SYNTHESIS OF L-2-DEOXYTHREOSE NUCLEOSIDES AS POTENTIAL ANTIVIRAL AGENTS

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Several nucleoside analogs are well-known to inhibit the reverse transcription (RT) process in HIV by terminating the synthesis of a proviral DNA strand. Recently, Herdewijn and coworkers showed that some L-2-deoxythreose nucleoside phosphonates (a; Figure 1) selectively inhibit HIV without affecting human DNA synthesis. These nucleoside analogues require two additional phosphorylation steps by cellular kinases before they are incorporated in the viral genome and lead to chain termination. It is believed that a cis orientation of the base and the 3'-substituent is a structural requirement for optimal conversion to the diphosphate. Inspired by these findings, we decided to

explore the antiviral properties of the threose-based nucleoside analogues (\mathbf{b} ; Figure 1), which are anticipated to have superior bioavailability compared to the parent phosphonates. Remarkably, such α -L-2'-deoxythreofuranosyl analogues (exhibiting a 1'R,3'R configuration) have not been reported before.

The synthesis of the desired T, U, C and A analogs (Scheme-1), involving a Vorbrüggen coupling and a Barton-McCombie deoxygenation, will be described.

Scheme-1

HO OAC ROOAC ROOAC ROOAC ROOAC B* = T, U,
$$C^{Ac}$$
, A^{Bz} B = T, U, C, A R = Benzyl (Bn), ρ -Methoxybenzyl (PMB)

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