

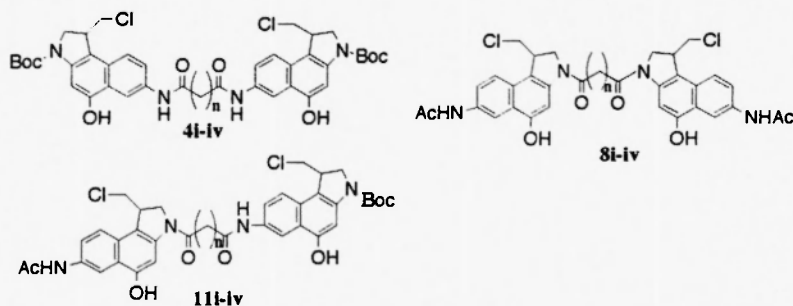
Graphical Abstracts

Heterocycl. Commun. 5 (1999) 497-502

Design and Synthesis of 1,2,9a-Tetrahydrocyclopropa[c]benz[e]indole-4-one (CBI) Dimers

Guofeng Jia, Hirokazu Iida and J. William Lown*

Department of Chemistry, University of Alberta, Edmonton, AB, Canada, T6G 2G2

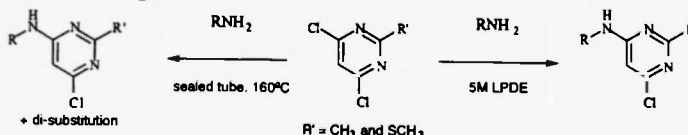


Heterocycl. Commun. 5 (1999) 503-508

Regiocontrolled Amination of Dichloropyrimidines in LiClO₄ - Et₂O Solutions.

James Garner and Adam McCluskey* Department of Chemistry, The University of Newcastle, University Drive, Callaghan, NSW, Australia 2308

Treatment of dichloropyrimidine with amines in 5M LPDE yields only the mono-aminated product allowing for the selective introduction of a second amine in subsequent synthetic steps.



Heterocycl. Commun. 5 (1999) 509-514

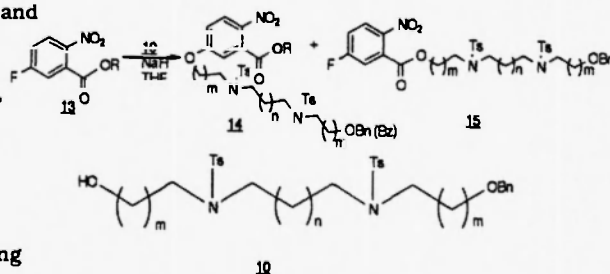
S_NAr REACTIONS OF METHYL AND ETHYL 2-NITRO-5-FLUOROBENZOATES IN THE SYNTHESIS OF PYRROLO-[2,1-c][1,4]BENZODIAZEPINE PRECURSORS

Hirokazu Iida,¹ Yukihiro Misumi,¹ Kiyoshi Matsumoto,^{*1} and J. William Lown,^{*2}

¹ Graduate School of Human and Environmental Studies, Kyoto University, Kyoto 606-8501, Japan

² Department of Chemistry, University of Alberta, Edmonton, AB, Canada, T6G 2G2

The title reaction was investigated as part of an effective synthesis of pyrrolo[2,1-c][1,4]-benzodiazepines possessing a long alkylamino unit at position 7.

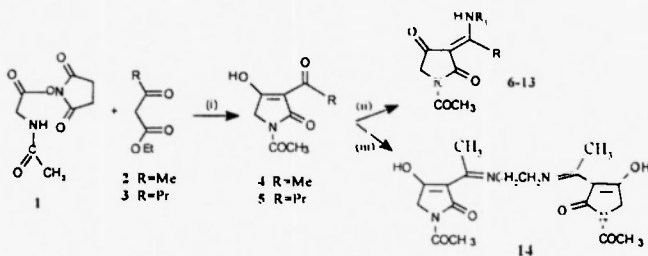


SYNTHESIS AND NMR SPECTROSCOPIC STUDIES OF NOVEL N-ACETYL-3-AMINOALKYL TETRAMIC ACIDS

Efstathios Gavrielatos^a, John Markopoulos^b, Olga Iglessi-Markopoulou^{a*}.

^a Laboratory of Organic Chemistry, Department of Chemical Engineering, National Technical University of Athens, Zografou Campus, 157 73 Athens, GREECE.

^b Laboratory of Inorganic Chemistry, Department of Chemistry, University of Athens, GREECE.



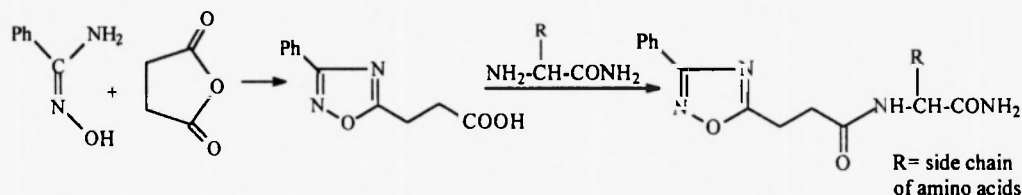
Scheme 1. (i) NaH, PhH, r.t., (ii) H₂NR₁, absolute ethanol, reflux 2.5 h. (iii) H₂NCH₂CH₂NH₂, absolute ethanol, reflux 2.5 h

SYNTHESIS OF NEW 1,2,4-OXADIAZOLES-DERIVED DIPEPTIDOMIMETICS, A POTENTIAL CLASS OF ANTIINFLAMMATORY DRUGS

R. F. Vieira¹, D. J. Brondani¹, A. R. de Faria¹, R. M. Srivastava² & A. C. Lima Leite^{1*}

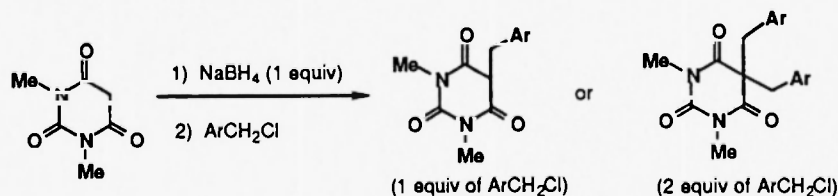
¹Laboratório de Planejamento, Avaliação e Síntese de Fármacos - LABSINFA, Laboratório de Tecnologia Químico-Farmacêutica - Departamento de Farmácia - CCS Universidade Federal de Pernambuco - Rua Prof. Artur Sa S/N, CDU, 50740-520, Recife-PE-Brasil ²Departamento de Química Fundamental - CCEN - UFPE - e-mail: acb02@elogica.com.br

A new series of 1,2,4-oxadiazoles containing pseudopeptide moiety as the side chain has been synthesized in good yield using a strategy of peptide synthesis.



SODIUM BOROHYDRIDE MEDIATED BENZYLATION OF 1,3-DIMETHYLBARBITURIC ACID

Lucjan Strekowski* and Mohamed A. Ismail, Department of Chemistry, Georgia State University, Atlanta, Georgia 30303, USA; Hanafi H. Zoorob, Faculty of Science, Al-Mansoura University, Al-Mansoura, Egypt

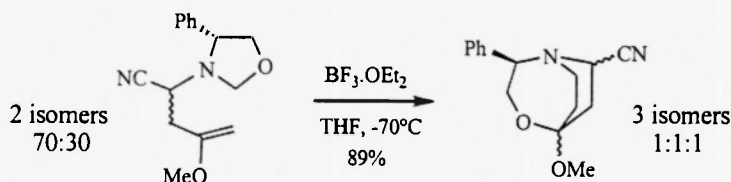


STEREOCHEMICAL EFFECTS IN THE EFFICIENT CYCLISATION OF *N*-(1-CYANO-3-METHOXY-BUT-3-ENYL)-4-PHENYLOXAZOLIDINE

David J. Aitken,* Laure Besson, Karolin Partogyan-Halim, Henri-Philippe Husson

Laboratoire de Chimie Thérapeutique-CNRS, Faculté de Pharmacie, Université Paris V, 4 avenue de l'Observatoire, 75270 Paris cedex 06, France, and

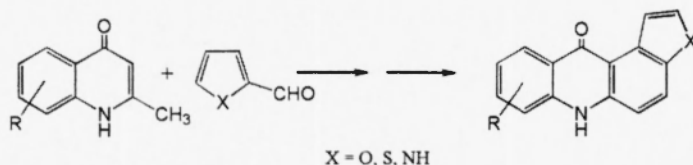
Laboratoire SEESIB-CNRS, Département de Chimie, Université Clermont-Ferrand II, 24 avenue des Landais, 63177 Aubière cedex, France



Conformational preferences may determine the stereochemical results

SYNTHESIS OF FURO- THIENO- AND PYRROLO- [3,2-a]ACRIDONES

S. Thamarai Selvi and P.S. Mohan, Department of Chemistry, Bharathiar University, Coimbatore-641 046, INDIA

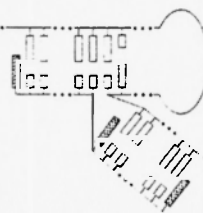


MULTIPLE LID STABILIZATION OF THE DNA THREE-WAY JUNCTION. INSERTION OF *N*³-(1-PYRENYLMETHYL)THYMIDINE.

Sherif A. El-Kafrawy and Erik B. Pedersen*

Department of Chemistry, University of Southern Denmark, Odense University, DK-5230 Odense M, Denmark.

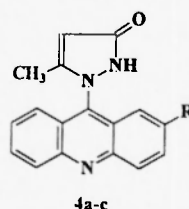
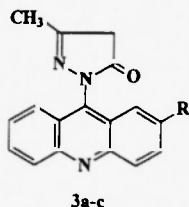
The DNA three-way junction is stabilized when stacking moieties cover the inclined arm at the branch point and the free ends of the duplex arms. *N*³-(1-pyrenylmethyl)thymidine, used as the stacking moiety, is easily synthesized by *N*³-alkylation of adequately protected thymidine.



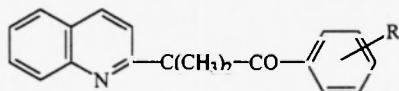
1-(ACRIDIN-9'-YL)-PYRAZOLIN-3 AND 5-ONES. A NEW CLASS OF HETEROCYCLES WITH POTENTIAL BIOLOGICAL ACTIVITY.

Ioan Cristea^a, Mariana M. Popovici, Maria T. Mendel^b, Ioan Silaghi-Dumitrescu^a and Erika Kozma^a^aDepartment of Organic Chemistry, "Babeş-Bolyai" University, Str. Arany Janos 11, 3400 Cluj-Napoca, Romania^bChemical and Pharmaceutical Research Institute, Str. Fabricii 126, 3400 Cluj-Napoca, Romania

1-(Acridin-9'-yl)-3-methylpyrazolin-5-ones **3a-c** were synthesised by cyclocondensation of 9-hydrazinoacridine derivatives and 1-(acridin-9'-yl)-5-methylpyrazolin-3-ones **4a-c** respectively, by arylation of 3-methyl-5-pyrazolone with 9-chloroacridine derivatives.



SYNTHESIS AND NMR SPECTRA OF 2-METHYL-2-QUINOLIN-2-YL-PROPIOPHENONES

Ryszard Gawinecki,^{**} Erkki Kolehmainen,^b Borys Osmialowski,^a Peter Palkovič^c and Maija Nissinen^b^a Department of Chemistry, Technical and Agricultural University, Seminaryjna 3, PL-85-326 Bydgoszcz, Poland;^b Department of Chemistry, University of Jyväskylä, P.O. Box 35, FIN-40351 Jyväskylä, Finland;^c Department of Chemistry, Comenius University, Mlynská dolina CH 2, SK-842 15 Bratislava, Slovakia

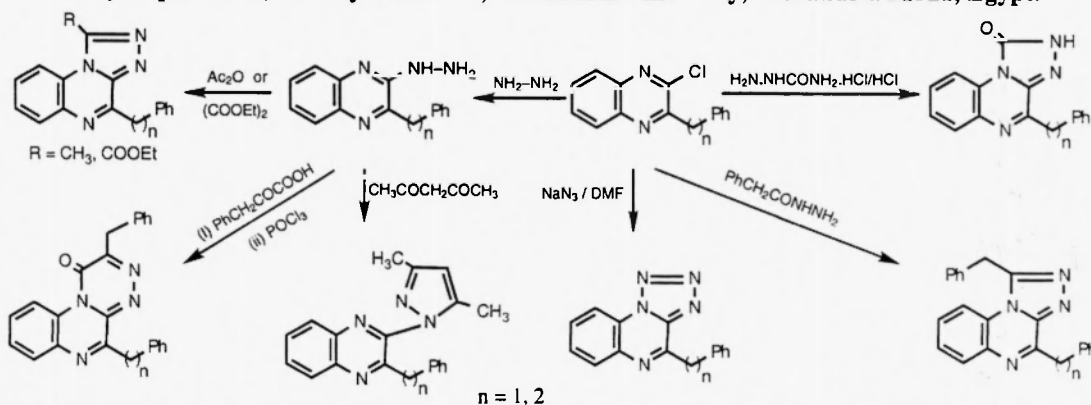
R = 4-CF₃, 3-CF₃, 3-Cl, 3-F, 4-Cl, 4-Br, 3-OMe, 4-F, H, 3-Me, 4-Me, 4-OMe

A series of 2-methyl-2-quinolin-2-yl-propio-phenones was synthesized. Structure of these compounds was studied by IR, ¹H, ¹³C and ¹⁵N NMR and X-ray methods.

SYNTHESIS AND ANTIMICROBIAL ACTIVITY OF CONDENSED AND UNCONDENSED QUINOXALINES

A. M. El Massry

Chemistry Department, Faculty of Science, Alexandria University, Alexandria 21321, Egypt.

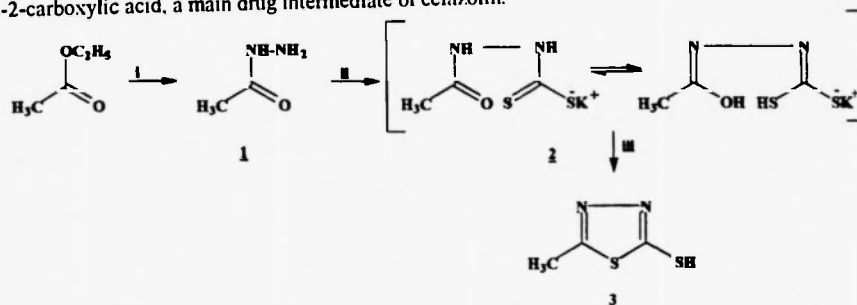


Heterocycl. Commun. 5 (1999) 565-568

Ranesh Chandra* and Narendra N. Ghosh

Ranesh Chandra* and Narendra N. Ghosh
Dr. B.R. Ambedkar Center for Biomedical Research, University of Delhi, Delhi-110007, India.

A simple convenient method of synthesis of 2-mercapto-5-methyl-1,3,4-thiadiazole (MMTD) is described which was then converted into 7-Amino-3-[[5-methyl-1,3,4-thiadiazole-2-yl]thio]methyl]-8-oxo-5-thia-1-azabicyclo[4.2.0]-oct-2-ene-2-carboxylic acid, a main drug intermediate of cefazolin.

Heterocycl. Commun. **5** (1999) 569-576

SYNTHESIS OF 4-SUBSTITUTED PYRIDIN-2(1H)-ONES, PYRIDINE-2(1H)-THIONES, RELATED DERIVATIVES AS ANALOGUES OF CARDIOTONIC DRUG OF MILRINONE

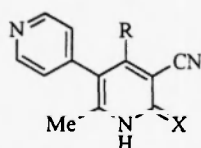
A. Krauze^{*a}, R. Vitolina^a, V. Garalene^b, H.-J. Jansch^c, G. Duburs^a

³Latvian Institute of Organic Synthesis, Riga, Aizkraukles 21, LV-1006, Latvia

^bLithuanian Institute of Cardiology, Kaunas, Lithuania LT-3007

^cArzneimittelwerk Dresden, Stadtbetrieb des Pharmazeutischen kombinats Germed, Dresden, BRD

A convenient method of synthesis of 4-substituted 5-(4'-pyridyl)pyridine-2(1H)-ones and -thiones was elaborated by Michael reaction of 4-pyridylacetone and 2-cyano-3-R-acrylamides (thioamides) with subsequent heterocyclization, dehydration and dehydrogenation. Their cardiovascular activity in vitro was screened.



X = O; R = 2-F₂HCOC₆H₄; 4-ClC₆H₄; 4-Pyridyl;

X = S; R = 4-Pyridyl

Heterocycl. Commun. 5 (1999) 577-584

Synthesis of substituted quinolines and heterocyclo[x,y-c]quinolines by the nucleophilic substitution and rearrangements of 4-chloro-2-methyl-3-nitroquinolines

A. I. khodair, M. M. A. Abbasi, El-Sayed I. Ibrahim, A. H. Soliman and El-Sayed H. El-Ashry** Chemistry*

Department, Faculty of Science, Suez Canal University, *Tanta University, and **Alexandria university, Egypt.

