

JOURNAL OF THE CHEMICAL SOCIETY

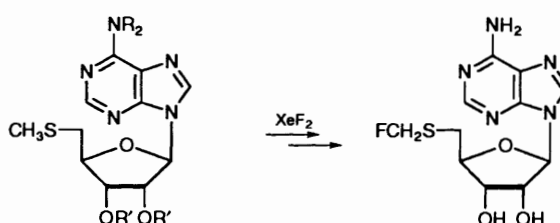
Perkin Transactions 1

Organic and Bio-organic Chemistry

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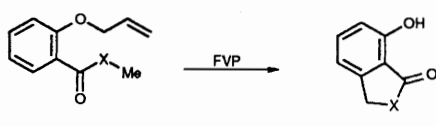
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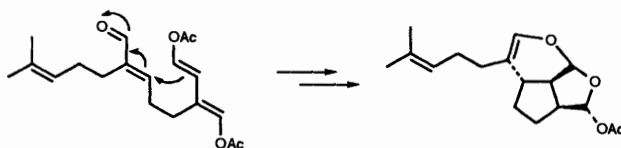
- 155 Pyrolysis of *O*-allyl salicylic amides and esters, and related compounds: formation of isoindolones and phthalides

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- 161 Preuplotin, a putative biogenetic precursor of the euplotins, bioactive sesquiterpenoids of the marine ciliated protist *Euplotes crassus*

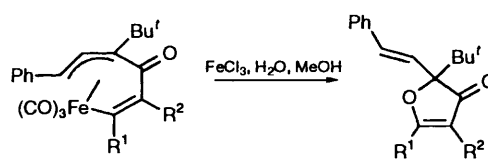
Graziano Guella, Fernando Dini, Antonio Tomei and Francesco Pietra



Preuplotin and its putative biogenetic descendant euplotin C are contained in *Euplotes crassus* together with euplotins A and B, likely adjuvant factors in adaptive radiation of this marine ciliate

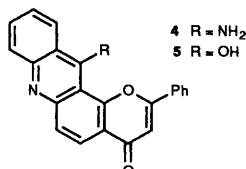
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173 The synthesis of pyranoacridinone inhibitors of protein tyrosine kinases

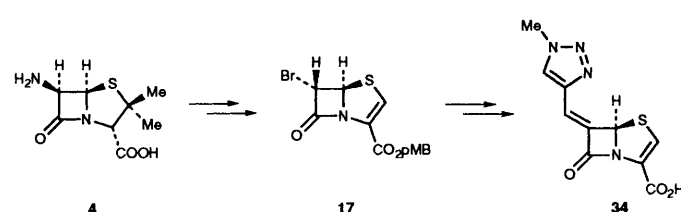
Paul W. Groundwater and Kevin R. H. Solomons



The synthesis of pyranoacridinone inhibitors of growth factor-stimulated cell proliferation is described

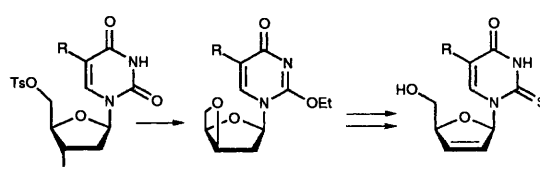
179 Synthesis of (5*R*)-(Z)-6-(1-methyl-1,2,3-triazol-4-ylmethylene)penem-3-carboxylic acid, a potent broad spectrum β -lactamase inhibitor, from 6-aminopenicillanic acid

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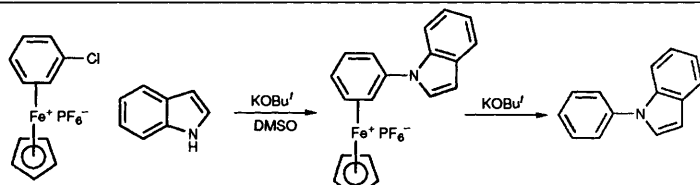
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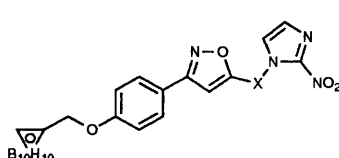
Richard A. Brown, Sharon I. S. Fernando and Roger M. G. Roberts



The use of 1,8-phenanthroline and potassium *tert*-butoxide as demetallating agents for [η -arene](η -cyclopentadienyl)Fe][PF6] complexes is described. The findings were used to develop an efficient one-pot arylation procedure

203 Tumour-targetted boranes. Part 2. Coupling of *closو-carboranes to substituted 2-nitro-imidazoles via 1,3-dipolar cycloaddition*

Martin Scobie, Mary F. Mahon and Michael D. Threadgill

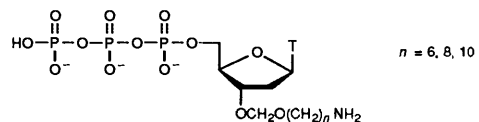


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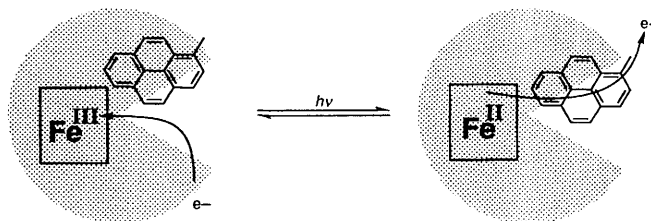
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Preparation and properties of thymidine 5'-triphosphates bearing an ω -aminoalkoxymethyl spacer arm at their 3'-O are described

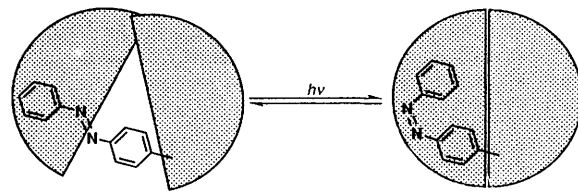
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The First Completely Chemical Synthesis of (6S)- N^5 -Formyltetrahydropteroyl Poly- γ -L-glutamic Acid Derivatives **A.L. Fitzhugh, R.K. Akee, F.C. Ruei, J.Wu and J.R. Klose**

Synthesis of 2,5-Dioxygenated Pyrazine 4-Oxides: Total Synthesis of a New Inhibitor of Superoxide Anion Generation, OPC-15161 **Y. Kita, S. Akai, H. Fujioka, Y. Tamura, H. Tone and Y. Taniguchi**

Conjugate Addition of the Phenyldimethylsilyl Group to α,β -Unsaturated Carbonyl Compounds using a Silyl Zincate in Place of the Silyl Cuprate **R.A.N.C. Crump, I. Fleming and C.J. Urch**

Enantioselective Addition of Diethylzinc to Aldehydes using Chiral Polymer Catalysts Possessing a Methylene Spacer **M. Watanabe and K. Soai**

Novel Insertion, Rearrangement and Addition Products from a Dihalocarbene Reaction with 5(10)-Unsaturated Steroids **J.F. Templeton, Y. Ling, W. Lin, R.J. Pitura and K. Marat**

The Synthesis of E Enol Ethers of Protected 4-Amino Aldehydes **S. Warren, E.W. Collington and S.K. Armstrong**

Cyclization of Isothiosemicarbazones. Part 10. A Novel Route to 2-Amino-[1,2,4,]triazolo[1,5-a]pyridine Derivatives **C. Yamazaki, Y. Miyamoto and H. Sakima**

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Cycloaddition Reactions between Carbonyl Oxides and Dicarbonyl Compounds: Isolation and Characterisation of Novel Polycyclic 1,2,4,6-Tetroxepane Derivatives **K.J. McCullough, T. Sugimoto, S. Tanaka, S. Kusabayashi and M. Nojima**

Enzymatic Cyclization of 2,3-Dihydrosqualene and Squalene-2,3-epoxide by Squalene Cyclases: from Pentacyclic to Tetracyclic Triterpenes **M. Rohmer and I. Abe**

Chemistry of the Mycalamides. Antiviral and Antitumour Compounds from a Marine Sponge. Part 4. Reactions of Mycalamide A and Alkyl Derivatives with Basic Nucleophiles **J.W. Blunt, A.M. Thompson, M.H.G. Munro and B.M. Clark**

Biosynthesis of Hyoscyamine involves an Intramolecular Rearrangement of Littorine **R.J. Robins, P. Bachmann and J.G. Woolley**

Synthesis of Functionalized Compounds Containing a Difluoromethylene Moiety **C.-M. Hu and J. Chen**

Diels-Alder Reactions Catalysed by Cation-exchanged Clay Minerals **R.W. McCabe, A. Mateer, K. Martin, S. Dyer and J.M. Adams**

Biosynthesis of Porphyrins and Related Macrocycles. Part 42. Pulse Labelling Experiments Concerning the Timing of Cobalt Insertion during Vitamin B12 Biosynthesis **A.R. Battersby, F.J. Leeper, N.P.J. Stamford, A. Prell, S.M. Monaghan, R.A. Vishwakarma and S. Balachandran**

Glycosides. Part 1. A New Synthesis of 1,2-trans O-Aryl Glycosides, via Tributyltin Phenoxides **S. Mottadelli, M.L. Gelmi and F. Clerici**

