Eur. J. Med. Chem. 37, 659-669 (2002).

## Preparations of heterospirostanols and their pharmacological activities

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(3β,20*S*,22*S*,25*R*)-22-Thiospirosol-5-en-3-ol **1** and (3β,20*S*,22*S*,25*R*)-22-seleno-spirosol -5-en-3-ol **2** were prepared from diosgenin **3** via 26-iodopseudodiosgenin **4** as a key intermediate.Diosgenone 5, solasodinone 6, (20S,22S,25R)-22-thio-spirosol-4-en-3-one 7, (20*S*,22*S*,25*R*)-22-selenospirosol-4-en-3-one **8** and(20*R*,22*S*,25*R*)-spirosol-4-en-3one 9 were prepared by Oppenauer oxidation of 3, solasodine 10, 1, 2 and  $(3\beta, 20R, 22R, 25R)$ -spirosol-5-en-3-ol 11, respectively. Oxidations of 5 and 6 with 2,3-dichloro-5,6-dicyano-1,4-benzoquinone (DDQ) provided corresponding dienone products, (20S,22S,25R)-spirosol-1,4-dien-3-one **12** and (20S,22S,25R)-22-thiospirosol -1,4-dien-3-one 13, respectively, while oxidation of 14 (C-20 diastereoisomer of 5) dienone 21-exo gave no product but vinyl product 15. 26-Thioacetylpseudodiosgenone 16 and 26-cyanoselenopseudodiosgenone 17 were prepared by treatment of 26-iodopseudo-diosgenose 18, which was obtained by Oppenauer oxidation of 4, with potassium thioacetate and potassium selenocyanate, respectively. Compounds 5 and 14 exhibited more than 80% inhibitions in INF- $\gamma$  productions at 10.0  $\mu$ M. Compounds 10 and 17 showed cytotoxic activities (IC50 = 6 and 5  $\mu$ M, respectively) against cancerous HCT 116 cell lines. 26-Cyanoselenopseudodiosgenin 19 and 17 had antiurease activities  $(IC50 = 12.4 \text{ and } 11.4 \mu M, \text{ respectively}), in which only the latter showed an$ inhibition zone (mean zone diameter = 12.2 mm) formed by *Bacillus subtilis 168 trp.*