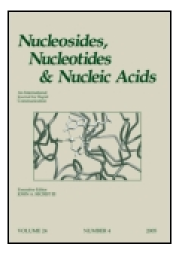
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A CONVENIENT APPROACH TO N-3 ALKYLATION OF 9-SUBSTITUTED GUANINES

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ABSTRACT: Aryl or *tert*-butyl substituent in the 6 position of 3,9-dihydro-3-[(2-hydroxy-ethoxy)methyl]-9-oxo-6-R-5*H*-imidazo[1,2-*a*]purine 1 directs the benzylation reaction partly into N-4 position to give 3. Cleavage of the third ring of 3 gives 3-benzylacycloguanosine 5, a first 3-aralkilo-9-substituted guanine.

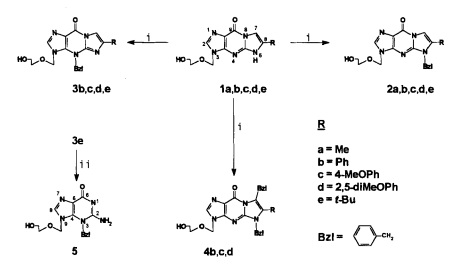
No direct N-3 alkylation of monomeric 9-substituted guanine has been noted so far. We have previously found that N-3 methylation of guanine moiety in guanosine, 2'-deoxyguanosine and 9-[(2-hydroxyethoxy)methyl]guanine (acycloguanosine) can be accomplished by a conversion into tricyclic $1,N^2$ -(prop-1-ene-1,2-diyl) derivatives, i.e. 3,9-dihydro-6-methyl-9-oxo-5*H*-imidazo[1,2-*a*]purine system. This tricyclic system which undergoes high yield N-4 methylation by means of cyclopropanation reagent¹ is easily split to 3-methylguaninederivatives.²⁻⁴ The type of alkylation reagent however, does not allow for a variety of N-3 substituents.

We present now further application of the tricyclic modification based on the observation that a substituent in the 6 position has an impact on the site of substitution with alkyl and aralkylhalides under alkaline conditions. Our attention was mainly focused on benzylation reactions of 3,9-dihydro-3-[(2-hydroxyethoxy)methyl]-9-oxo-6-R-5*H*-imidazo[1,2-*a*]purines 1. When 6-methyl derivative 1a in dry DMF was treated with K₂CO₃ followed by benzyl bromide, at room temperature for 5 hours, N-5 benzyl substituted 2a resulted as a single product in over 80 % yield. Under analogous conditions the presence of 6-aryl groups (R = phenyl, 4-methoxyphenyl,2,5-dimethoxyphenyl in compounds 1 b, c, d,

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respectively) led to a mixture of products which were separated on silica gel column in chloroform - methanol gradient followed by rechromato-graphy of mixed fractions in ethyl acetate - ethanol. In addition to preponderant N-5 derivatives2 b, c, d; N-4, 3 b, c, d (2-18 %); N-5,7, 4 b, c, d (17-25 %) and 7,0⁹ (3 %) substituted products were isolated. In the case of 6-*tert*-butyl substrate 1e, the N-4 product 3e became the major one (36 %). The structures of the compounds were assigned on the basis of ¹H and ¹³C NMR, UV and mass spectra. ¹H and ¹³C chemical shift values of benzyl methylene signal were particularly diagnostic (δ_{H} , δ_{C} ; N-4 : ~5.9, ~49.0; N-5 : ~5.4, ~45.9; N-5,7 : ~5.2, 4.4, ~45.7, 29.7). The removal of the third ring of 3e was accomplished with 30 % aq hydrogen peroxide in DMF at room temperature for 7 days. Crude product was acetylated with acetic anhydride in pyridine, separated by silica gel column chromatography in toluene - ethanol gradient and deblocked with methanolic ammonia to give 3-benzylacycloguanosine 5. Similarly 3-(4-nitrobenzyl)acycloguanosine was prepared.

The N-4 directing effect of 6-substituents was weaker in the case of alkylation. On methylation and ethylation of 1b the yields of N-4 substituted products were up to 10 %; 1e did not react at N-4.



i: $C_6H_5CH_2Br$, DMF dry, K_2CO_3 , r.t.; ii: 30% H_2O_2 , DMF, r.t.

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