Organic & Biomolecular Chemistry

An international journal of synthetic, physical and biomolecular organic chemistry

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IN THIS ISSUE

ISSN 1477-0520 CODEN OBCRAK 6(11) 1857-2020 (2008)





Cover See Takuzo Aida *et al.*, pp. 1871–1876. A photoresponsive chiral molecular plier composed of a ferrocene pivot and zinc porphyrin jaws is able to grab and twist guest molecules.

Image reproduced by permission of Takuzo Aida from *Organic & Biomolecular Chemistry*, 2008, **6**, 1871. Organic & Biomolecular Chemistry



Inside cover

See Robert Cichewicz *et al.*, pp. 1895–1897. Epigenetic mechanisms are used to regulate natural product biosynthesis. The application of small-molecule modulators against epigenetic targets uncovers silenced natural product pathways.

Image reproduced by permission of Robert Cichewicz from *Organic* & *Biomolecular Chemistry*, 2008, **6**, 1895.

CHEMICAL BIOLOGY

B41

Drawing together research highlights and news from all RSC publications, *Chemical Biology* provides a 'snapshot' of the latest developments in the chemical biology, showcasing newsworthy articles and significant scientific advances.



June 2008/Volume 3/Issue 6 www.rsc.org/chembiology

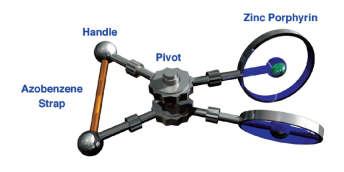
EMERGING AREA

1871

Chiral ferrocenes as novel rotary modules for molecular machines

Kazushi Kinbara,* Takahiro Muraoka and Takuzo Aida*

Ferrocene, a double-decker organometallic compound that generates angular motion, can be used as a unique rotary module for molecular machines.



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PERSPECTIVE

1877

Modulating the electronic properties of porphyrinoids: a voyage from the violet to the infrared regions of the electromagnetic spectrum

Yannick Rio, M. Salomé Rodríguez-Morgade* and Tomás Torres*

The accomplishments achieved so far in the porphyrinoid area relating to the absorption wavelengths of its different members, and the possibility of preparing novel systems with different colour properties and, consequently, new emerging applications, are discussed.

COMMUNICATIONS

1895

Epigenetic remodeling of the fungal secondary metabolome

Russell B. Williams, Jon C. Henrikson, Ashley R. Hoover, Andrlynn E. Lee and Robert H. Cichewicz*

Fungi treated with DNA methyltransferase and histone deacetylase inhibitors exhibit natural product profiles with enhanced chemical diversity demonstrating that small-molecule epigenetic modifiers are effective tools for rationally controlling the native expression of fungal biosynthetic pathways and generating new biomolecules.

1898

Phenylenediamine catalysis of "click glycosylations" in water: practical and direct access to unprotected neoglycoconjugates

Aurélie Baron, Yves Blériot, Matthieu Sollogoub and Boris Vauzeilles*

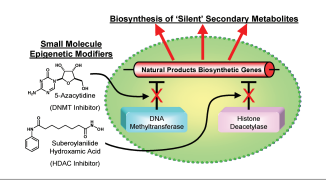
Phenylenediamine-catalyzed click chemistry leads to the efficient, practical, and column-free preparation of neoglycoconjugates from unprotected glucosyl azide, in pure water when aglycon solubility permits.

1902

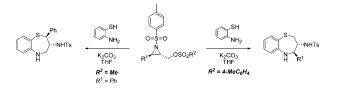
Regio- and stereocontrolled synthesis of novel 3-sulfonamido-2,3,4,5-tetrahydro-1,5-benzothiazepines from 2-(bromomethyl)- or 2-(sulfonyloxymethyl)aziridines

Michinori Karikomi, Matthias D'hooghe, Guido Verniest and Norbert De Kimpe*

A highly efficient regio- and stereocontrolled synthesis of 3-sulfonamido-2,3,4,5-tetrahydro-1,5-benzothiazepines has been accomplished starting from easily accessible aziridine substrates.









Gordon Research Conferences

GRC is holding over 190 meetings in 2008. A few upcoming meetings that may be of particular interest to *Organic & Biomolecular Chemistry* readers are listed below.

ISOTOPES IN BIOLOGICAL & CHEMICAL SCIENCES

February 17-22, 2008 Crowne Plaza, Ventura, CA Chair: Daniel M. Quinn

http://www.grc.org/programs.aspx?year=2008&program=isotopes

NUCLEIC ACIDS

June 1-6, 2008 Salve Regina University, Newport, RI Chairs: Paul Modrich & Jamie H. Cate

http://www.grc.org/programs.aspx?year=2008&program=nucacids

COMPUTATIONAL ASPECTS - BIOMOLECULAR NMR

May 18-23, 2008 Il Ciocco, Lucca (Barga), Italy Chair: Rafael P. Bruschweiler

http://www.grc.org/programs.aspx?year=2008&program=bio_nmr

COMPUTATIONAL CHEMISTRY New Computational Tools For 21st Century Chemistry

July 27 - August 1, 2008 Mount Holyoke College, South Hadley, MA Chair: Jed W. Pitera

http://www.grc.org/programs.aspx?year=2008&program=compchem

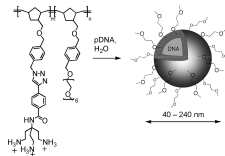
STEREOCHEMISTRY

July 27 - August 1, 2008 Salve Regina University, Newport, RI Chair: Chris H. Senanayake

http://www.grc.org/programs.aspx?year=2008&program=stereo

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1905



Dendronised block copolymers as potential vectors for gene transfection

Tony J. Wigglesworth, Francisco Teixeira Jr., Fabian Axthelm, Sara Eisler, Noemi S. Csaba, Hans P. Merkle, Wolfgang Meier and François Diederich*

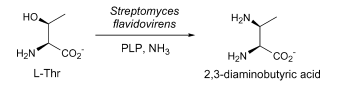
The modular synthesis of block copolymers bearing a dendronised cationic block for DNA complexation and a poly(ethylene glycol) block for encapsulation of the complex is reported. These materials strongly complex DNA and spontaneously form polyion complex micelles in the presence of plasmid DNA.

1912

Identification of a novel β-replacement reaction in the biosynthesis of 2,3-diaminobutyric acid in peptidylnucleoside mureidomycin A

Wai-Ho Lam, Kathrin Rychli and Timothy D. H. Bugg*

A pyridoxal 5'-phosphate dependent enzyme activity has been identified, that is responsible for conversion of L-threonine and ammonia into 2,3-diaminobutyric acid in *Streptomyces flavidovirens*.



1918

Nonenzymic polycyclisation of analogues of oxidosqualene with a preformed C-ring

Johan M. Winne, Pierre J. De Clercq,* Marco Milanesio, Philip Pattison and Davide Viterbo

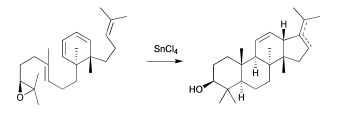
Cationic polycyclisations of partially constrained epoxypolyenes are reported, aspects of which bear interesting analogies with their biosynthetic counterparts involved in sterol formation.

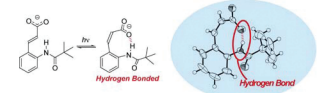
1926

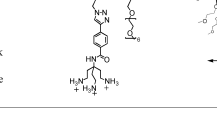
Manipulation of an intramolecular NH \cdots O hydrogen bond by photoswitching between stable E/Z isomers of the cinnamate framework

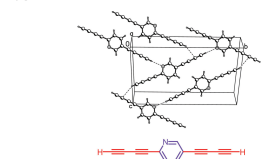
Takashi Matsuhira, Hitoshi Yamamoto,* Taka-aki Okamura and Norikazu Ueyama

An intramolecular NH \cdots O hydrogen bond was switched by E/Z photoisomerization of the cinnamate framework; the p K_a value of the carboxylic acid was decreased by using 313 nm photoirradiation.







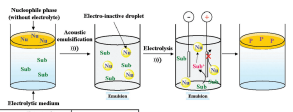


1938

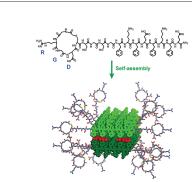
1944

Q

Q



Electrode: Sub Electron transfer Sub' Bulk solution: Sub' + Nu - P & Sub: Substrate, Nu: Nucleophile, Sub': Carbocation, P: P



1949 Ph $InCl_2OMe$ Ph Arl Arl Arl Lil

Carbon-rich molecules: synthesis and isolation of aryl/heteroaryl terminal bis(butadiynes) (HC=C-C= C-Ar-C=C-C=CH) and their applications in the synthesis of oligo(arylenebutadiynylene) molecular wires

Kara West, Changsheng Wang, Andrei S. Batsanov and Martin R. Bryce*

Terminal aryl/heteroaryl bis(butadiynes) (HC–C \equiv C–C \equiv C–Ar–C \equiv C–C \equiv CH) have been isolated and exploited in the synthesis of highly-conjugated oligo(arylenebutadiynylene)s.

A new approach to anodic substitution reaction using acoustic emulsification

Ryosuke Asami, Toshio Fuchigami and Mahito Atobe*

Anodic oxidation of the substrate successfully proceeded without affecting the oxidation of nucleophile and the carbocation generated was rapidly trapped by the coexisting electro-inactive nucleophile droplets before its decomposition.

A cyclic RGD-coated peptide nanoribbon as a selective intracellular nanocarrier

Yong-beom Lim, Oh-Joon Kwon, Eunji Lee, Pyung-Hwan Kim, Chae-Ok Yun and Myongsoo Lee*

A β -ribbon coated with cyclic Arg-Gly-Asp (cRGD) can encapsulate hydrophobic guest molecules, and deliver them into cells selectively.

Hydroindation of allenes and its application to radical cyclization

Naoki Hayashi, Yusuke Hirokawa, Ikuya Shibata,* Makoto Yasuda and Akio Baba*

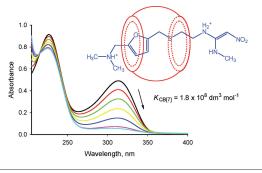
Hydroindation of allenes and radical cyclization of 1,2,7-trienes (allenenes) were accomplished by $HInCl_2$ with high regioselectivity to afford a variety of vinylic indiums which could be used for successive coupling reactions in a one-pot procedure.

1955

Cucurbit[7]uril host–guest complexes of the histamine H₂-receptor antagonist ranitidine

Ruibing Wang and Donal H. Macartney*

The histamine H_2 -receptor antagonist ranitidine forms very stable complexes with the cucurbit[7]uril molecule in aqueous solution over a wide pH range, reducing its acidity and increasing its thermal stability.

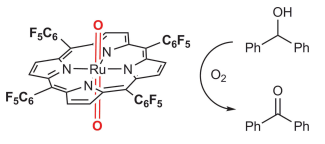


1961

The aerobic oxidation of alcohols with a ruthenium porphyrin catalyst in organic and fluorinated solvents

Vasily N. Korotchenko, Kay Severin and Michel R. Gagné*

Under appropriate activation conditions fluorinated ruthenium porphyrin complexes catalyze the aerobic oxidation of non-enolizable alcohols. Experiments to distinguish between auto-oxidation and metal catalysis are reported.





1966

Exciplex and excimer molecular probes: detection of conformational flip in a *myo*-inositol chair

Manikandan Kadirvel, Biljana Arsic, Sally Freeman* and Elena V. Bichenkova*

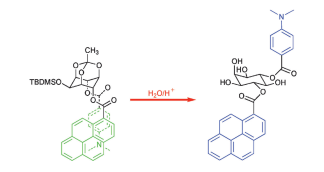
Green fluorescence of exciplex and excimer containing *myo*-inositol orthoesters is lost upon acid-induced deprotection to give penta-equatorial chairs with blue fluorescence. This ring flip provides an irreversible switch for the development of acid-sensitive fluorescent probes.

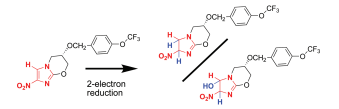
1973

Intermediates in the reduction of the antituberculosis drug PA-824, (6*S*)-2-nitro-6-{[4-(trifluoromethoxy)benzyl]oxy}-6,7-dihydro-5*H*-imidazo[2,1-*b*][1,3]oxazine, in aqueous solution

Robert F. Anderson,* Sujata S. Shinde, Andrej Maroz, Maruta Boyd, Brian D. Palmer and William A. Denny

Stepwise reduction of the nitroimidazole moiety in aqueous solution proceeds atypically at the imidazole ring followed by the nitro group.





Dynamic Stereochemistry of Chiral Compounds

This book provides an overview of fundamental concepts of asymmetric synthesis highlighting the significance of stereochemical and stereodynamic reaction control. Topics include kinetic resolution (KR), dynamic kinetic resolution (DKR), dynamic kinetic asymmetric transformation (DYKAT), and dynamic thermodynamic resolution (DTR). In-depth discussions of asymmetric synthesis with chiral organolithium compounds, atropisomeric biaryl synthesis, self-regeneration of stereogenicity (SRS), chiral amplification with chiral relays and other commonly used strategies are also provided. Particular emphasis is given to selective introduction, interconversion and translocation of central, axial, planar, and helical chirality.

A systematic coverage of stereochemical principles and stereodynamic properties of chiral compounds guides the reader through the book and establishes a conceptual linkage to asymmetric synthesis, interconversion of stereoisomers, molecular devices that resemble the structure and stereomutations of propellers, bevel gears, switches and motors, and topologically chiral assemblies such as catenanes and rotaxanes. Racemization and diastereomerization reactions of numerous chiral compounds are discussed as well as the principles, scope and compatibility of commonly used analytical techniques.

 More than 550 figures, schemes and tables illustrating mechanisms of numerous asymmetric reactions and stereomutations of chiral compounds

• Technical drawings illustrating the conceptual linkage between macroscopic devices such as turnstiles, ratchets, brakes, bevel gears, propellers or knots and molecular analogs

 More than 3000 references to encourage further reading and facilitate additional literature research

• A comprehensive glossary with stereochemical definitions and terms which facilitate understanding and reinforce learning

This book will be of particular interest to advanced undergraduates, graduates and professionals working and researching in the fields of synthetic organic chemistry and stereochemistry.

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Dynamic Stereochemistry of

Chiral Compounds

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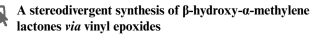
Author: Christian Wolf

Publication date: 14 December 2007 Publisher: RSC Publishing Format: Hardback ISBN: 9780854042463 Price: £49.90

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1981



Marion Davoust, Frédéric Cantagrel, Patrick Metzner* and Jean-François Brière*

The sulfonium ylide epoxidation of aldehydes furnished novel vinyl epoxides, which served as useful building blocks for a stereodivergent route towards β-hydroxy-α-methylene lactones. A formal synthesis of the antibiotic conocandin, displaying this vinyl oxirane motif, is also described.

1994

Novel polyoxazole-based cyclopeptides from Streptomyces sp. Total synthesis of the cyclopeptide YM-216391 and synthetic studies towards telomestatin

Jon Deeley, Anna Bertram and Gerald Pattenden*

Synthetic approaches to the contiguously linked poly-azole units and to the macrocycles in telomestatin and YM-216391 are described. The synthetic cyclopeptide YM-216391 is shown to be the enantiomer of the natural product isolated from Streptomyces nobilis.

2011

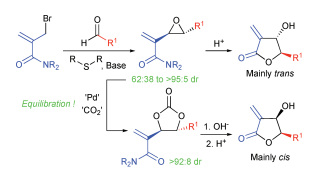
An enantioselective approach to (+)-laurencin

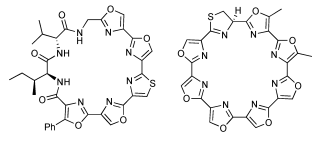
Vikrant A. Adsool and Sunil V. Pansare*

A concise enantioselective synthesis of an advanced intermediate for the marine natural product (+)-laurencin is described. Ring-opening of an ephedrine-derived spiro-epoxide, hemiacetal allylation and ring closing metathesis are the key steps in the synthesis.

MOLECULAR BIOSYSTEMS

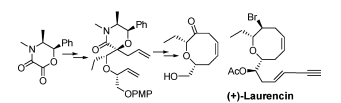
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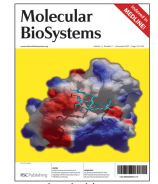






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Lee, Andrlynn E., 1895 Lee, Eunji, 1944 Lee, Myongsoo, 1944 Lim, Yong-beom, 1944 Macartney, Donal H., 1955 Maroz, Andrej, 1973 Matsuhira, Takashi, 1926 Meier, Wolfgang, 1905 Merkle, Hans P., 1905 Metzner, Patrick, 1981 Milanesio, Marco, 1918 Muraoka, Takahiro, 1871 Okamura, Taka-aki, 1926 Palmer, Brian D., 1973 Pansare, Sunil V., 2011 Pattenden, Gerald, 1994 Pattison, Philip, 1918 Rio, Yannick, 1877 Rodríguez-Morgade, M. Salomé, 1877 Rychli, Kathrin, 1912

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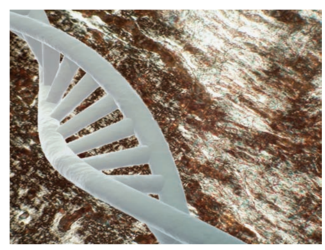
Chemical Biology

Potential phospholipid ordering in cell nuclei causes debate Liquid crystals in all of us

European researchers have sparked debate by claiming that cell nuclei could contain liquid crystals. The hypothesis is supported by experiments revealing that DNA transcription is possible in such environments, says the team from the UK and Germany.

George Attard from the University of Southampton, UK, and colleagues have shown that DNA can be transcribed into RNA within a liquid crystalline phase formed by phospholipids. Moreover, they found that isolated cell nuclei exhibit an optical property known as birefringence that is characteristic of liquid crystals. The researchers say that together these results 'raise the possibility that lipids might form organised structures in the nucleus in vivo.'

Attard says his research is 'off the wall', and adds that mainstream researchers are reluctant to accept his views. Indeed, Roel van Driel, an expert in nuclear organisation from the University of Amsterdam, the



Netherlands, is not convinced. He points out that Attard's group freezedried the studied nuclei, which will have caused major structural rearrangements. Therefore there is no evidence that living nuclei show birefringence, he says. Attard accepts this, but adds: 'We have x-ray data from non-freeze-dried nuclei which Birefringence suggests that phospholipids may form ordered structures inside the cell nucleus are consistent with long-range ordering.'

Van Driel also says that for living nuclei to show birefringence, chromatin – a complex of DNA and proteins – would have to be ordered on the length scale of the nucleus, which it is not. Attard counters that chromatin could adopt any degree of structural ordering, or none at all, within a liquid crystalline phase, but that these phases would still cause birefringence.

Despite criticism, Attard says that it is 'likely' that nuclei are in a liquid crystalline state. Cell nuclei are rich in phospholipids and these molecules are known to self-organise into structures – for example membranes – he explains. Based on the intermolecular forces, 'you would be more surprised to find that nuclei are not liquid crystalline rather than the reverse,' says Attard. Danièle Gibney

Reference

J Corsi et al, Chem. Commun., 2008, 2307 (DOI: 10.1039/b801199k)

In this issue

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Radical proposal for nitrate link to asthma

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Communicating with nature

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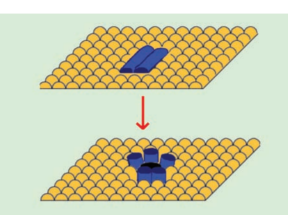
Research highlights

Break through study yields insights into viral mechanism **How does a virus bore a hole in a cell?**

Chemists in the US studying how viruses enter cells say their results could help in the search for new antiviral medicines.

Stanford University chemists Richard Zare and Soonwoo Chah used cell and virus models to investigate an early stage in viral infection. Using lipid spheres called vesicles as cell mimics they studied the way the vesicles rupture after exposure to PEP1, a helix-shaped peptide that resembles a peptide found in the hepatitis C virus. Zare says that 'it is important to understand how viruses interact with and break up the cell membrane. Knowledge of the exact sequence and duration of these steps is crucial to developing possible strategies for combating disease.'

When a virus invades a cell in the body, it fuses with the cell membrane and releases its genetic material into the host cell, turning into a 'chemical factory' that produces more copies of



Peptide PEP1 (blue) lies flat on a vesicle surface before rearranging to form a pore through the membrane

Reference

S Chah and R N Zare, *Phys. Chem. Chem. Phys.*, 2008, DOI: 10.1039/b802632g the virus. It is known that there is an intermediate stage between the time a virus merges with the cell membrane and the time it delivers its genetic contents into the cell. 'During this period, the host cell's fate hangs in the balance,' Zare explains. Cell infection is often irreversible once a virus has penetrated the cell – so this stage 'may represent an opportunity for drug development,' suggests Zare.

Zare and Chah investigated the intermediate stage more closely using surface plasmon resonance (SPR) microscopy to measure the lipids' optical properties. Since these properties are different for intact and ruptured vesicles, SPR allows the researchers to follow the rupture mechanism in real time. They found that, after introducing the peptide to the vesicles, the peptides first lie flat on the surface then switch to cross the membrane, forming pores. 'This attack causes the vesicles to transform into a lipid bilayer,' says Zare.

Richard Epand, a biochemist from McMaster University, Hamilton, Canada, is impressed. He says that although using a viral peptide and not the intact virus means there are some limitations, the work 'could contribute significantly to our understanding of viral fusion processes.'

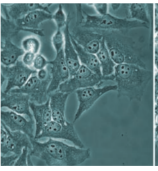
Michael Spencelayh

Nanostructures that puncture cell membranes prove selective Peptides provide fatal blow for cancer cells

Peptide nanostructures that punch holes in cancer cells are 'the first step towards efficient nanochemotherapeutics,' say chemists in Canada. Normand Voyer and colleagues at the University of Laval in Quebec have designed a series of modified peptide nanostructures that can puncture cancer cell membranes, leading to the cells' death.

The team explains that in the past decade, cancer cell resistance to chemotherapeutic agents has led to increased cancer deaths. We believe that nanochemotherapeutics can overcome this problem due to the particular properties of nanometresized compounds,' says Voyer.

Basing their structures on a membrane-disrupting peptide they had made previously, the researchers engineered analogues that would be selective for cancer cells. The engineered peptides are inactive until they reach cancer



When cancer cells (left) are treated with peptide nanostructures their cell membranes are destroyed (right) cell surfaces where they convert into an active cell membrane disruption agent. Since the enzyme that activates the peptides is overexpressed in prostate cancer cells, normal cells do not activate the peptide to the same extent, leading to the peptides' selectivity.

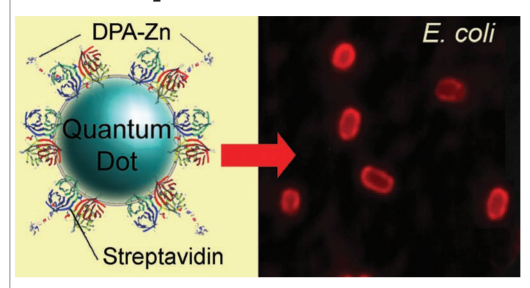
Vincent Rotello, an expert in the supramolecular chemistry of biological and materials systems at the University of Massachusetts, Amherst, US, is enthusiastic about the findings. 'While enzymatic activation has been used before for therapeutics,' he says, 'this peptidebased scaffold has great promise due to the modular nature of its construction.' This is because the amino acid building blocks used to assemble the peptides can be readily varied, which provides 'incredible control over the structure and dynamics of the eventual therapeutics,' Rotello adds.

Voyer explains that the work 'illustrates chemists' abilities to design novel nanometresized molecular architectures from scratch to address highly challenging problems.' Future efforts will be geared towards 'determining the mechanism of action of this new class of antitumour agents,' he adds. *Kathleen Too*

Reference

P L Boudreault *et al, Chem. Commun.,* 2008, 2118 (DOI: 10.1039/b800528a)

Nanoparticles target bacteria to illuminate infection in vivo **Probes spot the difference**



Fluorescent probes are shedding light on bacterial infection. Bradley Smith from the University of Notre Dame, US, and colleagues have made fluorescent probes that can distinguish between different mutants of the same bacterial species and can be used to observe the bacteria in vivo.

The team made the probes by attaching the bacteria-targeting ligand zinc(II) dipicolylamine (Zn-DPA) to a fluorescent nanoparticle called a quantum dot. Zn-DPA targets bacterial cells because it has a strong affinity for the phospholipids in their outer cell membranes. However, when attached to a relatively large quantum dot, it appears the ligand is unable to reach the phospholipids in some bacteria, leading to selective binding.

Smith's team showed that Zn-DPA-quantum dots can stain a rough strain of *Escherichia coli* intensely, but not smooth *E. coli* strains or Gram-positive bacteria – a group of bacteria that can be stained by crystal violet dye. They suggest this is because Grampositive bacteria have thick cell walls with pores too small to allow the quantum dots to pass through. Similarly, the smooth *E. coli* strains are surrounded by a polysaccharide layer which prevents the Zn-DPA-quantum dots reaching Zinc (II) dipicolylamine coated quantum dots (left) are a selective stain for some bacteria such as a rough strain of Escherichia coli (right)

Reference

W M Leevy et al, Chem. Commun., 2008, 2331 (DOI: 10.1039/b803590c) the phospholipids in the membrane beneath.

The team says it should be possible to exploit the probes' selectivity in highly sensitive multicoloured staining schemes to identify bacterial species and mutant strains rapidly in contaminated samples.

The group also tested the feasibility of using the probes for in vivo imaging of bacterial infection in mice. They found the bacterial fluorescent signal to be 10-fold greater than the background autofluorescence. But, admit the scientists, it is only 1.5 times greater than when bacteria are labelled with Zn-DPA attached to an organic fluorophore rather than a quantum dot. The observed fluorescence is limited because maximum tissue penetration is achieved when a fluorophore's excitation and emission wavelengths are both between 650 and 900nm - these quantum dots emit at 800nm but excite below 500nm.

Smith plans to develop the optical imaging method so that it can be used to evaluate antibiotic therapy in animals. 'The challenge is to make very bright and highly selective near-infrared imaging probes that also exhibit favourable pharmacokinetics and low toxicity,' says Smith. *Freva Mearns*

News in brief

This month in Chemical Science

Swellable gels fix bad backs

This month's Instant insight sees Brian Saunders and Tony Freemont discuss a new approach for treating back pain using injectable microgel implants.

Carbon nanotubes wear coats to deliver drugs

Polymer coated carbon nanotubes could find a new use in drug delivery, claim Korean scientists.

Fungi wake up to new natural products

Re-awakening 'silent' metabolic pathways in fungi has revealed a new range of natural products to US scientists.

How mouldy is your house?

Concerns about mould growing in houses are on the increase, claim mycologists in France.

See www.rsc.org/chemicalscience for full versions of these articles.

This month in Chemical Technology

People power

In this month's interview, Duncan Graham explains just how important people are for the future of science.

On-chip suction stops worm wiggling

US scientists have developed a microfluidic method for immobilising worms, allowing them to be used in high throughput studies of disease.

Making sense of DNAzymes

In this month's Instant insight, Itamar Willner and colleagues discuss the applications of DNAbased enzymes.

See www.rsc.org/chemicaltechnology for full versions of these articles.

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Noise control as competitive RNA interferes with gene expression **Knowledge out of chaos**

Scientists have upset gene expression to study its randomness and discover how the cell reduces this variability.

VATIONAL SCIENCE FOUNDATION

An organism's genetic code does not simply equate to a certain outcome. Noise in gene expression can result in physical differences in genetically identical populations. Synthetic biologists who want to construct and study gene networks need to understand this noise for their own experiments to be valid, and new work from America explores just that.

In gene expression, genes are read and translated into protein products using small RNA molecules overseen by a large complex called a ribosome. Because there are so many of these small RNAs in a cell, variations in their relative levels can affect protein production. Andrew Ellington, at the University of Texas at Austin, US, and colleagues decided to investigate this phenomenon further.



Variations in the amounts of different RNA molecules in cells can affect protein production The group made small ribosome competing RNA (rcRNA) molecules that were designed to compete with cell RNA for the ribosome and affect gene expression. Their aim was to use the rcRNAs as a tool in gene expression noise studies to introduce noise controllably using different amounts and types of rcRNA.

When the researchers added the rcRNAs to *Escherichia coli* cells they found that their rcRNAs do generate noise, causing fluctuations in the production of a fluorescent protein by the bacteria. The team used its rcRNA approach to show that DNA sections called operons are highly effective at reducing noise as they eliminate the relative RNA fluctuations between genes.

Jim Collins, co-director of the Centre for BioDynamics at Boston University, US, is very impressed with the new tool. He describes the work as 'an excellent example of how synthetic biology techniques can be used to gain insight into fundamental biological principles.' *Laura Howes*

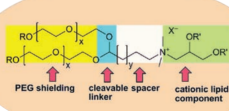
Reference

J J Tabor et al, Mol. BioSyst., 2008, DOI: 10.1039/b801245h

Polymeric lipids ensure DNA vectors meet their targets Plastic coats wrap up gene delivery

UK chemists have used smart polymers to deliver DNA into cells. Based on pH-sensitive poly(ethylene glycol) (PEG) lipids, the polymers can be used as a removable protective coat for gene delivery systems.

Gene delivery systems, or vectors, have to protect their DNA cargo from enzymes, cross cell membranes and yet still release a therapeutic dose of intact DNA inside the target cell. Viral vectors can deliver genes into cells, however, they can provoke an immune response which limits their therapeutic use. One of the problems often associated with nonviral gene delivery systems is that 'the efficiency is too low and that the vectors are not sufficiently stable. particularly in vivo,' says Helen Hailes a reader in chemical biology



The pH-sensitive coating shields the vector until the PEG region (yellow) is removed by hydrolysis at the linker (blue)

Reference

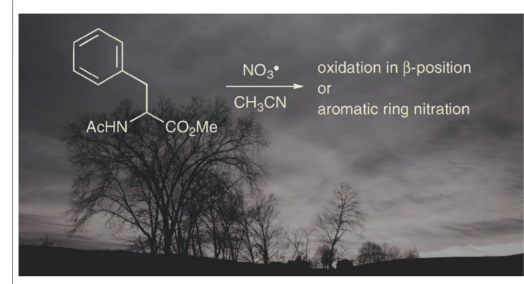
J B Wong et al, Mol. BioSyst., 2008, DOI: 10.1039/b719782a at University College London, UK. To overcome this problem, Hailes and her colleagues have developed acid-cleavable PEG lipids to shield the DNA in a non-viral vector. The vector consists

of a targeting peptide and cargo DNA with the PEG lipids as covering. This coating stabilises the particles, protects the DNA from nuclease enzymes, provides water solubility and facilitates transport through the cell membrane. Once the vector is inside the cell, the lower pH triggers hydrolysis and shedding of the coating, releasing the cargo DNA. As different PEG lipid structures are hydrolysed at different pH, this offers a method of controlling the pH dependence of DNA release, suggests Hailes.

The team demonstrated the new system's effectiveness by using it to transfer DNA coding a bioluminescent enzyme into different cell types, and then measuring the enzyme's activity. 'The PEG lipids seem to provide cell specific properties,' comments Antonio Villaverde, an expert in non-viral gene therapy at the Autonomous University of Barcelona, Spain. 'This could be an interesting element to favour cell targeting in delivering such constructs.'

Looking to the future, Hailes says that the team hopes to 'design different tunable features into the lipids for a range of delivery applications.' She adds that this could include using the system to deliver small interfering RNA – short strands of RNA that can be used to interfere with gene expression. *Russell Johnson*

Reaction between atmospheric species could explain their role in airway disease **Radical proposal for nitrate link to asthma**



Australian researchers have discovered that nitrate radicals irreversibly damage amino acids. This raises the possibility that the radicals play a role in respiratory disease, they claim.

Nitrogen dioxide and ozone have been linked to airway diseases such as asthma, although their exact role is not clear. In the atmosphere, these gases can react to form extremely reactive nitrate radicals, leading Uta Wille and Duanne Sigmund at the University of Melbourne, Victoria, to question whether there is a link between these radicals and respiratory illness.

As part of their research, the duo has found that nitrate radicals irreversibly damage aromatic amino acids, forming compounds including β -nitrate esters, β -carbonyl and aromatic nitro-compounds. 'The reaction forms oxidised products,' explains Wille,

Nitrate radicals will react with aromatic amino acids. The radicals form in the atmosphere and build up overnight 'some of which have been found in polluted air and are associated with immune stimulation.' By analysing these products the researchers proposed a mechanism for the oxidation and nitration reactions.

'The next step is to study the radicals' role in damage to proteins, peptides and carbohydrates – molecules that line the cells of the respiratory tract and so are in direct contact with the atmosphere,' says Wille. Ultimately the team wants to check whether the radicals can migrate through the cell membrane, 'where they could cause damage inside the cell,' Wille adds.

Malcolm Forbes, an expert in free radical chemistry at the University of North Carolina at Chapel Hill, US, welcomes the research, and says: 'The results will have a significant impact on research into oxidative damage to proteins, particularly in regard to respiratory illness. The challenge now is to correlate these results with in vivo studies to assess the real impact to society.' *Russell Johnson*

Reference

D C E Sigmund and U Wille, *Chem. Commun.*, 2008, 2121 (DOI: 10.1039/b803456g)

In the current issue of Research Articles...



Fast-lysis cell traps for chemical cytometry Paul J Marc *et al, Lab Chip*, 2008, **8**, 710 (DOI: 10.1039/b719301g)

Intracellular applications of analytical SERS spectroscopy and multispectral imaging

Igor Chourpa et al, Chem. Soc. Rev., 2008, **37**, 993 (DOI: 10.1039/b714732p)

Simultaneous determination of glycated haemoglobin, a long term biomarker of diabetes mellitus, and total haemoglobin by isotope dilution and HPLC-ICP-MS

M Estela del Castillo Busto *et al, J. Anal. At. Spectrom.*, 2008, **23**, 758 (DOI: 10.1039/b718008j)

Highly efficient quenching of excimer fluorescence by perylene diimide in DNA

Nicolas Bouquin *et al, Chem. Commun.*, 2008, 1974 (DOI: 10.1039/b802193g)

Measuring the simultaneous effects of hypoxia and deformation on ATP release from erythrocytes

Andrea Faris and Dana M Spence, *Analyst*, 2008, **133**, 678 (DOI: 10.1039/b719990b)

The importance of surfaces in single-molecule bioscience Mari-Liis Visnapuu *et al*, *Mol. BioSyst.*, 2008, **4**, 394 (DOI: 10.1039/b800444g)

Quantitative SERRS for DNA sequence analysis

Duncan Graham and Karen Faulds, *Chem. Soc. Rev.*, 2008, **37**, 1042 (DOI: 10.1039/b707941a)

Porphyrin-bile acid conjugates: from saccharide recognition in the solution to the selective cancer cell fluorescence detection Jarmila Králová *et al, Org. Biomol. Chem.*, 2008, **6**, 1548 (DOI: 10.1039/b717528k)

Read more at www.rsc.org/chembiology

Interview

The protein detective

Kathryn Lilley tells Michael Smith how curry and beer could be the downfall of biomarkers



Kathryn Lilley

Kathryn Lilley is director of the Cambridge Centre for Proteomics, located in the Systems Biology Centre, Cambridge, UK. Her group is involved with several important collaborative proteomics projects, in particular involving the model organisms Arabidopsis and Drosophila.

What inspired you to become a scientist?

My grandfather was an amateur gardener and my mother picked up his love of plants. I think it was her love and enthusiasm for nature and plants that made me very interested in biology.

I always had a battle as to whether to become a musician, as that's my other great passion. I suppose I realised that you could be a professional scientist and an amateur musician but it would be very difficult to live your life the other way around!

Why did you choose to specialise in proteomics?

After my PhD, which involved a lot of protein sequencing, an opportunity came up to run a protein sequencing facility at the University of Leicester.

About that time, there were some major papers published on mass spectrometry of proteins, particularly from Mathias Mann's group at the European Molecular Biology Laboratory in Heidelberg. Although we purchased our own mass spectrometers in Leicester to support peptide and DNA synthesis, they were not the right type to carry out these new methods of protein identification. I was frustrated as I really wanted to carry out these wonderful new proteomics technologies.

Next, the opportunity came to move to Cambridge to set up a proteomics laboratory with funding from the BBSRC and with it, the idea that I could do all this wonderful scalable protein identification. After running a core facility for many years in Leicester, in Cambridge I got the opportunity to get back in touch with how to answer interesting biological questions.

What biological questions are you interested in?

In the early days of the facility, most projects centred around knowing the differences in protein abundances between mutant and wild types or treated and untreated states. If we can see which proteins are changing in their expression or posttranslational state, it gives us an insight into what's going on inside a cell.

I got fed up with identifying the same sets of proteins and wanted to probe into the lower abundance fraction of proteins. I also wanted to get information on where they resided in a cell and who with!

What are you working on at the moment?

Organelle proteomics. The first stage of many proteomics experiments is to take the cells and mash them up. Usually you add a healthy dose of detergent.

This means you lose all the spatial information about your proteins within a cell. I think this is a very important thing to study because where proteins are and what they associate with is going to give us a huge amount of information that we can't get from just looking at their abundance.

We are starting to look at components from signalling pathways – how they move around the cell upon signalling events and how this may change under different situations, including development and differentiation. We've spent a few years trying to fine tune the methodologies. We're not there yet but we've gone quite some way to be able to produce technology that is robust both in terms of identifying proteins associated with certain organelles and mapping onto that the position of protein complexes.

What do you think about biomarkers as an ultimate aim of proteomics?

Proteomics is a very attractive way of finding biomarkers but it is fraught with issues.

The main issue is the dynamic range of protein concentrations within plasma, which currently no techniques can cover.

Secondly, if you do find biomarkers in a certain set of proteins that are always up-regulated in someone who is suffering from a cold, how discriminatory is that? I can't see that there is going to be any one biomarker that will tell you which disease is present.

Finally, blood plasma is really a mirror of your general state of health and what you've been up to in the last day or two. There is a lot of person-inperson variability depending on your health and whether you've had a curry and several beers the night before! In a population set, I think it's going to be very difficult to find an abundance change that is disease-specific.

What advice would you give to someone considering a career in science?

Looking back on my own career, I became too specialised too soon. What I lost touch with very quickly, and I wish I hadn't, was maths. More and more in biology, we are making quantitative measurements. The way in which we deal with those measurements is controlled by statistical and mathematical tools. We need that know-how to be able to design our experiments properly and to see whether we believe the data that's coming out. My advice is to avoid losing touch with other scientific disciplines. Try and stay broadly focused.

Instant insight **Communicating with nature**

Bacteria have invented a potentially global language - quorum sensing. Kim Janda of the Scripps Research Institute in La Jolla, US, translates

Over the course of history, humans have developed countless ways to communicate with each other, and over 6800 languages have been catalogued. Despite these advances, our verbal relations with other species remain somewhat limited to gestures and shouts. as can be seen in the case of dogs and their owners. On a microscopic level, bacteria can communicate with one another through a different language, one based on small molecules, using a mechanism known as quorum sensing (OS). In contrast to humans' limited verbal communication capacities, QS allows communication and interaction with other bacterial species, and even other organisms such as mammals.

Bacterial language relies on the exchange of small chemical signals, called autoinducers. Through this exchange, bacteria monitor their density and regulate gene expression in a populationdependent manner. This allows them to coordinate their behavior and function, equipping the bacterial communities for competition or cooperation with multicellular organisms. A classic example is the symbiosis between the Hawaiian bobtail squid Euprymna scolopes and the luminescent bacterium Vibrio fischeri. In this relationship, the bacteria provide the squid with luminescence, allowing it to blend in with the moonlight while feeding, and so avoid casting shadows on the sea floor which would alert both predators and prey. At the same time, the bacteria also benefit, as they receive nutrients and safety.

QS has traditionally been referred to as a communication mechanism between bacteria within one species. However, research is emerging that implicates a role for QS in interspecies communication and competition, and such systems have been proposed to exist in a wide variety of bacteria. Particularly relevant to interspecies



The Hawaiian bobtail squid has a symbiotic relationship with a luminescent bacterium

Reference

C A Lowerv. T J Dickerson. and K D Janda, Chem. Soc. Rev. 2008, DOI: 10.1039/b702781h

communication is the autoinducer 2 (AI-2)-based QS system, which has been suggested to function in over 50 bacterial species. Recently, it was shown that Actinomyces naeslundii and Streptococcus oralis, two bacteria responsible for oral plaque formation, require AI-2 production to initiate plaque development.

But communication amongst bacterial species is not always so cooperative; certain autoinducers and their byproducts have been shown to have cytotoxic effects on other bacteria. Pseudomonas aeruginosa is especially adept at this intercellular competition, in that at least two autoinducer-derived molecules exhibit detrimental effects towards other bacteria, most notably *Staphylococcus* aureus. This activity may give P. aeruginosa a competitive advantage over S. aureus in the lungs of cystic fibrosis patients, a clinical setting plagued

by infections due to these two pathogens.

In addition to helping bacteria organise their behaviour and functions, recent research suggests OS is a means for bacteria to interact with other organisms. Similar to bacterial interspecies relations, OS systems may mediate this interkingdom signalling either through host cell recognition of bacterial signals or through the unregulated action of an autoinducer on the host cell. Several studies have detailed the effects of AHL (N-acylhomoserine lactone)based signalling molecules on human cells - the responses ranging from immuno-activation to cell death.

A potential communication language between humans and Escherichia coli has also been described. E. coli responds to two human-derived small molecule signals, adrenaline and noradrenaline, to regulate virulence expression. For this same purpose, E. coli also employs a small molecule of its own production, termed AI-3. Based on the role of E. coli in the gastrointestinal tract, and the overlap between bacterial recognition of AI-3 and adrenaline. AI-3 has been suggested to play a role in maintaining intestinal homeostasis.

Because QS can mediate so many relationships, it may represent a global language that spans every kingdom of life. Human interpretation may impart a deeper knowledge of bacterial lifestyles and provide the opportunity for an appropriate response, at least one of which would be developing pharmacological interventions for bacterial infection.

Read more in the tutorial review 'Interspecies and interkingdom communication mediated by bacterial quorum sensing' in issue 7, 2008 of Chemical Society Reviews.

Chemical Biology

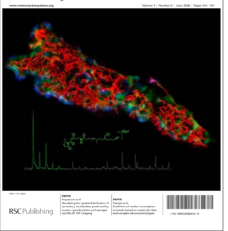
Essential elements

Emerging Investigators

Highlighting the brightest new researchers in the field, issue 6 of Molecular BioSystems (MBS) is not to be missed. The 20 full research papers, seven communications and two reviews are written by outstanding young scientists at the chemicaland systems-biology interfaces. The issue features novel methods to visualise and manipulate protein function in living cells, the development of chemical techniques to monitor specific protein post-translational modifications, new insights into metabolomics and much, much more.

All the contributors were personally recommended by MBS editorial or advisory board members as young scientists whose work has the potential to

Molecular **BioSystems**



influence the future directions of these fields. All submissions were subjected to full peer review and the result is an issue showcasing

work in some of the most fascinating and important areas of biology.

We intend to run future issues of this kind so watch this space. Finally, MBS extends a big thank-you to all the Emerging Investigators themselves for making this such an excellent collection of papers. We wish them every success in their future careers and - in the words of Tom Kodadek, the MBS editorial board chair - 'Clearly the future of this exciting area of biology is in good hands!'

Find out more at www.molecularbiosystems.org

And watch out for a related theme issue from ChemSocRev (www.rsc.org/chemsocrev) in July; issue 7 will be a thematic issue examining the interface of chemistry with biology.

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Pioneers in Miniaturisation Prize

Leading the way in miniaturisation, Lab on a Chip has teamed up with Corning Incorporated to again host the Pioneers in Miniaturisation Prize. Spanning a variety of disciplines, this prize recognises outstanding achievements and significant contributions by a younger scientist to the understanding and advancement of micro- and nanoscale science.

As a leading-edge science and technology organisation, Corning Incorporated is keen to reward, recognise and encourage the development of miniaturisation in the chemical and biological sciences and promotes interdisciplinary research required for the most significant innovations in this area.

The recipient of the award will receive a US\$5000 bursary to support their continued contribution to the field. A deadline for applications has been set for 31st August 2008. Following the final decision, which will be made by committee, a winner will be announced at the µTAS 2008 conference, in San Diego, CA, US.

For more information visit www.rsc.org/loc

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