A. Toropov |
| | Venezuela: Franklin Vargas |

Figure 19. ARKIVOC Editorial Board of Referees: members from outside W. Europe, N. America, Japan.

An important part of the philosophy of ARKIVOC is to honor chemists from around the world. Distinguished scientists, including chemists, have long been honored in their own countries and a small number, mainly from a few Western developed countries, are honored internationally. Arkivoc has chosen to recognize chemists from around the world and, so far we have produced 85 commemorative issues (Figure 20).

| Country | Year | Chemist |
|-----------|------|-------------------|
| Argentina | 2003 | Rossi |
| | 2003 | Ruveda |
| | 2005 | Ledekremer |
| Austria | 2001 | Sauter |
| Australia | 2001 | Cameron |
| | 2004 | Rickards |
| Belgium | 2003 | Hoomaert |
| | 2007 | Krief |
| Belarus | 2008 | Kulinkovich |
| Bulgaria | 2009 | Conference |
| Botswana | 2007 | Abegaz |
| Brazil | 2004 | Gottlieb |
| Canada | 2001 | Tee |
| | 2002 | Muchowski |
| | 2009 | Sorensen |
| China | 2003 | Huang |
| | 2004 | Yuan |
| Croatia | 2002 | Sunko |
| Finland | 2001 | Pihlaja |
| | 2009 | Lonnberg |
| France | 2006 | Lattes |
| | 2008 | Queguiner |
| | 2008 | Solladie- Cavalle |
| Germany | 2004 | Krohn |
| | 2007 | Adam |
| | 2007 | Anders |
| | 2007 | Tietze |
| Greece | 2003 | Varvoglis |

| Country | Year | Chemist |
|---------|-------------|------------------|
| Holland | 2004 | Zwanenburg |
| Hungary | 2003 | Bernath |
| | 2004 | Antus |
| | 2008 | Szantay |
| India | 2001 | Govindachari |
| | 2002 | Kessar |
| | 2003 | Chatterje |
| | 2004 | Sukh Dev |
| | 2005 | Narismhan |
| | 2005 | Anand |
| | 2005 | Rao |
| 2005 | Swaminathan | |
| Israel | 2001 | Hassner |
| Italy | 2002 | Spinelli |
| | 2004 | Tortorella |
| | 2006 | Bartoli |
| | 2006 | ICHC Conf. |
| | 2006 | Med. Chem. Conf. |
| | 2009 | Vivona |
| Japan | 2003 | Fukumoto |
| | 2009 | Tomoya |
| Kuwait | 2008 | Al- Awadi |
| Latvia | 2006 | Lukevics |
| Mexico | 2003 | Regional Issue |
| | 2005 | Juaristi |
| | 2008 | Contreras |
| NZ | 2006 | Coxon |
| Norway | 2001 | Undheim |

| Country | Year | Chemist |
|-----------|------------|----------------|
| Pakistan | 2007 | Rahman |
| Poland | 2004 | Makosza |
| | 2007 | Epstajn |
| Romania | 2002 | Nenitzescu |
| | 2005 | Balaban |
| Russia | 2001 | Voronkov |
| | 2003 | Trofimov |
| | 2004 | Chupakhin |
| | 2004 | Konovlov |
| | 2005 | Minkin |
| | 2005 | Zefirov |
| | 2008 | Beletskaya |
| 2009 | Pozharskii | |
| Slovakia | 2005 | Fisera |
| Slovenia | 2001 | Tisler |
| | 2003 | Stanovnik |
| S. Africa | 2002 | Bull |
| Sweden | 2008 | Norin |
| Ukraine | 2005 | Regional Issue |
| UK | 2000 | Jones |
| | 2000 | Meth- Cohn |
| | 2002 | Lloyd |
| | 2002 | Rees |
| USA | 2001 | Abramovitch |
| | 2001 | Thyagaragan |
| | 2002 | Karabatsos |
| | 2002 | Padwa |
| | 2003 | Shine |
| 2007 | Joullé | |

Figure 20. 2000- 2009: 85 Chemists honored from 33 countries.

Our impact factor has risen steadily and has now overtaken that of several other journals (Figure 21); while it is still well below that of the best journals in organic chemistry, we hope that the increase in our impact factor will continue.

We have also tried to improve contacts between chemists worldwide by running an annual heterocyclic and synthetic conference at the University of Florida. At this conference we offer a dozen plenary lectures from some of the world's most distinguished chemists (Figure 22) together with short courses on various aspects of organic chemistry, 20-30 invited lectures, some 60 posters, and a full social program. Figure 23 shows the program for the 2008 conference.

| Year | Arkivoc | Molecules | JHetC | Heterocycles | Synthetic Comm. | Synthesis | JOC |
|------|---------|-----------|-------|--------------|-----------------|-----------|-------|
| 2000 | - | 0.182 | 0.781 | 1.015 | 0.828 | 2.193 | 3.689 |
| 2001 | - | 0.223 | 0.746 | 0.970 | 0.912 | 1.985 | 3.280 |
| 2002 | - | 0.408 | 0.701 | 1.045 | 0.802 | 2.201 | 3.217 |
| 2003 | 0.392 | 0.911 | 0.711 | 1.082 | 0.853 | 2.074 | 3.297 |
| 2004 | 0.418 | 0.676 | 0.814 | 1.064 | 0.965 | 2.203 | 3.462 |
| 2005 | 0.694 | 1.113 | 0.735 | 1.070 | 0.860 | 2.401 | 3.675 |
| 2006 | 0.800 | 0.841 | 0.776 | 1.077 | 1.001 | 2.333 | 3.790 |
| 2007 | 1.253 | 0.940 | 0.813 | 1.066 | 0.977 | 2.257 | 3.959 |

Figure 21. ARKIVOC Impact Factor.

| Name | Year | Country | Name | Year | Country | Name | Year | Country | Name | Year | Country |
|---------------|------|---------|---------------|--------|---------|--------------|------|---------|--------------|------|----------|
| E. Anders | 2002 | Germany | V. Farina | 2008 | Belgium | G. Mehta | 2007 | India | J. Schwarz | 2006 | USA |
| J. Armstrong | 2004 | USA | J. Froehlich | 2000 | Austria | N. Meanwell | 2007 | USA | E. Scriven | 2000 | USA |
| J. Backval | 2008 | Sweden | G. Fu | 2006 | USA | B. Maryanoff | 2006 | USA | J. Sisko | 2002 | USA |
| J. Bakke | 2003 | Norway | A. Furstner | 2005/7 | Germany | G. Mehta | 2006 | India | B. Snider | 2007 | USA |
| J. Barluenga | 2002 | Spain | B. Ganem | 2006 | USA | A. Meijere | 2008 | Germany | V. Snieckus | 2000 | Canada |
| M. Begtrup | 2005 | Denmark | G. Gribble | 2003 | USA | O. Meth-Cohn | 2000 | UK | B. Stanovnik | 2001 | Slovenia |
| J. Bergman | 2007 | Sweden | R. Grigg | 2002 | UK | V. Minkin | 2001 | Russia | G.S.R. Rao | 2003 | India |
| I. Beletskaya | 2002 | Russia | R. Grubbs | 2003 | USA | M. Mitchell | 2001 | USA | R. Taylor | 2006 | UK |
| S. Benner | 2001 | USA | K. Hafner | 2000 | Germany | G. Molander | 2001 | USA | T. Tidwell | 2004 | Canada |
| S. Blechert | 2003 | Germany | S. Hecht | 2003 | USA | C.J. Moody | 2007 | UK | L. Tietze | 2004 | Germany |
| N. Bodor | 2002 | USA | P. Hodgson | 2005 | UK | G. Molander | 2005 | USA | B. Trofimov | 2000 | Russia |
| D. Boger | 2005 | UK | P. Jacobi | 2008 | USA | T. Mukaiyama | 2000 | Japan | B. Trost | 2006 | USA |
| M. Brimble | 2005 | NZ | G. Johnson | 2002 | USA | G. Newkome | 2001 | USA | J. Vollhardt | 2007 | USA |
| J. Bristol | 2007 | USA | J. Joule | 2005 | UK | P. Ornstein | 2004 | USA | S. Volante | 2006 | USA |
| M. Butters | 2003 | UK | C. Kappe | 2008 | Austria | A. Padwa | 2004 | USA | S. von Unge | 2003 | Sweden |
| D. Comins | 2000 | USA | J. Kiely | 2000 | USA | G. Pattenden | 2008 | UK | J. Stoddart | 2007 | USA |
| P. Canfalone | 2001 | USA | S. King | 2008 | USA | W. Pearson | 2002 | USA | A. Whittle | 2007 | UK |
| D. Curran | 2002 | USA | P. Knochel | 2007 | Germany | N. Petasis | 2002 | USA | S. Weinrab | 2008 | USA |
| A. Czarnik | 2004 | USA | S. Kobayashi | 2003 | Japan | G. Queguine | 2004 | France | P. Wipf | 2008 | USA |
| N. Kimpe | 2003 | Belgium | H. Kroto | 2008 | USA | A.V. Rao | 2005 | India | P. Wuts | 2002 | USA |
| M. deLong | 2005 | USA | P. Lam | 2008 | USA | C. Ramsden | 2001 | UK | Y. Yamamoto | 2001 | Japan |
| S. Denmark | 2005 | USA | R. Larson | 2007 | USA | C. Rees | 2004 | UK | H. Yamamoto | 2006 | USA |
| W. Dolbier | 2006 | USA | R. Larock | 2005 | USA | H. Reissig | 2007 | Germany | M. Yus | 2003 | Spain |
| S. Dondoni | 2000 | Italy | J. Macor | 2004 | USA | M. Reetz | 2006 | Germany | T. Zhang | 2008 | USA |
| P. Dunn | 2001 | UK | M. Makosza | 2001 | Poland | J. Sanders | 2004 | UK | | | |
| J. Ellman | 2004 | USA | M. Martinelli | 2000 | USA | J. Schantl | 2002 | Austria | | | |

Figure 22. Plenary lectures at Flohet, 2000-2008.

| Conference Program | | Registration Fees | | | | | | | | | | | | | | | | | | | |
|---|-----------|--|--|--|----------|----------|----------------|-----------|----------|----------------------------------|---------|---------|---|----------|----------|--|---------|---------|--|---------|---------|
| Sunday, March 9, 2008 6:00 pm-9:00 pm Registration and Welcome Reception 6:30 pm Buffet Supper | | <table border="1"> <thead> <tr> <th></th> <th>Standard</th> <th>Academic</th> </tr> </thead> <tbody> <tr> <td>Conference Fee</td> <td>\$1200.00</td> <td>\$600.00</td> </tr> <tr> <td>Conference Dinner/Wine Reception</td> <td>\$50.00</td> <td>\$50.00</td> </tr> <tr> <td>Guest pass for Flohet-9 Accompanying Persons (per person)</td> <td>\$250.00</td> <td>\$250.00</td> </tr> <tr> <td>Copy of "Handbook of Heterocyclic Chemistry"</td> <td>\$85.00</td> <td>\$85.00</td> </tr> <tr> <td>Specimen Hard copy of an Arkivoc Issue</td> <td>\$15.00</td> <td>\$15.00</td> </tr> </tbody> </table> <p><i>Academic rates apply to full-time employees at an undergraduate teaching institution</i></p> <p>For inquiries: Contact Vicki Tyson, Conference Organizer E-mail: flohet@arkat-usa.org Tel: (001) 386-684-9303 or (001) 352-316-0208 Fax: (001) 386-684-1433 or (001) 352-392-9199</p> | | | Standard | Academic | Conference Fee | \$1200.00 | \$600.00 | Conference Dinner/Wine Reception | \$50.00 | \$50.00 | Guest pass for Flohet-9 Accompanying Persons (per person) | \$250.00 | \$250.00 | Copy of "Handbook of Heterocyclic Chemistry" | \$85.00 | \$85.00 | Specimen Hard copy of an Arkivoc Issue | \$15.00 | \$15.00 |
| | Standard | Academic | | | | | | | | | | | | | | | | | | | |
| Conference Fee | \$1200.00 | \$600.00 | | | | | | | | | | | | | | | | | | | |
| Conference Dinner/Wine Reception | \$50.00 | \$50.00 | | | | | | | | | | | | | | | | | | | |
| Guest pass for Flohet-9 Accompanying Persons (per person) | \$250.00 | \$250.00 | | | | | | | | | | | | | | | | | | | |
| Copy of "Handbook of Heterocyclic Chemistry" | \$85.00 | \$85.00 | | | | | | | | | | | | | | | | | | | |
| Specimen Hard copy of an Arkivoc Issue | \$15.00 | \$15.00 | | | | | | | | | | | | | | | | | | | |
| Monday, March 10, 2008 8:00 am Registration 8:25 am Opening Remarks 8:30 am-10:10 am Plenary Lectures 1 and 2 9:30 am Excursion for Accompanying Persons 10:10 am-10:40 am Coffee Break 10:40 am-12:20 pm Alternatives: (a) Short Course I; or (b) Invited Lectures 1-4; or (c) Invited Lectures 5-8; Lunch 12:20 pm-1:40 pm Alternatives: (a) Short Course II; or (b) Invited Lectures 9-12; or (c) Invited Lectures 13-16; 1:40 pm-3:20 pm Coffee Break 3:20 pm-3:50 pm Plenary Lectures 3 and 4 3:50 pm-5:30 pm Poster Session 5:30 pm-9:00 pm Buffet Supper 6:30 pm-8:00 pm | | Registration and Payment Three easy ways to pay... A completed registration form should be submitted online at http://registration.arkat-usa.org/ . Credit Card We accept VISA, MasterCard, American Express and Discover. Bank Transfer You will be required to provide information including proof of transfer, ensuring that the delegate's name and company details are also provided to the bank with the transfer details and all bank fees are added to the amount due. Check Checks should be made payable to ARKAT USA, Inc. and should be in US Dollars. <small>Please quote Ref: ARKAT USA, Inc. conference. Fees reduced 10% for IUPAC Members (this does not include books)</small> | | | | | | | | | | | | | | | | | | | |
| Tuesday, March 11, 2008 8:30 am-10:10 am Plenary Lectures 5 and 6 10:10 am-10:40 am Coffee Break 10:40 am-12:20 pm Alternatives: (a) Short Course III; or (b) Invited Lectures 17-20; or (c) Invited Lectures 21-24; Lunch 12:20 pm-1:40 pm Alternatives: (a) Short Course IV; or (b) Invited Lectures 25-28; or (c) Invited Lectures 29-32; 1:40 pm-3:20 pm Coffee Break 3:20 pm-3:50 pm Plenary Lectures 7 and 8 3:50 pm-6:30 pm Poster Session Continued 6:30 pm-7:15 pm Wine Reception 7:15 pm-9:15 pm Conference Dinner (ticket required) | | Introduction to FloHet - 9 Eight previous Florida Conferences held each March for the years 2000 through 2007 inclusive, brought together the academic and industrial communities with an abundance of heterocyclic and synthetic chemistry reflecting the current interest in the subject. The program holds particular interest for the industrial chemical community where pharmaceuticals, agrochemicals and colorants usually contain at least one heterocyclic ring. This, the Ninth Florida Heterocyclic and Synthetic Conference, continues in the tradition of its highly successful predecessors. Registration will start on Sunday, March 9 th with a Welcome Reception. On Monday the 10 th through Wednesday the 12 th , the Conference will feature twelve plenary lectures given by academic and industrial experts from around the world together with invited lectures and short courses on heterocyclic topics. A poster session combined with a buffet supper will be held on the evening of Monday, March 10 th . A wine reception and conference banquet are scheduled for the evening of Tuesday, March 11 th . The conference closes with a farewell party on the evening of Wednesday, March 12 th . | | | | | | | | | | | | | | | | | | | |
| Wednesday, March 12, 2008 8:30 am-10:10 am Plenary Lectures 9 and 10 10:10 am-10:40 am Coffee Break 10:40 am-12:20 pm Alternatives: (a) Short Course V; or (b) Invited Lectures 33-36; or (c) Invited Lectures 37-40; Lunch 12:20 pm-1:30 pm Plenary Lectures 11 and 12 1:30 pm-3:10 pm Closing Remarks 3:10 pm-3:20 pm Short Excursion or Short Course VI 3:30 pm-5:30 pm Farewell Party 6:30 pm-9:00 pm | | Plenary Speakers Jan Backval , Professor, University of Stockholm, Sweden <i>"Catalysis in the Synthesis of Heterocycles"</i> Vittorio Farina , Senior Research Fellow, J&J Pharmaceutical R&D, Belgium <i>"Practical Synthesis of New Pharmaceutical Ingredients"</i> Peter Jacobi , Professor, Dartmouth College, NH, USA <i>"Synthetic Studies Toward Furanosteroids"</i> C. Oliver Kappe , Professor, University of Graz, Austria <i>"Microwave-Assisted Synthesis of Heterocycles"</i> Steve King , Divisional Vice President, Abbott Laboratories, IL, USA <i>"Highly Efficient Synthesis of a New Drug"</i> Harry Kroto , Professor, Florida State University, USA <i>"Architecture in NanoSpace"</i> Patrick Lam , Director, Bristol-Meyers Squibb, CT, USA <i>"C-X Bond Cross-Coupling via Boronic Acid"</i> Armin de Meijere , Professor, University of Goettingen, Germany <i>"New Heterocycles from Small Ring Compounds"</i> Gerald Pattenden , Professor, University of Nottingham, UK <i>"Total Synthesis using Biosynthetic Speculation"</i> Steven M. Weinreb , Professor, Pennsylvania State University, USA <i>"Development of New Methodology for Alkaloid Total Synthesis"</i> Peter Wipf , Professor, University of Pittsburgh, USA <i>"From Strained Rings to Heterocycles: The Bicyclobutane Route to Pyrrolidines"</i> Tony Zhang , Research Fellow, Eli Lilly, Indianapolis, USA <i>"Roles of Palladium in the Process Development of Heterocyclic Drug Candidates"</i> | | | | | | | | | | | | | | | | | | | |
| Cancellations/Refunds: Should you be unable to attend and cancel in writing no later than January 26 th , 2008. ARKAT USA will refund your registration less \$250.00 processing fee. Unfortunately, refunds are not possible after January 26 th , 2008. Substitutions can be made at any time. | | | | | | | | | | | | | | | | | | | | | |

Figure 23. Conference Program at Flohet in 2008.

We hope that all chemists will help our efforts to extend the hand of friendship to organic chemists all around the world. We now have a rather high rejection rate for Arkivoc, but if you submit a manuscript we will try to provide you with constructive criticism that may be helpful even if it is rejected. We hope that you will access ARKIVOC and, when relevant, cite ARKIVOC; the subject index will help you find suitable papers.

We also hope that you will consider coming to the Flohet conference; it gives excellent value and all profits go to support ARKIVOC. We only regret that we cannot reduce fees or give any awards because the whole operation of ARKIVOC and Flohet occurs without major support and is only made possible by the unstinting efforts of a large number of community-minded chemists.

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