Synthesis and Cytotoxicity of 3,4-Diaryl-2(5H)-Furanone Derivatives

Kim Yong^o, Nam NguyenHai, You YoungJae, Ahn ByungZun

College of Pharmacy, Chungnam National University, Taejon 305-764, Korea

2(5H)-Furanone is a common moiety incorporated in a number of drugs with diverse biological activities such as antitumor, antifungal, antibacterial and antiinflammatory. One of them, a 3,4-diaryl-2(5H)-furanone analogues, in which the two aromatic rings are tethered directly into the 2(5H)-furanone ring, a biomoiety found in a number of drugs with diverse biological activities were synthesized and evaluated against for their cytotoxicity in a small panel of cancer cell lines. Four of ten compounds in this series, e.g. 3-(3,4,5-trimethoxyphenyl)-4-(4-methoxyphenyl)-, 3-(3,4,5-trimethoxyphenyl)-4-(3-hydroxy-4-methoxyphenyl)-, 3-(3,4,5-trimethoxyphenyl)-4-(3-amino-4-methoxyphenyl)-, and 3-(3,4,5-trimethoxyphenyl)-2(5H)-furanones, were found to have potent cytotoxic activities with ED₅₀ values of less than 20 nM in most of the cell lines tested.

[PD1-30] [10/19/2001 (Fri) 14:00 - 17:00 / Hall D]

Epoxidation of [25R]-1,4,6-Spirostatriene-3-one

Kim haksoon^o, Ma EunSook

Catholic University of Daegu

The [25R]-5-spirosten-3β-ol was oxidized with dichlorodicyanobenzoquinone(DDQ) to give [25R]-1,4,6-spirostatriene-3-one. Treatment of the triene with alkaline hydrogen peroxide afforded the [25R]-1,2-epoxy-4,6-spirosta

dien-3-one. Epoxdation of the triene with m-chloroperoxybenzoic acid produced the [25R]-6,7-epoxy-1,4-spirostadien-3-one. These products were reduced with litium metal and ammonium chloride in liquid ammonia, to yield [25R]-6-spirosten-1,3-diol and [25R]-1-spirosten-3,6-diol, respectively. [25R]-1,2-epoxy-4,6-spirostadien-3-one was reduced with lithium metal in absolute ethanol to give [25R]-1-ethoxy-4,6-spirostadien-3-one.

[PD1-31] [10/19/2001 (Fri) 14:00 - 17:00 / Hall D]

Aminoacyl adenylate analogues as inhibitors of aminoacyl-tRNA synthetase

Jang SunYoung^o, Park SangWoo, Lee YeonSook, Kang TaeHee, Song NanKyue, Kwak JinHwan, Kim SeokIn, Kim Sunghoon, Choi Sooyoung

ImaGene Co., Ltd. Central Research Institute, 81508 Sung Kyun Kwan University, Suwon, Institute for Biomedical Research, Han Dong University, Pohang, College of Pharmacy, Seoul National University, Seoul

The emergence of drug resistant *Staphylococcus aureus* poses a significant health treat to human. Thus there is a critical need to develop new antimicrobial agents with novel mode of action. Aminoacyl-tRNA synthetases(ARSs) are essential in protein biosynthesis, catalyzing the attachment of amino acids to their cognate tRNA prior to the ribosome. Selective inhibition of bacterial ARS has proved to be a successful strategy for the production of antibacterial compounds. Pseudomonic acid(generic name: mupirocin) is a potent inhibitor of isoleucyl-tRNA synthetase. Recently, structure-activity relationship