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A. A GENERAL PROCEDURE FOR THE CONVERSION OF RIBONUCLEOSIDES TO 2'-DEOXYNUCLEOSIDES

B. A FACILE METHOD FOR
DETERMINATION OF ANOMERIC CONFIGURATION

bу

O JOHN SCOTT WILSON

A THESIS

SUBMITTED TO THE FACULTY OF GRADUATE STUDIES AND RESEARCH

ÎN PARTIAL FULFILMENT OF THE REQUIREMENTS FOR THE DEGREE

OF DOCTOR OF PHILOSOPHY

IN CHEMISTRY

EDMONTON, ALBERTA FALL, 1981

THE UNIVERSITY OF ALBERTA

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A GENERAL PROCEDURE FOR THE CONVERSION OF RIBONUCLEOSIDES TO 2'-DEOXYNUCLEOSIDES

> A FACILE METHOD FOR DETERMINATION OF ANOMERIC CONFIGURATION

submitted by JOHN SCOTJ WILSON in partial fulfilment of the requirements for the degree of Doctor of Philosophy in Chemistry.

ABSTRACT

A general and efficient procedure for the preparation of 2'-deoxynucleosides has been developed. Selective protection of the 3'- and 5'- hydroxyl functions of the ribonucleosides was achieved through preparation of $3',5'-\underline{0}-(1,1,3,3-tetraisopropyldisilox-1,3-diyl)$ nucleoside derivatives. The 2'-hydroxyl group of the blocked nucleoside was functionalized under mildly basic conditions using phenyl chlorothionocarbonate and $4-\underline{N}$, \underline{N} -dimethylaminopyridine in acetonitrile. Treatment of the 2'-phenylthionocarbonate derivative obtained with $tri-\underline{n}$ -butylstannane in the presence of a free radical initiator at 80°C gave the 2'-deoxynucleoside derivative in excellent yield. Deprotection was effected under neutral conditions using tetra- \underline{n} -butylammonium fluòride to give the 2'-deoxynucleoside. High yields of 2'-deoxyadenosine, 2'-deoxyguanosine, 2'deoxyuridine, 2'-deoxytubercidir and 2'-deoxytoyocamycin were realized using this reaction sequence. Application of the procedure to other secondary hydroxyl groups was equally successful as steroid and carbohydrate examples were also deoxygenated in high yield. Acid sensitive substrates were amenable to this method of deoxygenation as demonstrated by the synthesis of methyl 2-deoxy-3, $5-\text{di}-\underline{0}-\underline{p}-\text{toluyl-}\beta-D-\underline{erythro}-\text{pentofuranoside}$ from methyl $\beta-D$ -ribofuranoside.

The similarity and consistency of the 1H NMR spectra *of the 3',5'- $\underline{0}$ -(1,1,3,3-tetraisopropyldisilox-1,3-diyl)= nucleosides indicated that a relatively fixed sugar conformation existed in these compounds. A representative number of these 3',5' blocked derivatives were prepared from β -ribo, α -ribo, β -arabino and α arabinonuclaside substrates. It was found that for β-ribonucleoside derivatives, the spin-spin coupling values for H-1' to H-2' were generally less than 1.5 Hz. For α -ribo and β -arabinonucleoside derivatives, spinspin coupling values for H-1' to H-2' were generally greater than 3.5 Hz. The ^{1}H NMR spectra of the $\alpha\text{-arabino-}$ nucleoside derivatives unexpectedly revealed a large H-1' to H-2' coupling value. This was thought to arise from a change in the sugar conformation of these derivatives relative to the other examples of the 3',5'-0-(1,1,3,3-tetraisopropyldisilox-1,3-diyl)nucleosides studied. A single crystal X-ray analysis of 3',5'-0-(1,1,3,3-tetraisopropyldisilox-1,3-diyl)cytidine was determined to verify assumptions made from NMR spectra on the sugar conformation of these derivatives.

ACKNOWLEDGMENTS

I am grateful to all faculty members, non-academic staff and colleagues who have made my stay at the University of Alberta a meaningful experience. In particular, Dr. M.J. Robins gave me continual support and encouragement and was a constant source of ideas and suggestions.

Members of the support staff of the Chemistry Department are greatly appreciated for their unflagging assistance and dedication. I am especially grateful to Ms. Diane Dowhaniuk for her patience in typing this thesis.

Financial support from the National Research Council of Canada and the University of Alberta is gratefully acknowledged.

Finally, I would like to acknowledge the people in Lethbridge who gave me so much guidance and support. The members of the Chemistry Department of the University of Lethbridge were very good to me and will always be fondly remembered. My parents and grandparents placed no obligations upon me and were always a source of encouragement and affection. Ms. Darrell Sturrock was an inspiration in her strength and compassion.

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I. INTRODUCTION

A. <u>General Introduction and Survey of 2'-Deoxy-nucleoside Synthesis</u>.

The fundamental unit of higher organisms, the cell, is a complex structure of interconnected elements, each having specific functions. The nucleus is the control center within the cell and contains all genetic information essential for the integrated operation of cellular components. In 1871, Fredrich Miescher isolated a material from the nuclei of pus cells which he called "nuclein". This material was subsequently referred to as "nucleic acid" by Altmann and was found to be common to cells from a number of sources.

The nucleic acids provide the means for the storage and transmission of genetic information utilized in the synthesis of all cellular constituents. These macromolecules are comprised of polymeric chains of hucleotides linked by phosphodiester bonds. The nucleotide unit is composed of a purine or pyrimidine base, a furanose sugar moiety and a phosphoric acid residue. Levene and Jacobs introduced the term nucleoside to describe the carbohydrate derivatives of purines and pyrimidines isolated from the alkaline hydrolysis of yeast ribonucleic acids. This term has been expanded to include all of the compounds of synthetic or natural

 $\beta - \underline{D} - RIBOFURANOSYL$

2'-DEOXY-B-D-RIBOFURANOSYL
(2'-DEOXY-B-D-ERYTHROPENTOFURANOSYL)

origin which contain a heterocyclic base attached to the $\mathcal{L}\text{-l}$ position of a sugar.

There are two types of nucleic acids, the constituents of which differ primarily in the carbohydrate residue. In ribonucleic acid (RNA) the sugar moiety is D-ribose and in deoxyribonucleic acids (DNA) the sugar is 2-deoxy-D-ribose (2-deoxy-D-erythro-pentose). Leveneand co-workers 4.5.6 were responsible for the characterization of the sugar residues of RMA and DNA through their exhaustive studies from 1909 to 1933.

Five heterocyclic bases are ubiquitous to the nucleic acids and were first identified by Kossël and co-workers through acid hydrolysis of nucleic acids. The purines adenine $(\underline{1})$ and guanine $(\underline{2})$ and the pyrimidine cytosine $(\underline{3})$ are common to both DNA and RNA. Uracil $(\underline{5})$ is found only in RNA, while its 5-methyl derivative thymine $(\underline{4})$ is the complementary pyrimidine contained in DNA. The corresponding nucleosides in RNA are adenosine $(\underline{6})$, guanosine $(\underline{7})$, cytidine $(\underline{8})$ and uridine $(\underline{9})$ and in DNA are 2'-deoxyadenosine $(\underline{10})$, 2'-deoxyguanosine $(\underline{11})$, 2'-deoxycytidine $(\underline{12})$ and thymidine $(\underline{13})$.

The DNA present in the nucleus of all eukaryotic cells contains the information necessary for the biosynthesis and metabolism of the cell. Through the specific ordering of the nucleic acid units in these

macromolecules a unique code is established. This code is transcribed to smaller mRNA molecules which act as templates in code translation for subsequent protein (enzyme) synthesis. Chemical inhibition of the blo-synthesis of nucleic acids is one major basis of chemotherapy. Bacterial cells and tumor cells divide much more frequently than the cells of the host organism and thus are expected to be affected more severely by an antimetabolite. Routes which are both general and efficient for the synthesis of ribonucleoside and 2'-deoxynucleoside antimetabolites are of considerable importance in this area.

The position of the sugar-base attachment in nucleosides obtained from nucleic acids was first established by ultraviolet spectral comparisons. 8,9,10 Unequivocal conformation of the location of the sugar residue was presented by Todd and co-workers 11 through the synthesis of $^{9-\underline{D}}$ -mannopyranosyl adenine (15). Oxidation of this compound with periodate gave a dialdehyde (14) identical to that obtained from adenosine (6).

Todd and co-workers 12 also confirmed the assignment of the β configuration of nucleosides by demonstration of the formation of <u>cis</u>-intramolecularly linked 5'-cyclo-nucleosides derived from cytidine (8) and adenosine (6). Total synthesis of the two major purine ribonucleosides was achieved by Todd's group as the final structure proof. 13 The historical development of the identification and characterization of the nucleosides has been reviewed extensively. $^{14-19}$

The first synthesis of a nucleoside was reported by Fischer and Helferich 20 in 1914 through the coupling of the silver salt of theophylline with tetra-0-acetyl- α -0-glucopyranosyl bromide. The glycosylpurine derivative obtained was later shown to be 7-substituted. This general coupling procedure was utilized by Todd and co-workers 13 in 1948 to synthesize adenosine (6) and guanosine (7), using 2,3,5-tri-0-acetyl-0-ribofuranosyl bromide (17) as the sugar element. The silver salt of 2,8-dichloroadenine (16) was used as the purine component since both the natural purine nucleosides could be made from the 2,8-dichloroadenosine derivative (18) obtained.

Attempts to condense silver salts of oxopyrimidines with halo-sugar derivatives as described above did not give the expected pyrimidine nucleosides, but rather the $\underline{0}$ -glycosylpyrimidines. At the $\underline{0}$ -glycosylpyrimidines.

SCHEME II

$$NH_2$$
 NH_2
 NH_2

demonstrated that 2,4-dialkoxypyrimidines could be employed as coupling reagents in synthesis of pyrimidine nucleosides. The reaction of tri-0-acetyl-0-ribo-furanosyl bromide (17) with 2,4-diethoxypyrimidine (19) provided an intermediate (20) from which both uridine (9)²² and cytidine (8)²³ were obtained.

SCHEME III

Until 1948 the nucleic acids were thought to be composed of the four basic nucleoside monomers (6-13) only. The discovery of 5-methylcytosine in a sample of calf thymus DNA by Hotchkiss²⁴ prompted a surge of reinvestigation. However, it was the isolation of the first nucleoside antibiotic, cordycepin (21), from Cordyceps militaris by Cunningham et al. 25 in 1951 that initiated the boom of chemical research in the nucleoside

toxic agent which demonstrates a broad spectrum of activity against bacterial strains through the inhibition of purine biosynthesis. The 5'-triphosphate derivative (22) has also been shown to be tytotoxic through the inhibition of nucleic acid biosynthesis. 26

The intense interest generated by the identification of a new class of antibiotics inspired researchers to devise efficient syntheses of potential biologically

active compounds from readily obtainable substrates. The employment of chloromercuri derivatives of purines by Davoll and Lowy 27 in place of the silver salts used previously improved yields of coupling reactions significantly. It was later demonstrated 24 that mercuri derivatives of pyrimidines were also effective condensing reagents for the preparation of pyrimidine nucleosides. This led to the first coupling-based synthesis of a 21-deoxynucleoside, thymidine (13), by Shaw and Warrener²⁹ in 1958. Baker and co-workers³⁰ reported the first successful synthesis of 2'-deoxyadenosine in the following year. The starting material used was 1,2-di-0-acety1-5- $\underline{0}$ -methoxycarbony1-3- $\underline{0}$ -(ptoluenesulfonyl)- \underline{D} -xylofuranose ($\underline{23}$) which was converted to the 1-halo derivative (24) prior to condensation with chloromercuri 6-benzamidopurine to give the blocked nucleoside (25). The epoxide (26) was formed upon treatment of (25) with sodium methoxide in methanol. Reaction of (26) with sodium ethylmercaptide afforded the 3'-ethylthic nucleoside (27). Treatment of (27)with thionyl chloride gave the 3'-chloronucleoside (28). Exposure of (28) to aqueous base resulted in a mixture of 2'-(30) and 3'-(27) ethylthic derivatives. episulfonium ion intermediate (29) was proposed to explain this migration. Desulfurization of (30) gave 2'-deoxyadenosine (10) in a overall yield of 0.5% from (23).

S C H E M E IV

The general methods of sugar-base coupling utilized successfully for the synthesis of ribonucleosides have limitations when applied to 2-deoxynucleosides. di-0-acyl-2-deoxyribofuranosyl chlorides are extremely labile unless stabilized with appropriate blocking groups. 31,32 Anomeric mixtures of the 2'-deoxynucleoside are always obtained in the condensation reactions. The mixtures of anomers obtained in these coupliffgs can be difficult to separate and are much more troublesome if the base component of the nucleoside contains an acidic function. This precludes the use of the Dekker 50 anion exchange chromatography method. Furthermore, the normally unwanted α anomer is frequently an equivalent or predominant product. Numerous reports of base-sugar couplings using various procedures for the synthesis of the naturally occurring 2'-deoxynucleosides and related analogues have appeared in the literature. 31-49

The stereoselectivity of coupling reactions is dependent on many factors. The method of condensation, choice of solvents, temperature and catalysts can all affect the ratio of anomers obtained. Through mechanistic analysis of reactions involving metal salts of purines, Baker proposed a trans rule which postulated that the "condensation of a heavy metal salt of a purine with an acylated glycosyl halide will form a nucleoside with a C-1 to C-2 trans configuration." This

behavior had been observed in pyrimidine synthesis as well. 52 The experimental facts were rationalized by the intervention of an acyloxonium ion (32) formed by participation of the adjacent trans 2-acyloxy group of one anomer (31) of the glycosyl halide. Attack of the heterocyclic base at C-l of (32) gives the trans 1,2-nucleoside (34). Presumably the cis anomer (33) can react directly by $S_{N}2$ displacement to give the same product (34).

SCHEME V

<u>33</u>

Sugar substrates which have the necessary participating group at the C-2 position normally give exclusive formation or high ratios of the predicted anomer. Conversely, those substrates which lack this participating C-2 group, such as 2-deoxyglycosyl halides, give rise to anomeric mixtures upon condensation with a base.

Bardos and co-workers 53 reported that the anomeric ratio of products obtained from reacting pertrimethylsilylated thiopyrimidine bases with 2'-deoxyglycosyl halides could be controlled under certain conditions. By using the purified α anomer of the sugar component and azeotropically removing the trimethylsilyl chloride (TMSC1) as it was produced, excellent yields of a pure β anomer were realized in this silyl Hilbert-Johnson reaction. However, other workers have found that parallel results were not obtained when different bases were coupled. 45

Ryan, Acton and Goodman 54 have reported the synthesis of 2'-alkylthiopurine nucleosides through a condensation reaction using an alkyl 1-thio- α -D-arabinofuranoside as the sugar precursor. The stereospecificity of the reaction was thought to result from the formation of an episulfonium ion between C-1 and C-2. Trip 55 attempted to extend this result to achieve a fully stereoselective synthesis of B-2'-deoxynucleosides. However the poor yields obtained in the reductive removal of the alkyl-

thio group limited the usefulness of this method.

Since ribonucleosides are readily available from natural and synthetic sources as β anomers, they are attractive substrates for 2'-deoxynucleoside synthesis. However, there have been major difficulties involved in obtaining the 2'-deoxynucleosides or suitable precursors such as 2'-halo or 2'-mercapto derivatives. Direct displacement of leaving groups by an external nucleophile at the 2'-position of ribonucleosides is hindered sterically by the base and electronically by the electron deficient C-1' and β oriented base dipole. Furthermore, since the 2'- and 3'-hydroxyl groups are very similar in reactivity, selective functionalization is very difficult. Any intermediate which has both positions "symmetrically" activated will give a predominance of the 3'-derivative from external nucleophilic attack owing to the noted inhibition of reaction at C-2'.

Todd and co-workers 56 were the first to report the synthesis of a pyrimidine 2'-deoxynucleoside from the corresponding ribonucleoside. They found that treatment of 5'-0-acetyl-2'-0-p-toluenesulfonyluridine (35) with sodium iodide gave the 2'-iodo derivative (37) with retention of the <u>ribo</u> configuration. Subsequent hydrogenolysis and deacetylation gave 2'-deoxyuridine (38). It was shown that the displacement reaction proceeded via an 0-2+2'-anhydropyrimidine intermediate (36). Al-

though Todd obtained an overall yield of less than 10% for this conversion of uridine to 2'-deoxyuridine, the novel method attracted the interest of numerous groups. 57-59

R = Ac Ts = p-toluénesulfonyl

Efficient outes for synthesis of the cyclonucleoside intermediates were developed which utilize the 2',3'-diol, structure and hence eliminate the problem of selectively functionalizing two very similar hydroxyl groups. The cyclic 2',3'-0-carbonate (39) and 2',3'-0-thionocarbonate (40) derivatives have been used by several groups 60-64 to give excellent yields of these pyrimidine 0-2+2'-cyclonucleosides. The formation of 2',3'-acetoxonium ion (41), accessible via a variety of procedures, also leads to high yields of the 0-2+2'-cyclonucleosides in the pyrimidine series. 65,66

R'- COPh

comprehensive review of pyrimidine cyclonucleoside formation has been published recently. 67

Holý reported a 56% yield of thymidine from 5-methyluridine utilizing the following reaction sequence. The 0-2+2'-cyclonucleoside (43), obtained from 5-methyluridine (42) via the cyclic 2',3'-0-carbonate derivative (39) was benzoylated to give the 3',5'-di-0-benzoyl-0-2+2'-cyclo-5-methyluridine (44). Reaction of (44) with dry hydrogen chloride in dimethyl-formamide gave the 2'-chloro derivative (45). Mild reduction of (44) using tri-n-butylstannane followed by debenzoylation gave thymidine (13).

Attempts to devise efficient syntheses of purine 2'-halo and 2'-deoxynucleosides have proved to be much more difficult. Ikehara and Tada⁶⁹ effected purine 5-8+2'-cyclonucleoside formation by treatment of the 8-bromo derivative (46) with thiourea. Intramolecular displacement of the total group at C-2' gave the desired product (47).

It was found, however, that Raney nickel desulphurization was very inefficient and low yields of 2'-deoxy-adenosine ($\underline{10}$) were obtained. Analogous reactions involving 8-oxoadenosine ($\underline{48}$) also proved to be disappointing since the C-8-oxy group was extremely difficult to remove. 70

SCHEME VIII

Extensive work on the preparation of purine 2'- and 3'-halonucleosides via 2',3'-acatoxonium ion intermediates (41) has been pursued by both fatt and co-workers 71 and Robins and co-workers. Above as stated previously, the 2'-halo derivatives (49) are formed as minor products accompanying the predominant 3'-halo derivatives (50). In the case of adenosine, these procedures give ratios of (49) to (50) from approximately 1:9 to 1:6 depending on conditions used.

SCHEME IX

HO OH
$$\frac{(CH_3)_2CCOX}{OAc}$$
 RO OAC $\frac{6}{50}$ $\frac{6}{50}$

Attempts to synthesize purine 2'-deoxynucleosides through S_N^2 displacement have been disappointing. Sporns 74 treated 3'-0-methyl-5'-0-pivaloyladenosine with 5 equivalents of thionyl chloride in pyridine at 120°C for 1 hour to obtain a 16.5% yield of the 2'-chloro derivative. Ranganathan 75 reported that displacement of the trifluoromethanesulphonyl group was possible at the 2'-position of adenosine. The triflyl derivative

ŗ

 $(\underline{52})$ was prepared in 70% yield from the 3',5'-diprotected precursor $(\underline{51})$. Reaction with potassium thioacetate at room temperature gave the <u>arabino</u>-thio derivative $(\underline{53})$ in 55% yield. Raney nickel desulphurization gave the 2'-deoxyadenosine, although in low overall yield from $(\underline{51})$.

S C H É M E X

THP = TETRAHYDROPYRANYL

Tf = TRIFLUOROMETHANESULPHONYL

Recently Robins and Muhs ⁷⁶ reported the first successful conversion of a pyrrolo[2,3-d]pyrimidine nucleoside to its 2'-deoxy analogue. The antibiotic tubercidin (54) was converted to 2'-deoxytubercidin (55) in an eight step procedure that utilized a key episulfonium ion migration analogous to the Baker route to 2'-deoxy-adenosine. The 27% overall yield obtained ⁷⁶ exemplies the difficulty associated with the syntheses of purinetype 2'-deoxynucleosides.

<u>54</u>

<u>55</u>

B. Recent Developments Pertinent to 2'-Deoxynucleoside
Synthesis.

A general and efficient route for preparation of purine 2'-deoxynucleosides has been elusive. The use of S_N^2 displacement reactions have been limited due to the intrinsic complications of the molecules as discussed in Chapter I.A. The possibility of utilizing an S_N^1 type process is negated since cation formation at C-2' is precluded by the adjacent electron deficient anomeric carbon. However, generation of an anion at C-2' is possible. In studies on the chromous ion induced elimination of vicinal halo acetates, Moffatt and coworkers 77 reported an anomalous result for the cischalo acetate of uridine (56). A large amount of glycosidic cleavage of this normally stable nucleoside was observed upon treatment with chromous acetate.

Moffatt hypothesized that this occurred as a result of the formation of the C-2' radical, (57) which collapsed either to the 2',3'-olefin (58) or with glycosidic cleavage. However, $\operatorname{Hol}\hat{y}^{59}$ has demonstrated that the free radical mediated reduction of 2'-halouridines with $\operatorname{tri-n-butylstannane}$ proceeds to the 2'-deoxynucleoside in high yield without glycosyl cleavage. Adachiet al. 78 have studied the electrochemical reduction of (56). They also observed a significant proportion of glycosidic cleavage. The proposed mechanism in this

SCHEME XI

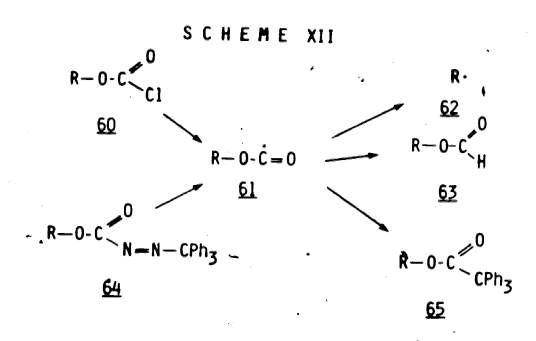
Elimination of the base to form the 1'-2' unsaturated sugar provides a plausible explanation. The similarity between these results of Moffatt⁷⁷ and Adachi⁷⁸ would suggest a common mechanism. This type of elimination of the heterocyclic base was observed also by Robins and Sporns.⁷⁹ They found that the generation of an anion at C-2' of an adenosine derivative using the dissolving metal reduction procedure of Ireland and co-workers⁸⁰ led exclusively to glycosidic cleavage.

Therefore, homolytic cleavage methods appeared to offer the only feasible approach to efficient deoxygenation of nucleosides at C-2'. This would be consistent with observations that deoxygenation of ribonucleotides to the corresponding DNA components effected by ribonucleotide reductase enzymes in nature are free radical mediated processes. 81

Photochemical deoxygenations of alcohols (as esters) to alkanes have been investigated by a number of groups recently. Pete and co-workers 82 reported that prolonged ultraviolet irradiation of carboxylic esters in aqueous hexamethylphosphoramide led to good yields of the corresponding alkanes. Application of this method to carbohydrates was successful for the deoxygenation of primary and secondary hydroxyl groups. 83,84 Other derivatives such as sulphonates and thioesters 86,87 have been photochemically reduced to the deoxygenated analogues. However, the strong absorbance of ultraviolet light by the heterocyclic base in nucleosides precludes this approach.

A chemical method was sought that would allow a sufficiently low transition state energy for the homolysis process to overcome the need for high temperatures. The possibility of decarbonation of an alkylcarbonyloxy radical (61) was examined since the generation of carbon dioxide should be energetically favourable. Kuivila and Walsh⁸⁸ had reported that such fragmentations did not occur unless the derived alkyl radical (62) was especially

stabilized. They found that formates $(\underline{63})$ were the major products from reaction of chloroformates $(\underline{60})$ with tri- \underline{n} -butylstannane at 110° C. Beak and Moje 89 obtained a 22% yield of toluene from benzylchloroformate and less than 1% cyclohexane from cyclohexylchloroformate using this method. Barton 90 generated radical $(\underline{61})$ through the photolysis of the azo-ester $(\underline{64})$, however, only recombination products $(\underline{65})$ were observed with no apparent loss of carbon dioxide.



More recently Jackson and co-workers 1 reported that treatment of primary and secondary chloroformates (60) with trivisopropylsilane in the presence of a radical initiator at 140°C led to good yields of alkanes. The greater strength of the silicon hydrogen bond (compared to the two hydrogen bond) makes this reagent less susceptible to hydrogen abstraction by the alkylcarbonyloxy

radical (61). The higher temperature employed facilitates the alkyl-oxygen homolysis process. The requirement of higher temperatures, however, precludes application of this method to sensitive substrates.

In 1975, Barton and McCombie 90 reported that thiocarbonyl esters of secondary alcohols ($\underline{66}$) underwent homolytic cleavage upon treatment with tri- \underline{n} -butylstannane to give the deoxygenation product.

The proposed mechanism, illustrated in Scheme XIII, involved attack of the tri-n-butylstannyl radical on sulphur of $(\underline{66})$ to form a stabilized radical intermediate The radical could react by two pathways: Homolysis of the alkyl-oxygen bond would form the thermodynamically stable carbonyl group and is designated as pathway A. Hydrogen capture by the newly formed alkyl radical (68)would lead to the deoxygenated product (69). The driving force for the reaction was postulated to arise from the greater stability of the carbonyl over the thiocarbonyl function; hydrogen capture would form (70) via pathway Subsequent hydrolytic processing of (70) would lead to the starting alcohol (71). The isolation of benzyl ether products (72) from treatment of thionobenzoate esters with $tri-\underline{n}$ -butylstannane presumably arises from the dethiation of (70) followed by a second hydrogen capture.92

The energy difference between the two mechanistic

SCHEME XIII

pathways A and B is apparently quite small. Hence, the proportion of deoxygenation product is dependent on the relative stabilities of intermediate (67) and the alkyl radical (68) generated by its homolysis. Primary alcohol derivatives reacted exclusively via pathway B due to the relative instability of a primary alkyl radical. Barton found that pathway A was favoured over pathway B for secondary alcohol thionoesters at higher temperatures. High dilution, whereby immediate hydrogen capture by intermediate (67) was less likely, also favoured pathway A.

Other thiocarbonyl derivatives examined by Barton 90 showed similar reactivity towards tri-n-butylstannane and also lead to deoxygenation products. Ring substituted thionobenzoates showed little advantage over the parent derivative. However, the methyldithiocarbonate (methyl-xanthate) group (73) eliminated the possibility of benzyl ether formation. Deoxygenation yields reported for all these derivatives were from 70-85% using steroid and simple carbohydrates as model compounds. Thioimidazolide derivatives (74) also were deoxygenated. They were

$$R-O-C$$
 $R-O-C$
 73
 $R-O-C$
 74
 N

synthesized easily under relatively mild conditions. Barton noted, however, that the thioimidazolide reductions were less consistent and were dependent on the substrate used. He attributed this behavior to the formation of imidazole in the reaction. The decomposition of tri-n-butylstannane to hexa- \underline{n} -butyldistannane is a process catalyzed by the presence of secondary amines. 93

Applications of Barton's methods to more demanding synthetic problems have met with varied success. In the total synthesis of hirsutene, Tatsuta and co-workers 94 prepared the dithiocarbonate derivative of a hirsutene precursor and reported a 90% conversion to the deoxygenated product upon treatment with tri-n-butylstannane. Rasmussen 95 recently reported that an 87% yield of a 4-deoxypyranoside (75) was obtained via reduction of the thioimidazolide derivative (74).

SCHEME XIV

$$R = O - C$$

Im

R'= C(CH) 2

Acton et al. 92 found that the 2-thionobenzoate derivative of 5-0-benzoyl-3-deoxy-3-C-[(benzoyloxy)methyl]-1-0-methyl- β -D-allofuranose (76) gave predominant formation of the corresponding benzyl ether (77) upon reduction. Brimacombe et al. 96 noted similarly poor results using this method. Unpublished results from our laboratories also have confirmed that benzyl ether formation is a consistent problem in this reaction. Acton's group reported that reduction of the methyldithiocarbonate derivative of the dibenzylated analogue of (78) gave a 2:1 ratio of deoxygenated product (79) to starting alcohol (80).

SCHEME XV

76 R = COPh, X = Ph:

78

$$R = CH2Ph, X = SMe: \frac{79}{80}$$

In addition to the dual reaction pathways, there are limitations of Barton's method arising from the preparation of the thionocarbonate derivatives. The strongly basic conditions used for the introduction of thiobenzoyl and methyldithiocarbonate groups are incompatible with several useful blocking groups employed in carbohydrate and nucleoside chemistry. An alternative synthesis of thionobenzoates was reported by Barton. 90 However, it involves the inconvenient generation and use of imidoyl chlorides and hydrogen sulfide. 90 As well, these thionobenzoate derivatives are the most prone to give high ratios of by-products relative to deoxygenation.

Barton 97 also examined the reduction of cyclic thionocarbonates $(\underline{81})$ with tri- \underline{n} -butylstannane. It was found that homolytic opening of cyclic thionocarbonate formed from one primary and one secondary hydroxyl group gave the alkane arising from secondary radical formation exclusively. The more stable radical and resulting alkane was generated. This method was applied to the 2',3'-cyclic thionocarbonate of $6-\underline{N},5'-\underline{O}$ -diacetyladenosine $(\underline{82})$, obtained in 46% yield by the procedure of Goodman and co-workers. 98 Treatment of a solution of $(\underline{82})$ in dimethylacetamide with tri- \underline{n} -butylstannane at 175° C was followed by basic hydrolysis and acetylation. The product mixture was found to contain 60% of the 2'-deoxy $(\underline{83})$ and 29% of the 3'-deoxy $(\underline{84})$ derivatives. Isolation and removal of the acetyl groups gave 2'-deoxyadenosine

 $(\underline{10})$ in ~22% overall yield from adenosine $(\underline{6})$. This yield is comparable to other methods of 2'-deoxyadenosine synthesis discussed in Chapter I.A. $^{69,71-73,75}$ Differing ratios of 2'- and 3'-deoxynucleosides would be expected upon application of this route to other substrates. The promise of this approach is limited by the low overall yield obtained and required separation of isomers.

Thus, a general method of 2'-deoxygenation was lacking although significant progress toward solving the problem had been made primarily by the work of Barton and his co-workers.

A very serious obstacle encountered in the quest for a general 2'-deoxygenation procedure was the lack of a method for the selective 3',5' protection of the trihydroxy nucleosides. A variety of methods exist by which primary alcohols may be derivatized in the presence of secondary hydroxyl groups. 99 Hence the 5'-hydroxyl group can be protected selectively with no difficulty. The problem lay in differentiating between the 2'- and 3'-hydroxyl functions. Since both secondary groups are very similar in terms of reactivity and steric environment, methods which involve partial rather than total derivatization of the hydroxyl groups give mixtures of . 2',5' and 3',5' isomers. 100-102 Although resolution of these mixtures is usually possible by chromatographic means, the overall yield of 3',5' blocked nucleoside is normally well-below 50%. Monoester derivatives of this cisglycol system undergo rapid migration under slightly basic conditions. 102-104 This process, illustrated in Scheme XVII for the case of the acetyl function, involves a cyclic orthoester intermediate (85) and gives rise to an equilibrium mixture of products. Reese and Sulston 105 reported that $5'-\underline{0}$ -acetyl-2',3'- $\underline{0}$ -methoxyethylideneadenosine (86) could be opened under mildly acidic conditions to give a mixture of 2',5' (87) and 3',5' $di-\underline{0}$ -acetyladenosine (88). The 3',5' isomer (88) could

SCHEME XVII

be crystallized from solution, which shifted the equilibrium to favour this isomer. Good yields of crystalline (88) were realized. However, re-equilibration proceeds upon dissolution in solvents required for derivatization of the 2'-hydroxyl group.

Moffatt and co-workers 106 reported high yields of

 $2'-\underline{0}$ -tosyladenosine ($\underline{90}$) from adenosine ($\underline{6}$) via <u>in situ</u> generation of an activated stannyl acetal (ketal) intermediate ($\underline{89}$). Ranganathan prepared $3',5'-di-\underline{0}$ -tetrahydropyranyl- $2'-\underline{0}$ -tosyladenosine ($\underline{91}$) from $2'-\underline{0}$ -tosyladenosine ($\underline{90}$). Treatment of ($\underline{91}$) with sodium amalgam gave $3',5'-di-\underline{0}$ -tetrahydropyranyladenosine ($\underline{51}$) in 45% yield from adenosine.

SCHEME XVIII

Nucleoside cyclic 3',5'-monophosphates (92) have been used occasionally as selectively blocked substrates. 107 However, the acidically ionized phosphate function is not amenable to use in most organic solvents. Preliminary investigations into the utilization of cyclic phosphate triesters (93) and cyclic phosphonates (94) did not appear to be promising. 108 The possibility of developing a selective carbocyclic protecting group was pursued briefly in our laboratory.

In 1978, Markiewicz and Wiewiorowski 109 reported a reagent for the selective preparation of 3',5' protected nucleosides. They had observed that triisopropylsilyl chloride reacted with primary alcohols 10³ times faster than with secondary alcohols. When 1,3-dichloro-1,1,3,3-tetraisopropyldisiloxane (95) was allowed to react with uridine in the presence of imidazole, 3',5'-0-(1,1,3,3-tetraisopropyldisilox-1,3-diyl)uridine (96) was obtained in 70-80% yield. The analogous adenosine derivative (97) also was produced in comparable yield.

The disiloxyl group was removed smoothly using a soluble fluoride ion salt, but was relatively stable to mild acidic and basic conditions.

SCHEME XIX

HO OH
$$+$$
 Si O Si OH

HO OH $+$ Si OH

 0 OH

 0 OH

 0 Si OH

 0 OH

This realization of an efficient and facile means to selectively protect nucleosides at 0-3' and 0-5' was essential to the development of a general 2'-deoxy-genation procedure.

These findings of Markiewicz have been substantiated in our laboratory and a representative variety of 3',5'-tetraisopropyldisiloxyl nucleosides have been prepared in high to quantitative conversion yields.

The ¹H NMR spectra of all of the 3',5' blocked nucleoside derivatives were very similar. The anomeric proton coupling constant was small in all cases. This

indicated that a relatively fixed sugar conformation existed in these compounds. It previously had been determined by ¹H NMR analysis in our laboratories that all of the nucleoside 3',5'-cyclic monophosphate derivatives exist in consistently rigid conformations that could be utilized to distinguish between α and β anomers the precursor nucleosides. ¹¹⁰ Difficulties involved in preparing those derivatives, however, have severely limited practical application of this unequivocal method of anomeric determination.

Similarities and consistency of the ¹H NMR spectra of the nucleoside 3',5'-cyclic monophosphate and 3',5'-tetraisopropyldisiloxyl compounds suggested that the latter easily prepared derivatives might be useful for this anomeric configuration determination. A high field ¹H NMR and single crystal X-ray structure determination was pursued in order to evaluate this possibility.

RESULTS AND DISCUSSION

A. A General Procedure for the Conversion of Ribonucleosides to 2'-Deoxynucleosides.

In 1975, Barton and McCombie 90 reported that secondary alcohols could be reductively deoxygenated through the homolytic cleavage of thionoester derivatives. This method was discussed at length in Chapter I.B. We examined the applicability of this novel approach to the problem of 2'-deoxygenation in purine nucleosides. The model compound chosen for our investigation was 3'-0-methyladenosine (98). A facile methylation procedure using diazomethane and stannous chloride had been developed in these laboratories which gave mixtures of 2'-and 3'-monomethylated nucleosides. 111 Use of the 3'-0-methyl compound eliminated the problem of selective blocking of the 2',3' diol system. The $3'-\underline{0}$ -methy⁴ group was expected to exert minimal steric and electronic interference at the 2' hydroxyl function and thus would allow clear evaluation of the deoxygenation method. The primary 5'-hydroxyl function was blocked selectively with the tert-butyldiphenylsilyl group 112 to give the 3' and 5' protected nucleoside (99). This silyl protecting group can be removed easily with fluoride anion 113 but is stable to mild acid and base conditions. more its nonpolar and bulky substituents serve to increase

the lypophilicity of the nucleoside.

S C H E M E XX

When (99) was treated with thiobenzoylimidazolide under the strongly basic conditions of the Barton procedure, 114 two products were obtained. The major component was found to be the 6-N-thiobenzoylated derivative (100) and the minor compound the 6-N,2'-O-dithiobenzoylated nucleoside (101). Hence the 6-amino group on the base was protected prior to thiobenzoylation. Treatment of (99) with dimethylformamide dimethylacetal in dimethylformamide 115 gave the 6-N-dimethylaminomethylene derivative (102). This N-6 blocking group is

difficult to remove selectively since conversion to the $6-\underline{\mathtt{N}}$ -formyl product occurs on exposure to acid. Subsequent basic conditions are required to regenerate the free 6-amino function. However, few methods exist for selective protection of a relatively unreactive amino group with a base stable protecting group in the presence of a free hydroxyl function. When (102) was subjected to the previously noted thiobenzoylimidazolide procedure, the $2'-\underline{0}$ -thionobenzoyl nucleoside (103) was obtained in 64% yield.

S C H E M E XXI

Slow addition of a dilute solution of (103) in toluene to a refluxing solution of $tri-\underline{n}$ -butylstannane in toluene resulted in the formation of two major products. These were identified as the 2'-0-benzyl ether (104) and the 2'-deoxy derivative (105). The viability of this deoxygenation process was verified by isolation of 2'-deoxy-3'-0-methyladenosine (106) after treatment of (105) with tetra-n-butylammonium fluoride · (TBAF) followed by acidic workup and anion exchange chromatography on Dowex 1 \times 2 (OH $^-$). 50 However, it should be noted that (104) was a major product of the reaction. Barton 90 reported that the yield of deoxygenated product increased as the concentration of the reacting species decreased. In our reductions of nucleoside derivatives, benzyl ether formation always occurred to a large extent irrespective of changes in addition procedures, concentrations and temperatures.

Methyldithiocarbonate (methylxanthate) derivatives were also reported to give deoxygenated products upon treatment with $tri-\underline{n}$ -butylstannane. When $(\underline{99})$ was dissolved in THF and treated with two equivalents of sodium hydride, excess carbon disulphide and methyl iodide according to the Barton procedure, multiple product formation was observed by TLC. Similarly, reaction of $(\underline{102})$ under these conditions resulted in a mixture of compounds.

Although no single compound was isolated for characterization, purine methylation at N-1 and Dimroth rearrangement to the $6-\underline{N}$ -methyl products are known to occur under similar conditions. If N-1 in view of the complex mixtures obtained by this procedure, this approach was abandoned.

Thus Barton's methods of deoxygenation, while conveniently successful with steroids and simple carbohydrate derivatives, were not directly applicable to nucleosides. The forcing basic conditions used to generate the thiocarbonyl derivatives were not selective and required fully protected substrates. Although the 3'-O-methyl ether and 6-N-dimethylaminomethylene groups could be used in model studies, the difficulties associated with removal of these functions precluded their use as protecting groups. Furthermore, the low yield of 2'-deoxygenated product obtained from the reduction of the thionobenzoate was unsatisfactory.

The problem of 2'-deoxygenation of nucleosides was separated into two interdependent studies. The first concern was to find an alcohol derivative which would undergo efficient alkyl-oxygen homolysis and could be synthesized under mild conditions. The second was to establish a procedure for the appropriate selective 3' and 5' protection of nucleosides. The protecting group(s) must be able to withstand the conditions required for

the above derivatization and homolysis at C-2' but must be subject to removal under relatively mild conditions after the reduction step. Since purine 2'-deoxynucleosides are very susceptible to acid hydrolysis, blocking groups that require acidic conditions for removal were excluded.

A reagent for activation and removal of the 2'-oxygen atom was desired that would functionalize alcohols under normal acylation conditions. Originally we examined the possibility of using thiobenzoyl chloride: Dithiobenzoic acid was prepared according to the procedure of Houben 117 through the reaction of phenylmagnesium bromide with carbon disulphide. The magenta colored ethereal solution obtained was very unstable toward air. Reaction of this product with thionyl chloride gave a complex mixture which did not yield the desired thiobenzoyl chloride. Dithiobenzoic acid was stabilized through salt formation (Zn, Pb). However, regeneration of the free acid was required prior to reaction with thionyl chloride. The extreme sensitivity of dithiobenzoic acid and thiobenzoyl chloride to oxygen and/or water was a serious limitation and hence this reagent was abandoned.

A chlorodithiocarbonate derivative appeared to offer more potential as an acylating reagent owing to increased stability and ease of preparation. Phenyl

chlorodithiocarbonate 118 (107) and n-butyl chlorodithiocarbonate 119 (108) were prepared by treatment of thiophosgene with the appropriate thiol. Both were stable distilled liquids. Cholesterol ($\underline{109}$) was chosen as a simple model secondary alcohol. When dried cholesterol was dissolved in pyridine and treated with (107) or (108)at room temperature the solution gradually became coloured and acylation was very slow. The addition of 4-N, Ndimethylaminopyridine (DMAP), a base demonstrated to be a powerful acylation catalyst, 120 made no observed difference. It was necessary to perform the reaction at 45°C for a shorter time to achieve optimum product formation. The 3-phenylxanthate (110) and 3-n-butylxanthate (111) derivatives of cholesterol were obtained in ~80% yields through multiple addition of the appropriate reagent.

109 R - H

110 R - CS₂Ph

111 R - CS₂nBu

112

Treatment of (110) or (111) with tri-n-butylstannane in refluxing toluene, according to the procedure of Barton 90 gave cholestene ($\underline{112}$) in 75-80% yield. A minor but significant amount (10-15%) of cholesterol ($\underline{109}$) was obtained also. No reaction was observed when these reductions were attempted in refluxing benzene. ever, addition of the free radical initiator 2,2'-azobis-(2-methylpropanitrile) (more commonly referred to as azobisisobutyronitrile (AIBN)) to the benzene solution resulted in cholestene formation at 80°C without concomitant Mormation of cholesterol. Furthermore, it was found that product formation was unaffected by the concentrations of substrate and $tri-\underline{n}$ -butylstannane under these conditions. It had been noted by Barton that without initiator present the yields of reduced products were inversely proportional to the concentrations of reactants. 90

Phenylselenylthiocarbonyl chloride (113) and phenyl chlorothionocarbonate (114) also were examined as acylating reagents. Treatment of diphenyldiselenide with sodium borohydride in ethanol 121 followed by introduction of thiophosgene gave (113) in low yield. The orange-red liquid was distilled prior to use. This reagent (113) solidified on contact with pyridine. Treatment of cholesterol with (113) in pyridine gave no observed formation of product. Variation of reaction temperatures and solvents had little effect and this approach was abandoned.

The procedure of Miyazaki 122 was employed for the

preparation of $(\underline{114})$ from phenol and thiophosgene. *

The resulting yellow liquid was stable and was purified in good yield by vacuum distillation.

$$C1-C \lesssim S$$

$$SePh$$

$$C1-C \lesssim OPh$$

$$\frac{113}{114}$$

When a solution of cholesterol in pyridine was treated with ($\underline{114}$) at room temperature moderate conversion to the phenylthionocarbonate derivative ($\underline{115}$) was observed. The reaction solution became coloured more slowly than the analogous reactions with the chlorodithiocarbonate reagents ($\underline{107}$) and ($\underline{108}$). Acylation was possible at room temperature as opposed to 45°C. A 96%

yield of (115) was obtained when the reaction was performed at room temperature in dichloromethane using 3-4 equivalents of pyridine. Thus, in terms of acylating potential, (114) was superior to other derivatives examined. Treatment of (115) with tri-n-butylstannane in refluxing toluene without an initiator resulted in almost no reaction occurring. Repeating this treatment of (115) at 80°C in the presence of AIBN resulted in a product distribution indistinguishable from that of the xanthates (110) and (111) under identical conditions. In all three cases cholestene was the major product. However, a steroidal by-product was formed also. The mass spectrum of this by-product showed a molecular ion of m/z 401 which is indicative of the presence of a nitrogen atom. The base peak at m/z 368 (arising from the formation of $\Delta^{3,5}$ cholestadiene) was common to all the cholesterol derivatives with a labile group at C-3. We speculated that this by-product could result from reaction of the steroidal radical formed and AIBN. the reductions of (110), (111) or (115) were performed in refluxing toluene using di-tert-butylperoxide as the initiator, cholestene was the sole product. A near quantitative conversion to cholestene was indicated by TLC and an 84% isolated yield was obtained after chromatography on alumina followed by crystallization.

We evaluated reactions of (110), (115) and 3β -

<u>109</u> X - OH

112 X - H

110 X - 0CS₂Ph

115 X - OCS(OPh)

116 X - OCSPh

117 X - OCH₂Ph

118

cholesterylthionobenzoate 90 (116) with tri-n-butylstannane under different conditions in an attempt to arrive at a plausible explanation for the differences observed. As discussed in Chapter I.B., Barton found that product to by-product ratios in the reduction of thionobenzoates were very sensitive to reactant concentrations. In our studies on (116) we also found this to be true. Treatment of (116) with tri-n-butylstannane in refluxing benzene with or without added AIBN gave the 3benzyl ether derivative (117) and cholesterol (109) as the respective major and minor products. reaction was conducted in refluxing toluene, equal amounts of (117) and cholestene (112) were obtained as well as a minor amount of (109). Slow addition of the reactants to toluene at 110° C gave (112) as the major product. However, minor quantities of (117) and (109) were always formed. The postulated intermediate benzylic radical $(\underline{67})$ would also be stabilized by the α -sulphur substituent. Owing to this stabilization, the lifetime of (67) might permit bimolecular hydrogen transfer from the stannane to compete with the desired unimolecular carbon-oxygen homolysis (8 scission). This prediction is consistent with the finding that at 80°C the benzyl ether (117) which results from hydrogen transfer is the major product observed. Barton's observations that high dilution and higher temperatures

favoured deoxygenation 90 also are consistent with the intermediacy of $(\underline{67})$.

Reaction of the phenyldithiocarbonate (110) with tri-n-butylstannane in the absence of a free radical initiator proceeds at 110°C but not at 80°C. Thus it would appear that the activation energy for initiation of reaction of these derivatives is higher than that of the thionobenzoates. The $\alpha\text{-dithio}$ radical intermediate (118) would be presumed to be less stabilized than the benzylic thio radical $(\underline{67})$. The formation of by-product cholesterol in the uninitiated reaction of dithiocarbonate (110) with stannane at 110°C was assumed to arise from hydrogen transfer to (118) in an \sim analogous manner to that occurring with (67). However, the observations that the reaction is independent of concentration and proceeds at a lower temperature in the presence of AIBN mitigate against this pathway as the sole route for formation of cholesterol. It might be more plausible to suggest that formation of the alcohol proceeds independently from the pathway for the alkane. These processes might not share (or else might diverge in early pathways from) the same initiation. intermediate. The involvement of an intermediate such as (118) was not substantiated. A more detailed study of all the products formed in the reaction would be required for a morp defined mechanistic analysis. Since

the mechanistic implications of cholesterol formation were of interest to us for practical considerations only, a more detailed study for theoretical evaluation was not pursued. It is noteworthy that heating a solution of the phenyldithiocarbonate (110) and AIBN in toluene at 110°C for extended periods gave no evidence of reaction in the absence of tri-n-butylstannane. Therefore, interaction between the redistilled tin hydride and the thiocarbonyl compound is apparently necessary for the overall thionocarbonate ester cleavage, and alcohol regeneration is not simply a deacylation artifact.

p

Reaction of the phenylthionocarbonate (115) with tri-n-butylstannane in the presence of AIBN proceeded at 80°C and gave product formation indistinguishable from that of the phenyldithiocarbonate (110) under the same conditions. However, essentially no reaction of (115) and tri-n-butylstannane occurred at 110°C in the absence of initiator. Reaction did occur at 140°C in the absence of AIBN to give cholesterol as the major product and cholestene as a minor product. Subjection of (115) to refluxing xylene in either the presence or absence of AIBN gave neither cholesterol nor cholestene although some decomposition of the starting material was noted upon prolonged heating. It would applear that the activation energy for initiation of reaction of (115) is higher than that of (110), but that the subsequent

deoxygenation process is similar. It is possible that the difference in reactivity results from the greater stability of an α -dithio radical intermediate (118) compared to an α -thio- α -oxy radical intermediate (119).

A procedure employing phenylthionocarbonate derivatives was chosen for our deoxygenation of alcohols
for several reasons. - Phenyl chlorothionocarbonate (114)
is a stable, effective acylating reagent that does not
require strongly basic conditions to derivatize hydroxyl
groups. The phenylthionocarbonate function is stable
to aqueous work-up procedures and is amenable to chromatographic purification methods. Reaction of these derivatives with tri-n-butylstannane occurred under mild
conditions in the presence of radical initiator and
led exclusively to the desired alkane products.

SCHEME XXII

120 R - H

121 R = CS(OPh)

122

The deoxygenation procedure was applied to 5α , 6α -epoxy-3 β -cholesterol (120) following the procedure outlined for cholesterol. The intermediate phenylthio-nocarbonate (121) was obtained in 94% yield and reduction with tri-n-butylstannane gave 5α , 6α -epoxy-cholestane (122) in 78% overall yield from (120).

The 5'- $\underline{0}$ -(tert-butyldiphenylsilyl)-3'- $\underline{0}$ -methyladenosine derivative (99) again was chosen as a suitable model to test the applicability of this deoxygenation method to nucleosides. Treatment of a solution of (99) in pyridine with phenyl chlorothionocarbonate ($\underline{114}$) caused the reaction solution to slowly become coloured and gave a 60% yield of the 2'- $\underline{0}$ -phenylthionocarbonate derivative ($\underline{123}$). The yield was not optimized, but no 6- \underline{N} -acylated products were observed. Treatment of a solution of ($\underline{123}$) in toluene with tri- \underline{n} -butylstannane at 75°C in the presence of AIBN resulted in almost quantitative conversion to $\underline{\#5}$ '- $\underline{0}$ -(tert-butyldiphenylsilyl)-2'-deoxy-3'- $\underline{0}$ -methyladenosine ($\underline{124}$) by TLC. Thus, an effective means to remove the 2'-hydroxyl group from

S C H E M E XXIII

nucleosides was at hand. The remaining major obstacle was the selective protection of nucleosides at 0-3' and 0-5' to allow specific derivatization and removal of the 2'-hydroxyl function.

Recommunication by Markiewicz and Niewiorowski 109a reported the first selective and stable 3',5' protecting group for nucleosides. We prepared this 1,3-dichloro-1,1,3,3-tetraisopropyldisiloxane (95) (TPDSC1) by a modified procedure improved from that reported by Markiewicz. Treatment of a solution of adenosine (6) in pyridine with TPDSC1 (95) gave 3',5'-0-(1,1,3,3-tetraisopropyldisilox-1,3-diyl)adenosine (97) (3',5'-0-tetraisopropyldisilox-1,3-diyl)adenosine (97) (3',5'-0-

SCHEME XXIV

TPDSadenosine) in 80% yield. When the 3',5'-0-TPDS derivative (97) was dissolved in pyridine and treated with phenyl chlorothionocarbonate (114) at room temperature, the reaction solution slowly became coloured. However, only a trace of the 2'-0-phenoxythiocarbonyl nucleoside (125) was formed. Multiple additions of reagent (114) to the pyridine solution and prolonged reaction times gave ~20% yields of the 2'-thionocarbonate derivative (125). It appeared that pyridine catalyzed decomposition of the reagent (114). The rate of acylation of sterically accessible alcohols such as cholesterol by reagent (114) in the presence of pyridine was sufficiently rapid to compete with the decomposition process. However, this was not the case with the hindered 3',5'-0-TPDS nucleosides. A variety of solvents and catalytic bases were tested for use in the acylation reaction of 3',5'-0-TPDSadenosine (97) with phenyl chlorothionocarbonate (114). Treatment of a suspension of 3',5'-0-TPDSadenosine (97) in anhydrous acetonitriles with 2 equivalents of 4-N, N-dimethylaminopyridine (DMAP) and 1.1 equivalents of reagent (114) resulted in quantitative conversion to the 2'-Q-phenoxythiocarbonyl nucleoside (125) as observed by TLC. The 1 H NMR spectrum of the isolated 2'-thionocarbonate (125) (Table 1) showed a marked downfield shift for the 2' H resonance in relation to that of the 2'-hydroxyl derivative (97).

1

This is consistent with reported 1 H NMR chemical shifts of C-H groups with ester and thionoester substituents. 123 It is noteworthy that we were unable to synthesize the analogous 2'-0-phenyldithiocarbonate nucleoside derivative (126) under any conditions tried using the phenyl chlorodithiocarbonate reagent (110).

Treatment of a solution of the $2'-\underline{0}$ -phenylthionocarbonate derivative (125) in toluene with tri-n-butylstannane at 75°C-in the presence of AIRN gave quantitative conversion to 2'-deoxy-3',5'-0-TPDSadenosine (127). Removal of the TPDS group from (127) was effected by subsequent treatment of the toluene reduction solution with tetra- \underline{n} -butylammonium fluoride (TBAF) at 80°C. The 2'-deoxyadenosine (10) obtained was purified by anion exchange chromatography on Dowex 1 \times 2 (OH $^{-}$). 50 All steps of the reaction sequence $(\underline{6})+(\underline{97})+(\underline{125})+(\underline{127})+$ (10) proceeded efficiently without crystallization or chromatographic purification of the preceding compound. The individual reactions were performed on the crude residues obtained from evaporation of the dried organic layer after liquid-liquid partition work-up of the preceding reaction. The four step reaction sequence from adenosine (6) gave 2'-deox x adenosine (10)in 78% overall yield. This is markedly superior to any prior synthetic method for the 2'-deoxygenation of a purine nucleoside. 30,69,71-73,75,76

When uridine (9) was treated with TPDSC1 (95) in pyridine, $3',5'-\underline{0}$ -TPDSuridine ($\underline{96}$) was obtained in excellent yield. Reaction of (96) with phenyl chlorothionocarbonate (114) in acetonitrile in the presence of DMAP proceeded quantitatively to give the $2'-\underline{0}$ phenoxythiocarbonyl-3',5'-0-TPDS derivative (128). It had been reported previously that the 0-2+2'cyclonucleoside of uridine (36) was formed in high yield upon heating either the cyclic 2',3'-0-carbonate 60-62 or thionocarbonate derivatives. 63,64 However, treatment of the 2'-thionocarbonate derivative (128) in toluene at 80°C in the presence of $\text{tri-}\underline{n}\text{-butylstannane}$ and AIBN resulted exclusively in 2'-deoxygenation to give the 2'-deoxy-3',5'- $\underline{0}$ -TPDS derivative ($\underline{129}$). No cyclonucleoside product was observed in this reaction. Treatment of the protected 2'-deoxynucleoside (129) with

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<u>96</u> R - OH

128 R - OCS (OPh)

129 R - H

TBAF was followed by chromatography on AU-4 charcoal to give 2' deoxyuridine (38) in 68% overall yield from uridine (9). This yield is slightly superior to prior procedures involving 0-2+2'-cyclonucleoside interconversions. 67

Application of our feduction method to guanosine (7) required only minor alterations. The insolubility of guanosine in pyridine restricted its conversion to 3',5'-0-TPDSguanosine (130) upon treatment with TPDSC1 $(\underline{95})$. When guanosine was dissolved in DMF/pyridine (10:1) and treated with TPDSC1 (95) a good conversion to $3';5'-\underline{0}$ -TPDSguanosine ($\underline{130}$) occurred. Analysis of the reaction mixture by TLC indicated that formation of a minor nucleoside by-product had occurred that migrated slightly faster than the 3',5'-0-TPDS derivative (130) in all solvent systems examined. product was not investigated due to the inherent difficulty associated with purification of guanosine derivatives. To avoid possible complications the $3'.5'-\underline{0}$ -TDPS derivative ($\underline{130}$) was isolated and crystallized in 70% yield prior to reaction with phenyl chlorothionocarbonate (114). Treatment of a suspension of (130) in acetonitrile with reagent (114) and DMAP gave $2'-\underline{0}$ -phenoxythiocarbonyl-3',5'- $\underline{0}$ -TPDSguanosine (131) in 94% yield. Reduction of the 2'-thionocarbonate (131) with tri-n-butylstannane following the procedure

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$$HO \longrightarrow B$$

$$HO \longrightarrow IPr_2SI \longrightarrow R$$

$$IPr_2$$

outlined previously and subsequent treatment of the reduction solution with TBAF gave 2'-deoxyguanosine $(\underline{11})$. The crude 2'-deoxyguanosine was applied to a column of Dowex 1 X 2 (OH⁻) resin and eluted using tetraethylammonium bicarbonate buffer. The overall yield of 2'-deoxyguanosine $(\underline{11})$ from guanosine $(\underline{7})$ was 60% and the yield of $(\underline{11})$ from 3',5'- $\underline{0}$ -TPDSguanosine $(\underline{130})$ was 85%.

The reaction conditions used to prepare 2'-deoxy-adenosine ($\underline{10}$) from adenosine ($\underline{6}$) were applied to tubercidin without alteration. The overall yield of 2'-deoxytubercidin ($\underline{55}$) from tubercidin ($\underline{54}$) was 68%. Differences between yields obtained for 2'-deoxy-adenosine ($\underline{10}$) and 2'-deoxytubercidin ($\underline{55}$) arise primarily from the different solubilities of the two products. The recovery of 2'-deoxytubercidin ($\underline{55}$) from crystallization was lower than that obtained for 2'-deoxyadenosine ($\underline{10}$).

Treatment of toyocamycin (vengicide) ($\underline{135}$) with TPDSC1 ($\underline{95}$) in pyridine gave an 89% yield of 3',5'- $\underline{0}$ -TPDStoyocamycin ($\underline{136}$). The excellent crystallization properties of this derivative ($\underline{136}$) allow the high recovery. Reaction of 3',5'- $\underline{0}$ -TPDStoyocamycin with phenyl chlorothionocarbonate ($\underline{114}$) in acetonitrile with DMAP followed by reduction of the 2'-phenylthionocarbonate derivative ($\underline{137}$) with tri- \underline{n} -butylstannane in toluene

gave an 87% yield of the blocked 2'-deoxy derivative (138) from toyocamycin (135). The 2'-deoxy-3',5'-0-TPDS-toyocamycin (138) was treated with TBAF and purified by chromatography on AU-4 charcoal to give a 69% overall yield of 2'-deoxytoyocamycin (139) from toyocamycin (135). Passage of 2'-deoxytoyocamycin (139) through a column of Dowex 1 X 2 (OH⁻) resin, using 40% methanol/water for elution gave 2'-deoxysangivamycin (141) in quantitative yield. This procedure for the conversion of toyocamycin (135) to sangivamycin (140) is more convenient and efficient than the method using hydrogen peroxide in aqueous base reported by Tolman et al. 124

Treatment of cytidine $(\underline{8})$ with TPDSC1 $(\underline{95})$ in pyridine gave 3',5'-0-TPDScytidine $(\underline{142})$ in excellent yield. However, reaction of $(\underline{142})$ with reagent $(\underline{114})$ under the usual conditions gave only partial conversion to the desired 2'-0-phenoxythiocarbonyl-3',5'-0-TPDS-cytidine $(\underline{146})$. Moderate overall conversion of $(\underline{8})$ to this derivative $(\underline{146})$ was obtained using six equivalents of DMAP in place of the normal two equivalents. It was assumed that the difference in behavior of cytidine $(\underline{8})$ compared to the other nucleosides, studied was attributable to the reactive 4-amino group on the base. The selective \underline{N} -4 acetylation procedure of Fox and coworkers $\underline{125}$ was followed to give $\underline{4}$ - \underline{N} -acetylcytidine $(\underline{143})$. Treatment of $(\underline{143})$ with TPDSC1 $(\underline{95})$ in pyridine

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gave $4-\underline{N}$ -ace yl-3',5'- \underline{O} -TPDScytidine ($\underline{144}$) in excellent yield. Surprisingly, this derivative ($\underline{144}$) was totally unreactive toward phenyl chlorothionocarbonate ($\underline{114}$) despite numerous attempts to promote reaction through variation of the reaction conditions. Treatment of $4-\underline{N}$ -acetyl-3',5'- \underline{O} -TPDScytidine ($\underline{144}$) in acetonitrile with acetic anhydride in the presence of DMAP gave $4-\underline{N}$ -acetyl-2'- \underline{O} -acetyl-3',5'- \underline{O} -TPDScytidine ($\underline{145}$) in good yield. The cause(s) of the lack of reactivity of the $4-\underline{N}$ -acetyl derivative ($\underline{144}$) toward reagent ($\underline{114}$) remain(s) unclear.

TPDScytidine (146) to the standard reduction procedure resulted in a complex mixture of products. Although a minor product of the mixture was found to be the 2'-deoxy-3',5'-0-PDScytidine derivative, the major contituent was unreacted thionocarbonate (146). Extended reaction times with multiple additions of tri-n-butyl-stannane resulted in further decomposition of reagents. The catalytic decomposition of tri-n-butylstannane by amines was discussed in Chapter I.A. Hence the 2'-thionocarbonate derivative (146) was acetylated with acetic anhydride and pyridine to give 4-N-acetyl-2'-0-phenoxythiocarbonyl-3',5'-0-TPDScytidine (147). This acetylation could be accomplished directly by subsequent addition of excess acetic anhydride to the acetonitrile

Treatment of $(\underline{147})$ with $tri-\underline{n}$ -butylstannane and AIBN at 80° C in toluene resulted in good conversion to the 2'-deoxygenated compound $(\underline{148})$. The $3',5'-\underline{0}$ -TPDS group was removed from the 2'-deoxy derivative $(\underline{148})$ in the usual manner and treatment of the $4-\underline{N}$ -acetyl derivative with sodium methoxide in methanol followed by anion exchange chromatography on 2-deoxycytidine 2-deoxycytidine 2-deoxycytidine 2-deoxycytidine 2-deoxycytidine 2-deoxycytidine 2-deoxycytidine 2-deoxycytidine 3-deoxycytidine 3-deoxycytidine

Few methods exist which allow successful deoxygenation of hindered alcohols on acid sensitive carbohydrate derivatives. Treatment of methyl $\beta - \underline{D}$ -ribofuranoside $\sim (149)$ in pyridine with TPDSC1 (95) gave an excellent , yield of methyl 3',5'- $\underline{0}$ -TPDS- β - \underline{D} -ribofuranoside (150). Acidic work-up conditions had to be stringently avoided since this glycoside is easily hydrolyzed. Chromatography could be performed efficiently on neutral silica if contact time with the absorbant was kept to a minimum. The $2-\underline{0}$ -phenoxythiocarbonyl-3,5- $\underline{0}$ -FPDS derivative (151) was prepared in the usual manner. The standard reduction conditions gave clean conversion to methyl 2 $deoxy-3,5-\underline{0}-TPDS-\beta-\underline{0}-erythro-pentofuranoside.$ Isolation and purification of this blocked derivative and the deprotected methyl 2-deoxy- β - \underline{D} - \underline{e} rythro-pentofuranoside

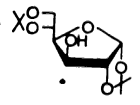
SCHEME XXVIII

were not pursued extensively owing to the acid sensitivity of these compounds. Protection of hydroxyl groups of 2'-deoxynucleosides as esters stabilizes the glycosyl bond against acid hydrolysis. 126 The crude residue of the methyl 2-deoxyfuranoside obtained from evaporation of the reduction and deprotection solution was treated with anhydrous pyridine. Treatment of this mixture with excess p-toluyl chloride gave the known methyl 2-deoxy-3,5-di-0-p-toluyl- β -p-erythro-pentofuranoside 127 (152), which was purified on neutral silica without serious

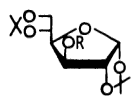
decomposition. The overall yield obtained for the conversion of methyl $\beta-\underline{D}$ -ribofuranoside (149) to methyl 2-deoxy-3,5-di- $\underline{0}$ -p-toluyl- $\beta-\underline{D}$ -erythro-pentofuranoside (152) was 58%. This method should find synthetic applicability in the carbohydrate field for deoxygenation of hindered and acid sensitive sugars.

Finally, our deoxygenation procedure was utilized for removal of a 3-hydroxyl function of a simple protected sugar derivative. Treatment of $1,2;5,6-di-\underline{0}$ -isopropylideneglucofuranose (153) in acetonitrile in the presence of DMAP with phenyl chlorothionocarbonate (114) gave the 3- $\underline{0}$ -phenoxythiocarbonyl derivative (154) in excellent yield. Treatment of this derivative (154) with tri- \underline{n} -butylstannane and AIBN in toluene at 80°C gave the 3-deoxysugar (155) in 85% overall yield from the starting alcohol (153). This result is comparable to results obtained by Barton 90 for deoxygenation of the same sugar model.

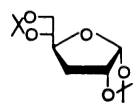
S C H E M E XXIX



153



154 R = CS(OPh)



155

The deuterolysis of 9-(2-chloro-2-deoxy-β-D-arabinofuranosyl)adenine (156) using tri-n-butyltin deuteride was studied previously in our laboratory. 128

The ratio of 2'-deoxy-2'-deutero-ribo (157a) to 2'-deoxy-2'-deutero-arabino nucleoside epimers (157b) produced in this free radical mediated dehalogenation reaction was -85:15. The observed bias in product formation was presumed to arise from the favoured attack on the bulky deuterostannane by the less hindered ribo face of the 2' radical (which is trans to the heterocycle at (1). Equivalent product ratios were betained from the epimeric 2'-chloro-arabino (156) and 2'-chloro-ribo adenosine derivatives (158). Hence, the free radical intermediate was assumed to be separated from the departing chlorine atom.

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We were interested to determine whether the present deoxygenation process with the 2'-phenylthionocarbonate derivatives would result in a similar stereochemical outcome. Treatment of a pyridine solution of $9-\beta-\underline{D}$ arabinofuranosyladenine (159) with TPDSC1 (95) gave a single product (by TLC). It was not apparent by 1H NMR * spectroscopy whether the product was the 8-membered trans-fused 3',5'- $\underline{0}$ -TPDS derivative ($\underline{160}$) or the 10membered cis 2',5'-0-TPDS derivative (161). A sample of this product was acetylated for Wurther 1H NMR analysis. The marked downfield shift of the signal for the 2'hydrogen allowed the unambiguous assignment of the 2'-O-acetyl structure (162).. This confirmed that 9-(3,5-0-TPDS- $\beta-\underline{D}$ -arabinofuranosyl) adenine (160) was the product of the silylation reaction. The 2'-phenylthionocarbonate derivative (163) was prepared by the standard method. The H NMR downfield shift of the peak for H-2' (see Table 5) reconfirmed our assignment of the 3',5'-0-TPDS structure (160).

Treatment of a toluene solution of the thionocarbonate ($\underline{163}$) with tri- \underline{n} -butyltin deuteride and AIBN
at 80°C was followed by addition of TBAF. Anion exchange chromatography on Dowex 1 X 2 (\underline{OH}^-) gave the
2'-deoxy-2'-deuteroadenosines ($\underline{157a}$, $\underline{157b}$). The ratio of $\underline{r1bo}$ to $\underline{arabino}$ deutero C-2' epimers was ~88:12 as
evaluated by integration of the peaks for the H-2' and

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H-2" protons measured at 400 MHz.

When the identical conditions of deuterolysis and deblocking were applied to the <u>ribo</u> 2'-thionocarbonate derivative (125) the ratio of <u>ribo</u> to <u>arabino</u> deutero C-2' epimers obtained was identical to that obtained from the <u>arabino</u> 2'-thionocarbonate derivative (163). This would imply that a separated alkyl radical intermediate is formed in this reaction.

As a control study, the 2'-chloro-2'-deoxy-3',5'-0-TPDS-arabino derivative (164) was prepared in the usual manner from 9-(2-chloro-2-deoxy-β-D-arabinofuranosyl)adenine (156). Reaction of the blocked 2'-chloro derivative (164) with tri-n-butyltin deuteride at 80°C in the presence of AIBN followed by treatment with TBAF and anion exchange chromatography gave the 2'-deoxy-2'deuteroadenosines (157a, 157b). The ratio of ribo to arabino deutero C-2' epimers again was ~88:12 as was obtained for both the epimeric ribo (125) and arabino $(\underline{163})$ 2'-thionocarbonate derivatives. It seems likely that all three derivatives, react via a common 2'-للهو-'yl radical intermediate in this hydrogenolysis process. The results obtained are comparable to those of our previous study. 128 The greater uniformity of product ratios obtained in the present work probably arises from the greater sensitivity of the 400 MHz 1H NMR analysis.

As discussed in our original study, the biosynthetic transformation of ribenucleotides to 2'-deoxyribo-nucleotides is thought to proceed by a free radical mediated enzymatic process. 128 The enzymatic conversions proceed with complete stereoselectivity and retention off configuration of attachment of the incoming H-2" to the ribo face. The present 2'-deoxygenation procedure allows a biomimetic conversion with greater than 85% stereoselectivity for the "natural" ribo epimer.

B. <u>A Facile Method for Determination of Anomeric</u> <u>Configuration</u>

A convenient and reliable method for determination of the anomeric configuration of an unknown natural product nucleoside or the product of a sugar base coupling synthesis has been elusive. There have been a number of empirical correlations made on various derivatives of anomeric nucleoside pairs. Small differences in spectral characteristics or other physical constants have been noted.

Originally, Hudson's rules of isorotation 129 for carbohydrate derivatives were applied to nucleosides. Ulbricht and co-workers 130 later found that the ORD/CD spectra of nucleoside anomers exhibited oppositely signed long-wavelength (" \bullet_{2u} ") transitions. This transition was found to be positive for α -purine and β -pyrimidine ribonucleosides and negative for the complementary anomers. However, both of these chiroptical methods were found to be subject to exceptions. 131

The formation of purine 12 3-N+C-5'- and pyrimidine 132 2-0+C-5'-cyclonucleosides has provided unequivocal proof of the cis orientation of the base and 6-5' (β -D-configuration). However, negative results from this chemical method could be misleading since certain β anomers do not form cyclonucleosides easily. 133 Furthermore, this method is not applicable to nucleosides with

modified bases which lack the required functionality.

The first identification of α -adenosine was accomplished via a novel method by Wright, Tener and Khorana. Synthetic α -adenosine and natural adenosine (β) were treated with periodic acid and the resulting dialdehydes were reduced with sodium borohydride. The two triols obtained had chirality remaining at C-1' only and were found to exhibit enantiomeric optical rotations. An extensive survey of oxidized-reduced nucleosides accomplished recently in these laboratories 135 revealed that triols resulting from α andmers have negative optical rotations while those from β anomers have positive rotations. However, exceptions have been observed with azapyrimidine (triazene) nucleosides.

 ^{1}H NMR spectroscopy has been utilized extensively for anomeric configuration determination. A number of empirical methods involving chemical shift differences have been published. $^{136},^{137}$ In general these deal with the anisotropic effects of the base on a specific site or function on the furanose residue. The most widely applied criterion involves chemical shift differences between the gem-dimethyl singlets of 2',3'-O-isopropylidene derivatives. 137 ^{1}H NMR spin coupling parameters have also been used to assign a trans H-1' to H-2' configuration when $J_{1'-2'} \leq 1\text{-}1.5$ Hz. 138

Other recent methods involve more advanced in-

for routine analysis. Relaxation time correlations for carbohydrate derivatives have been studied by Hall and co-workers ¹³⁹ and Guschlbauer and co-workers. ¹⁴⁰ A direct relation between the proximity of adjacent hydrogen atoms and their relative relaxation times was shown to exist. This allows calculation of the preferred conformation and anomeric configuration. ¹⁴⁰

¹³C NMR spectroscopy has also been investigated. The ¹³C NMR chemical shifts of carbon atoms are very sensitive to steric crowding. Pairs of anomers studied ¹⁴¹ had C-1' resonance frequencies at lower fields when the base was cis to the 2'-hydroxyl group relative to the trans orientation.

A survey of a number of methods with a critical evaluation of their limitations has been made. 110

Since essentially all of the methods have been found to have exceptions or severe limitations and the rules formulated were based on empirical observations, unequivocal identification of a single anomer often was impossible. Furthermore, most methods require measurement of a relative difference between anomeric derivatives and both anomers must be available for comparison.

Robins and MacCoss 110 described the first generally applicable method which allows the unequivocal deter-

mination of a single anomer. This method is based on the conversion of the ribonucleoside to its 3',5'cyclic monophosphate derivative and determination of its H-1' to H-2' NMR coupling constant. They found that the peak for H-1' appears as a singlet $(J_{1'-2'} \lesssim$ 0.7 Hz) for β-ribonucleoside 3',5'-cyclic monophosphates $(\beta-3',5'-cNMP's)$ and a moderate to strongly coupled doublet $(J_{1'-2'} \ge 3.5 \text{ Hz})$ for cis (a) anomers. The consistent ranges of the observed coupling constants result from the conformationally rigid cyclic nucleotides which are comprised of a tran's-fused six to five membered ring system. Since the geometry of the furanose moiety is fixed, $\phi_{H-1'-H-2'}$ is virtually the same for all β -3',5'-cNMP's and anisotropic effects of the base and other variable factors (previously utilized for anomeric determination) are of negligible consequence. Thus no exceptions have been observed with this method. Furthermore, since each anomer is uniquely defined, there is no need for both amomers to be available. This method, however, has not found widespread use due to the difficulty in the synthesis of the 3',5'-cNMP's.

We observed that the ¹H NMR spectra of the readily available 3'.5'-0-(1,1,3,3-tetraisopropyldisilox-1,3-diyl) (3',5'-0-TPDS) nucleoside derivatives were very similar to those of the 3',5'-cNMP's. It was concluded that the furanose ring in both types of derivatives must

be restricted to a similar conformation range. A representative number of 3',5'-O-TPDS nucleosides were prepared to determine if the conformational pattern observed was consistent and as reliable as that of the 3',5'-cNMP's.

Table I contains a compilation of the 1H NMR chemical shift data for the 3',5'-0-TPDS nucleosides with a trans 1'-2' configuration. The $3',5'-\underline{0}$ -TPDS derivative of methyl $\beta - \underline{D}$ -ribofuranoside is included also. The coupling constants for these derivatives are listed in Table 3. The coupling constants found for the 1'-2' and 3'-4' protons indicate that the furanose ring is in the ${}^3T_4 - {}^3E - {}^3T_2$ conformational range which is similar to that found in the 3',5'-cNMP's. 142 been estimated that $\varphi_{H-1\,'-H-2\,'}$ must be in the range of $90\text{--}105^\circ$ for a coupling of less than 1 Hz to occur in these furanosyl structures. A small anomeric coupling was observed for the B-ribofuranosides throughout the series. However, the anomeric proton coupling of the \sim 3',5'- $\underline{0}$ -TPDS derivatives of $\overline{9}$ - α - \underline{D} -arabinofuranosyladenine $(\underline{186})$ and $6-N-acetyl-9-(2-\underline{0}-acetyl-\alpha-\underline{D}-arabinofuranosyl$ adenine (191) are not consistent with the values obtained for the β (trans 1'-2')nucleosides. This indicates that a different furanose conformation has been adopted in the cases of $(\underline{186})$ and $(\underline{191})$.

The 3', 5'-cNMP's show much less variation in coupling parameters than the 3', 5'-0-TPDS derivatives

as expected from the trans-fused 6 to 5 vs. 8 to 5 membered ring systems. The anomeric proton coupling constants of the $3',5'-\underline{0}$ -TPDS derivatives showed only a slight variation with temperature, but solvent polarity had a more pronounced effect (see Table 5). We observed that the $J_{1,-2}$, coupling decreased for the 3',5'- $\underline{0}$ -TPDS- β - \underline{D} -ribofuranosides and increased for the 3',5'- $\underline{0}$ -TPDS- α - \underline{D} -ribofuranoside with increasing solvent polarity. This is consistent with a solvation-dependent conformational bias resulting from the steric demands of the bulky isopropyl groups on the TPDS fragment. It also emphasizes that the furanose ring is not geometrically locked as in the 3',5'-cNMP molecules. The altered furanose conformation observed for (186) and (191) also supports this conclusion. The $J_{1'-2'}$ values of the $2'-\underline{0}$ -phenoxythiocarbonyl-3',5'- $\underline{0}$ -TPDS derivatives in Table 3 show some variation in relation to their 2' hydroxyl precursors. Thus steric and/or electronic demands can alter the conformation of the furanose moiety in these compounds. In contrast, derivatization of the 2' hydroxyl group of the 3',5'cNMP's had a negligible effect on $J_{1'-2'}$.

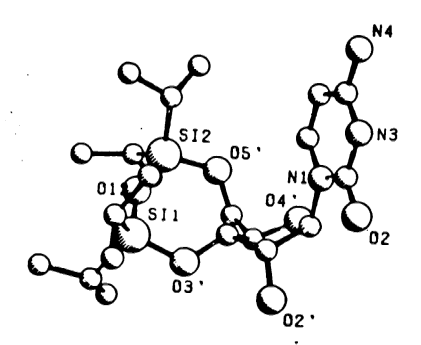
The chemical shift data for a number of cis 1'-2' examples of $3',5'-\underline{0}$ -TPDS nucleosides are compiled in Table 2 with the respective coupling constants listed in Table 4. The anomeric coupling constants of the α ribonucleosides and β arabinonucleoside de-

rivatives are very similar to those found in the 3'.5'-cNMP's. 110 A rationalization for the observed differences in the two types of <u>cis</u> 1'-2' nucleosides as well as a discussion of the strained conformations of the α -ribonucleoside cyclic monophosphates has been presented by Robins and MacCoss. 110

Although the 3',5'-0-TPDS derivatives have a less rigidly defined furanose conformation than the analogous 3',5'-cNMP's, a clear difference between anomers is observed. Ribofuranoside derivatives with $J_{1'-2'} \leq 1.5$ Hz can safely be assigned a β configuration since cis coupling values of this small a magnitude are geometrically precluded. It appears that cis anomers have $J_{1'-2'} \ge 3.5$ Hz. The β derivatives with an intermediate anomeric coupling value $(J_{1,-2}, \sim 2.5 \text{ Hz})$ should be subjected to other methods for a definitive identification. The ease of preparation of these 3',5'-0-TPDS derivatives makes this method very convenient as well as normally definitive for evaluation of the anomeric configuration of an unknown B-ribofuranoside. Furthermore, the 3',5'-0-TPDS group can be removed easily and efficiently with fluoride anion to regenerate the parent nucleoside.

In order to verify our 1 H NMR conclusions concerning the furanose conformation adapted by the 3',5'- $\underline{0}$ -TPDS derivatives and to examine the trioxadisila ring fragment we subjected the cytidine compound ($\underline{142}$) to

single crystal x-ray analysis. The computer generated structure is presented below. A Dreiding molecular



model of this compound had indicated that the C-5'-0-5' bond might assume the trans-gauche (tg) orientation as found for the 3',5'-cNMP's. 142 The 4'-5' and 4'-5" coupling constants did not contradict this assumption (see Table 3). However, a gg orientation for C-5'-0-5' also was compatible with the data and, of course, no information was available as to the conformation of the 3',5'-0-TPDS ring fragment. The x-ray analysis showed that the C-5'-0-5' bond was oriented gg for (141) in the solid state. The Si-0-Si bond

angle was expanded to 158° which allows formation of an unstrained 8-membered ring. This silicon-oxygensilicon bond expansion and accommodation of the four bulky isopropyl groups on the silicon atoms apparently result in a highly preferred conformation of the furanose in this trans-fused 8 to 5 membered ring system. The almost perfect ^3E envelope conformation observed for $(\frac{141}{4})$ has $\phi_{H-1'-H-2'}$ of ~90°C as anticipated from the NMR data. Table 6 illustrates the approximate relationship between estimated bond angles of adjacent hydrogen atoms in $(\underline{141})$ and the observed ^1H NMR spin-spin coupling values.

It is interesting to note that the methyl groups of the isopropyl substituents on silicon were nonequivalent by ¹³C NMR. The x-ray structure data revealed slightly different C-C bond lengths for the geminal methyls on each of the isopropyl methine carbons. A detailed analysis of the x-ray structure of (141) will be published in collaboration with James and Sawyer. ¹⁴³

В

ОН <u>159</u>

<u>160</u>

OAc

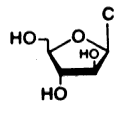
<u>162</u>

C1 <u>156</u>

<u>164</u>

N₃ 187

<u> 381</u>



<u>189</u>

190

TABLE 1

O-TPDS-8-D-tibonucleosides 1,111)
3',5'-
E H
fο
Data
Shift
Chemical
NMN

ane r cob
.05 3.92 5.60 7.30
m-7.52 ^m p-7.37 ^m .08 ^e 3.92 ^e 5.61 ^b 6.50 ^a
4.07m 3.98m - 6.43a 7.17
4.05 ^e 3.92 ^e 5.52 ^b 7.06 ^a -
4.06 - 5.35a 7.14B 7.42B 7.42B 7.42B
•
4.01 ^m 4.01 ^m - 7.02 ^a 7.17 ^m 7.51 ^m
4.10e 3.95e 5.70a 9.67a

Punod bonud	88 88 88 88 88 88 88 88 88 88 88 88 88	C-1,	C-2'	C-3	C-4.	, S-2	C-5"	но, .	NH ₂	Phenyl H'	8 Deuterated
168	7.82	5.86 ^b	4.55e	4.80e	4,13 ^m	4.15m	4.08 ^m	4.60m	6.32ª	1	Acetone
31	5.53 ^b 7.69 ^b	'5.54ª	4.14m	4.16 ^e	3.97 ^f	4.12e	3.92e	5.58b	; '		DMSO
128	5.85 ^b 7.80 ^b	6.00	6.08 ^b	4.60 ^e	4.16	4.20m	4.10m			7.19m 7.52m 7.37m	cDC13
142	5.68b 7.72b	5.57ª	3.92°	4.09°	4.01 ^f	4.17	3.92	5.61 ^b	7.:16 ^b	,	DMSO
144	7.23b 8.16b	5.62	₽80.4	4.08	4.08™	4.24e	3.94e	5.83 ^b	1.)	DWSO
146	5.87b 7.71b	5.79 ^b	6.08	4.84	4.03	7.1.4	3.94 ■	•	7.34ª	7.19 ^m . 7.52 ^m	DMSO
147	7.16	6.09ª	, 6.10 ^b	4.52e	4.20	¼.32™	4.06e	•	• • •	7.37 m	cDC13 *
172	5.55b 5.82e 5.82e	5.75ª	4.10 ^B	₹.16 ^m	4.05	4.00 m	4.00 m	5.75 E.8	•	E 7 9/. 2	DMSO

'-<u>O</u>-TPDS-β-<u>D</u>-ribonucleosides NMR Chemical Shift Data for 34 TABLE 1 (continued)

Com-	Base H	C-1,	C-2	C-3	* * - 0	· 2 - 2		H 0	NH ₂	Phenyl	. E	C-2' C-3' C-4' C-5' C-5" OH NH ₂ Phenyl H's Deuterated Solvent
174	5.50 ⁸ 5.60 ^b 7.52 ^b	5.85	4.16 ^m	4.16 ^m 4.25 ^e 4.16 ^m 4:18 ^m 3.95 ^e 4.20 ^m 7.20 ^a	4.16 ^m	4:18m	3.95e	4.20m	7.20			CDC13
170	8.32	4.52ª	3.96 b	3.96 ^b 4.13 ^e 3.84 ^m 4.04 ^m 3.87 ^m 5.20 ^b	3.84	4.04 1	3.874	5.20b	* 1	, ,	¢	DMSO
	1 .	5.047	4.52	4.52 4.33 4.00 4.16 4.10 5.12 4.90 5.20 6.85	# · · · ·	4.16 ^m	4.10 ^m	5.12	4.90ª 6.85ª		•	Acetone
176	8.25	5.25 ^b	4.68	4.68° 5.28° 4.05° 4.05° 4.05° 4.05° 4.05° 6.90°	4.05m	4.05m	4.05m	4.05m	6.90ª	. •		Acetone
150		4.79 ^b		3.97° 4.08° 3.84° 3.89° 3.76° 4.10°	3.84	3.89ª	3.76 ^m	4.10 ^m		1		DMSO

e = doublet of doublets f = doublet of triplets g = doublet of doublet of doublets m = multiplet iid) In all cases above, the resonances of the iPr H's on the disiloxyl group 1) Chemical shifts in δ ppm downfield from Me $_4$ Si(internal) d = quartet Appeared as a complex multiplet at \$ 1.05-1.10. a = singlet b = doublet c = triplet

TABLE 2

NMR Chemical Shift Data for $\alpha + D - ribo$, $\beta - D - a rabino, and <math>\alpha - D - a r$

Com- pound	Base H's	C-1,	C-2'	€-3	. 7-3	C-5 1	C-5"	но	NH 2	Deut.
180	8.16 ^a 8.26 ^a	6.38 ^b	4.34m	4.50°	4,20m	4.02e	3.91	5.79	7.30	DMSO
182	5.57b	6.02 ^b	4.18m	4.36e	4.13 ^f	4.03 ^e	3.87	5.40b	7.07 ^b	DMSO
184	5.67b 7.47b	6.01b	4.13 ^e	4.33	4.07 ^f	4.00°	.3.86°	5.62b	ı	DMSO
160	8 0 0 8 0 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	4.79 ^b 6.21 ^b	3.97e	4.08 ^c	3.84 ^m 3.80 ^m	3.89m 4.12e	3.76m 3.92°	4,07 ^m 5.80 ^b	7.30	DMSO
162	. 80 8. 60 9. 80 8. 80 8. 80 8. 80 8. 80 8. 80 8. 80 8. 80 8. 80 8. 80 8. 80 8. 80 8. 80 8. 80 8. 80 8. 80 8. 80 8. 80	6.56 ^b	5.64m	5.0811.	4.05m	4.05m	4.05m	ı		DMSO
164	8 14 8	6.50 ^b	4.66°	,61.4	3.94	4.23 ^e	4.10e	Y	5.95ª	CDC1,
188	8.10a 8.11a	6.40 ^b	m06.4	m06.4	3.94	4.24e	3.94m	· 1	7.36ª	DMSO
190	5.66b 7.44b	6.07 ^b	4.22d	, 4.07 ^e	3.68	3.98	3.92	5,63 ^b	7.11 ^b	DMSO
99	7.98 ^a 8.26 ^a	5.81 ^b	4.78	4.54	4.16 ^f	4.10e	4.04	r	5.91	CDC13
191	8.648	6.20b	6.26 ^m	4.61 ^m	4.61 ^{III}	3.98	3.98ª	•	3 I	DMSO

TABLE 3

(

97 - 0.9 5.0 8.0 3.2 2.6 12.6 4.7 135 - 1.6 5.0 8.0 3.2 2.6 13.0 - 130 - 1.6 5.0 8.0 3.2 2.6 13.0 5.2 131 - 2.6 5.8 7.7 2.0 2.3 12.5 4.7 132 3.7 1.5 8.6 3.2 2.7 13.0 - 133 3.7 1.5 8.6 3.2 2.7 13.0 5.0 136 - 1.3 5.5 8.4 2.8 2.2 11.8 - 168 - 1.5 5.0 7.5 3.5 2.0 11.5 - 168 - 1.5 5.0 7.5 3.5 2.0 11.5 - 168 - 1.5 5.0 7.5 3.5 2.0 11.5 - 168 - 1.5 5.0 7.5 2.6 2.6 13.2 4.5 168 - 1.5 5.3 10.0 m m - 168 - 1.5 5.3 10.0 13.2 <th>Com-</th> <th>J5-6 J1'-2' J2'-3' J3'-4' J4'-5' J4'-5" J5'-5"</th> <th>J1'-2'</th> <th>J21-31</th> <th>J31-41</th> <th>J4'-5'</th> <th>34'-5"</th> <th>J\$'-5"</th> <th>J2'-2'0H</th>	Com-	J5-6 J1'-2' J2'-3' J3'-4' J4'-5' J4'-5" J5'-5"	J1'-2'	J21-31	J31-41	J4'-5'	34'-5"	J\$'-5"	J2'-2'0H
- 0.9 5.0 8.0 3.2 2.6 12.6 - 1.2 5.5 8.7 4.1 m 13.0 - 2.6 5.8 7.0 m m m m m 3.6 a11) 5.5 7.7 2.0 2.3 12.5 3.7 1.5 5.2 8.6 3.2 2.7 13.0 - 1.3 5.5 8.5 m m m m - 1.5 5.0 8.4 2.8 3.2 17.8 - 1.5 5.0 7.5 3.5 2.0 11.5 8.0 s 4.7 8.4 2.6 2.6 13.2 8.0 s 5.3 10.0 m m m 7.6 s 5.8 2.2 2.7 13.0									
- 1.2 5.5 8.7 4.1 m 13.0 - 1.6 5.0 8.0 3.2 2.6 13.0 3.6 a11) 5.5 7.7 2.0 2.3 12.5 3.7 1.5 5.2 8.6 3.2 2.7 13.0 - 1.3 5.5 8.5 m m m m m m m m m m m m m m m m m m m	26	•	6.0	5.0	8.0	3.2	2.6	12.6	4.7
3.6 8.6 3.2 2.6 13.0 3.6 a ¹¹) 5.8 7.7 2.0 2.3 12.5 3.7 1.5 5.2 8.6 3.2 2.7 13.0 - 1.5 5.2 8.6 2.6 2.4 13.0 - 1.3 5.5 8.5 m m m - 1.5 5.2 8.4 2.8 3.2 12.8 - 1.5 5.2 8.4 2.8 3.2 11.8 8.0 e 4.7 8.4 2.6 2.6 11.5 8.0 e 5.3 10.0 m m m 7.6 e 5.3 2.2 2.7 13.0 7.6 e 5.3 2.2 2.7 13.0	125	, t	1.2			4.1	E	13.0	•
3.6 a ¹¹⁾ 3.6 a ¹¹⁾ 5.5 7.7 2.0 2.3 12.5 3.7 1.5 5.2 8.6 3.2 2.7 13.0 - a 5.0 8.5 m m m m m - 1.3 5.5 8.5 m m m - 1.5 5.2 8.4 2.8 3.2 12.8 - 1.5 5.0 7.5 3.5 2.0 11.5 8.0 a 4.7 8.4 2.6 2.6 13.2 8.0 a 5.3 10.0 m m 7.6 a 4.4 5.8 2.2 2.7 13.0	130	·	1.6	5.0	8.0	3.2	2.6	13.0	5.2
3.6	131	j	2.6	5.80	0	E	e	e .	ı
3.7 1.5 5.2 8.6 3.2 2.7 13.0 - 8.5 2.6 2.4 13.0 - 1.3 5.5 8.5 m m m - 1.5 5.2 8.4 2.8 3.2 12.8 - 1.5 5.0 7.5 3.5 2.0 11.5 8.0 s 4.7 8.4 2.6 2.6 13.2 8.0 s 5.3 10.0 m m m 7.6 s 4.4 5.8 2.2 2.7 13.0	132	3.6	(H.	5.3	7.7	2.0	2.3	12.5	4.7
- 8 5.0° 8.5 2.6 2.4 13.0 - 1.3 5.5 8.4 2.8 3.2 12.8 - 1.5 5.0 7.5 3.5 2.0 11.5 8.0 8 4.7 8.4 2.6 2.6 13.2 8.0 8 5.3 10.0 m m m m 7.6 8 4.4 5.8 2.2 2.7 13.0	133	3.7	1.5	5.2	, 60	3.2	2.7	13.0	
- 1.3 5.5 8.5 m m m m m m m m m m m m m m m m m m m	36		•	5.0	8.5	2.6	2.4	13.0	5.0
- 1.5 5.0 7.5 3.5 2.0 11.5 8.0 s 4.7 8.4 2.6 2.6 13.2 8.0 , s 5.3 10.0 m m m 7.6 s 4.4 5.8 2.2 2.7 13.0	37	1 .	1.3	5.5	8.5	E		e	•
8.0 s 4.7 8.4 2.6 2.6 13.2 8.0 m m m m m m m m m m m m m m m m m m m	99	1	1.5	.5.2	. 4.8	2.8	. 3.2	12.8	
8.0 s 4.7 8.4 2.6 2.6 13.2 8.0 , s 5.3 10.0 m m m 7.6 s 4.4 5.8 2.2 2.7 13.0	89	ı	1.5	5.0	7.5	3.5	2.0	11.5	ı
8.0 , s 5.3 10.0 m m m m 7.6 s 4.4 5.8 2.2 2.7 13.0	96	8.0	1. 1.	4.7	8.4	2.6	2.6	13:2	4.5
7.6 8 4.4 5.8 2.2 2.7 13.0	28	8.0	•	5.3	10.0	E	£	Ø	1
	42	7.6		4.4	5.8	2.2	2.7	13.0	. 4.4.

the lower limit of resolution for instrumentation used

TABLE 3 (continued).

NNR Spin-Spin Coupling Values for 3',5'-0-TPDS- β - \underline{D} -ribonucleosides

					ì	*	,	
Com-	J 5-6	J1'-2'	32:-3	3,-4,	14'-5'	J41-5"	J51-5"	J2'-2'0H
146	,	60	5.5	8.5	•	£	6	•
147		60	2.7	9.5	, E	, 6	13.8	
172	7.5	•	Œ	a	Ø	8	e	
(13-5	(J ₃₋₅ =2.5)						• *	
174	7.0	80	10.5	8.5	1.8	2.5	14.0	a
170	1	·.	4.5	8.5	E	£	12,5	0.4
178	,	2.3	5.6	.0	8. 7	3.0	12.5	•
176		2.0	5, 2	£	, e	È	, 8	E
150) 		4.5	7.5	3.0	5.9	11.5	4.2
			, 					-
1) Coupling in H:	ling in	, N	, , ,			ł	.	

TABLE 4

NMR Spin-Spin Coupling Values for α -D-ribo, β -D-arabino, and α -D-arabino 3' Q-TPDS-nucleosides¹

180 - 4.0 182 7.6 3.5 184 8.0 3.6 193 - 4.0 160 - 6.3	5.0 4.0 6.5	8.0 9.1 5.0 8.0		3.0.	12.5 %	F 3.
82 7.6 3.5 84 8.0 3.6 93 - 4.0 60 - 6.3	6.5			2:5	12.8	4.6
84 8.0 3.6 93 - 4.0 60 - 6.3	6.5	,		2.4		
60 - 6.3	6.5	,			•	4.4
6.3	8.0			4.0	10.5	e
			4.0	2.5	12.5	5.8
/ 0 = 70	Ē.			E		
164 - 6.0	8.0	, 8	0.4	3.0	13.0	. ,
188, - 6.6	· E	E	5,5	, , , E	13.0	· ,
20 7.4 6.2	7.0	7.8	3.8	3.0	13.0	5.8.
186 - 5.7	8.1	8.7	3.0	3.0	0.11	,
91 - 5.8			:		, E	*

Solvent Dependency of H-1'-H-2' Spin-Spin Coupling Values for Selected 3',5'-0-TPDS nucleosides.

Compound	J _{1'-2'}	Deuterated Solvent
	7	
97	0.9	DMSO
	1.4	CDC13
132	s 11)	DMSO
منر	1.5	CDC13
176	2.3	DMSO
	2.0	съзсосъз
	3.2	CDC13
184	3.5	DMSO
3	2.9	CDC1 ₃
186	6.2	DMSO
•	5.7	CDC13

i) Coupling in Hz

¹¹⁾ $J_{1,-2}$, $\lesssim 0.5 \text{ Hz (see Table 3)}$.

TABLE 6

Calculated and Observed Spin-Spin Coupling Values of 3',5'-0-(1,1,3,3-Tetraisopropyldisilox-1,3-diyl)cytidine

H atoms	Calcd. Tor- sion Angle ⁱ)	J _{Calcd} ii)	^J Observed
H-1'-H-2'	89.8	0	. 0
H-2'-H-3'	41.6	4.97	4.4
н-3'-н-4'	157.7	10-09	8.8
н-4'-н-5"	44.4	4.50	2.7
H-41-H-51	74.8	0.41	2.2,
OH-2'-H-2'	36:4	5.84	4.4

- i) Torsional angles calculated based on a 1.08Å carbon-hydrogen bond length and optimized sp³ geometry for the carbon atoms involved. Angles were obtained through a least-squares calculation from X-ray data and are given in degrees.
- ii) J_{H-H} Coupling values obtained from $J_{H-H}=10.5\cos^2\theta-1.2\cos\theta$. Coupling in Hz.

EXPERIMENTAL

A. General Procedures

Melting points were determined on a Reichert microstage apparatus and are uncorrected. Nuclear magnetic resonance (nmr) spectra were recorded on a Bruker WH-200 or Bruker WH-400 spectrometers operating in the FT mode, with Me_4Si as internal reference normally in $Me_2S0-\underline{d}_6$ unless specified otherwise. Ultraviolet (uv) spectra were recorded on a Cary 15 spectrophotometer and infrared (ir) spectra on a Nicolet 7199 FT(IR) instrument. Optical rotations were determined using a Perkin-Elmer Model 141 polarimeter with a 10 cm 1 ml microcell. Mass spectra (M.S.) were determined by the mass spectrometry laboratory of this department on an AEI MS-50 instrument with computer processing at 70 eV using a direct probe for sample introduction. Elemental analyses were determined by the Microanalytical Laboratory of this department or by Schwarzkopf Microanalytical Laboratory, wwwwoodside, N.Y. Evaporations were effected using Buchler rotating evaporators equipped with Dewar "dry-ice" condensers under water aspirator or mechanical oil pump vacuum at .40°C or cooler. Thin layer chromatography (TLC) was performed on E. Merck chromatographic sheets (silica gel 60 F_{254} , layer thickness 0.2 mm, catalogue 5775) with sample observation under UV light (2537 Å). Preparative

TLC was performed on glass plates coated with Merck silica gel PF 254. The solvents used for TLC were different ratios of methanol-chloroform (1:50, 1:20, 1:10) and the upper phase of EtOAc-nPrOH-H₂O (4:1:2). Silica gel column chromatography was performed using Mallin-ckrodt CC-7 (200 mesh) silica gel. Anion exchange chromatography was carried out on Dowex 1 X 2 resin in the hydroxide form.

The carbon used for chromatography was Barnebey-Cheney AU-4 charcoal. It was conditioned by the following treatments. It first was washed with methanol and then chloroform. After drying it was refluxed with 1 N aqueous HCl (with the acid solutions being replaced periodically) until the supernatant solution remained colourless, with water to neutrality, and then was refluxed with 10% aqueous NaOH (again with periodic replacement of the solution) until the supernatant remained colourless. The charcoal then was washed with water to neutrality, with methanol, and with chloroform. Finally it was air dried at room temperature.

All solvents used were of reagent grade and were distilled prior to use. Purification of most solvents and reagents was accomplished according to methods described in reference 144. Dimethylformamide (DMF) was purified according to the azeotropic distillation procedure described in reference 145. All dried solvents

were stored over Davison 4 Å molecular sieves purchased from the Fisher Scientific Company.

The reaction sequence developed for the 2'-deoxy-genation of nucleosides is a general four step procedure which is described in detail for adenosine. Subsequent experimental descriptions will refer to Procedure A: the silvlation step; Procedure B: the thioacylation step; Procedure C: the reduction step using n-Bu₃SnH; Procedure D: the desilvlation step. The standard work-up procedure refers to the partitioning of a reaction residue between ethyl acetate and water followed by washing the organic phase with cold 1 N HCl, H₂O, saturated NaHCO₃/H₂O solution, saturated NaCl/H₂O solution, drying (Na₂SO₄)? filtering, and evaporating the solvent. This procedure is detailed in the first reaction with adenosine.

Abbreviations used are: TPDSC1 = 1,1,3,3-Tetraiso-propyl-1,3-dichlorodisiloxane; PTC chloride = phenyl chlorothionocarbonate; DMAP = 4-N,N-Dimethylaminopyridine; and AIBN = Azobisisobutyronitrile. Skelly B represents the petroleum ether fraction which boils between 60-90°C. Upon distillation the fraction boiling at 63-65°C was recovered.

B. Syntheses

1,1,3,3-Tetraisogropy1-1,3-disifoxane (96a)

The procedure of Gilman and Clark 146 was followed resulting in a 35-40% direct yield of (95a) based of starting trichlorosilane. The by-product disopropyl-silanol obtained was converted to the disiloxane by addition of P_2O_5 and distillation of the supernatant. This gave a combined yield of 90% of (95a): bp 104° C/10 mm Hg; $n_D^{20} = 1.4335$; ir/neat/cm⁻¹ γ SiOH = 2110; M.S. m/z 246 (100%, M⁺), 203 (64%, M⁺-iPr); (lit. 109 bp 95° C/15 mm Hg).

1,1,3,3-Tetraisopropyl-1,3-dichlorodisiloxane (TPDSC1)-(95)

To 50 g (0.20 mol) of (95a) was added 500 ml of CCl₄, and 20% of the solvent was removed by distillation at atmospheric pressure. The dried solution was cooled to -20° C and stirred vigorously while dry Cl₂ was introduced for 1 h. The reaction vessel was then subjected to aspirator vacuum at 20 mm Hg for 5 min to remove dissolved HCl formed in the reaction. Introduction of Cl₂ then was continued for 1 h. The above procedure was repeated until the solution became yellow permanently. The CCl₄ was evaporated in vacuo and the residue was distilled at 68-70%C/0.05 mm Hg to give 48 g (75%) of (95): $n_D^{20} = 1.4550$; M.S. m/z 271 (M^+ -iPr); (lit. $n_D^{10} = 120^{\circ}$ C/15 mm Hg).

3β -Cholesteryl phenylthionocarbonate (115).

To a stirred solution of 3.87 g (10 mmol) of cholesterol (109) in 60 ml of dichloromethane was added 3 ml (37 mmol) of anhydrous pyridine and 2.0 ml (11 mmol) of phenyl chlorothionocarbonate (PTC chloride) (114). 122 After 2 h the solvents were evaporated in vacuo and the residue was subjected to the standard work-up procedure. Crystallization from acetone gave 5.0 g (96%) of (115): mp 162-163.5°C; M.S. m/z 369.3493 (85%, M⁺-0CSO ϕ = 369.3521), 368.3436 (100%, M⁺-HOCSO ϕ); Ahal. Calcd. for $C_{34}H_{50}O_{2}S$: C 78.11, H 9.64, S 6.12. Found: C 78.10, H 9.37, S 6.11.

Cholest-5-ene (112)

To 1.046 g (2 mmol) of ($\underline{115}$) was added 30 ml of toluene, 800 µl (3 mmol) of \underline{n} -Bu $_3$ SnH and 65 µl (0.4 mmol) of \underline{di} - \underline{tert} -butylperoxide. The solution was refluxed under nitrogen for 3 h before 3 ml of \underline{n} l \underline{M} solution of \underline{n} -Bu $_4$ NF in tetrahydrofuran was added. After refluxing for 4 h the solvents were evaporated \underline{in} vacuo and the residue was chromatographed on a column of alumina (20 g, 3 x 15 cm) using Skelly B as the eluant. Evaporation of the eluants and crystallization of the residue from ethanol gave 620 mg (84%) of ($\underline{112}$): mp 92-94°C; \underline{M} .S. \underline{M} /z 370 (100%, \underline{M}) 371 (30%, \underline{M} +1); \underline{Anal} . Calcul. for

C₂₇H₄₆: £ 87.49, H 12.51. Found: C 87.32, H 12.35; (1it. 90 mp 90-92°C).

$5\alpha, 6\alpha$ -epoxy- 3β -cholesteryl phenylthionocarbonate (121)

To 2.01 g (5 mmol) of 5α , 6α -epoxy- 3β -cholesterol 147 - (120) dissolved in 30 ml of dichloromethane was added 15 ml (19 mmol) of pyridine and 1.0 ml (5.5 mmol) of PTC chloride (114). The solution was stirred for 2 h before the solvent was removed in vacuo. After the standard work-up procedure the residue was crystallized from acetone to give 2.48 g (92%) of (121): mp 191-194°C; M.S. m/z 385.3453 (77%, M+-0tS0 ϕ), 384.3398 (100%, M+-H0CS0 ϕ); Anal. Calcd. for $C_{34}H_{50}O_{3}S$: C 75.79, H 9.35, S 5.95, O 8.91. Found: C 75.73, H 9.04, S 6.00, O 9.13.

5α,6α-Epoxycholestane (122)

To 1.078 g (2 mmol) of ($\underline{121}$) dissolved in 30 ml of toluene was added 800 µl (3 mmol) of n-Bu₃SnH and 65 µl (0.4 mmol) of di-tert-butylperoxide. The solution has refluxed under nitrogen for 3 h. The solvent was evaporated in vacuo and the residue was chromatographed on a column of alumina (20 gm, 3 x 15 cm) using Skelly B as the eluant. Evaporation of the appropriate fractions and crystallization of the residue from acetone gave 603 mg (78%) of ($\underline{122}$): mp 74-75°C; M.S. m/z 386.3552 (100%, M⁺ = 386.3549); Anal. Calcd. for C₂₇H₄₆O:

C 83.86, H 12.00, O 4.14. Found: C 83.81, H 11.86, O 4.05; (lit. 148 mp 74-75%C).

3',5'-0-(1,1,3,3-Tetraisopropyldisilox-1,3-diyl)adenosine (97)

To 267 mg (1-mmol) of dried adenosine ($\underline{6}$) suspended in 10 ml of anhydrous pyridine was added 320 μ l (1 mmol) of (TPDSC1) (95) and the mixture was stirred at room temperature for 3 h. Pyridine was evaporated in vacuo and the residue 🏂 partitioned between ethyl acetate and water. The organic phase was washed with 2 \times 20 ml of cold 1 $\underline{\text{N}}$ HC1, H_2O , saturated $\text{NaHCO}_3/\text{H}_2\text{O}$ solution, saturated ${
m NaC1/H}_2{
m O}$ solution, dried $({
m Na}_2{
m SO}_4)$, filtered, and evaporated. The resulting amorphous product was of sufficient purity for direct use in subsequent reactions. characterization, this material was chromatographed on a column of silica (10 g, 2×10 cm) using chloroform Sand 2% methanol/chloroform as eluants. Evaporation of the appropriate fractions and crystallization of the residue from CH_3CN gave 433 mg (85%) of (97): 99.5°C; uv (0.1 \underline{N} HC1) max 257 nm (ϵ 14,900); (0.1 \underline{N} NaOH) max 259 nm (ε 15,100); M.S. m/z 509.2485 (8.8%, $M^{+}[C_{22}H_{39}N_{5}O_{5}Si_{2}] = 509.2490), 466.1924 (100%, <math>M^{+}-iPr)$, 164.0573 (17%, BHCHO), 136.0621 (19%, B+2H); Anal. Calcd. for C₂₂H₃₉N₅O₅Si₂: C 51.83, H 7.71, N 13.74. C 51.65, H 7.92, N 13.73.

2'-0-Phenoxythiocarbonyl-3',5'-0-(1,1,3,3-tetraisopropyldisilox-1,3-diyl) tenosine (125)

To the vacuum-dried residue of crude (97) was added 15 ml of anhydrous acetonitrile and 250 mg (2 mmol) of 4- $\underline{N},\underline{N}$ -dimethylaminopyridine (DMAP). A 200 µl (1.1 mmol) \sim aliquot of PTC chloride (114) was added and the solution was stirred at room temperature for 16 h. Solvent was removed in vacuo and the residue was subjected to the standard work-up procedure. The resulting product was sufficiently pure to be used directly in the reduction step. Purification of this material was achieved by chromatography on a column of silica (10 g, 2 x 10 cm) using chloroform and 1.5% methanol/ chloroform as eluants. Evaporation of the appropriate fractions gave 586 mg (90%) of (125) as an oil with uv (EtOH) max 259 nm, 228 nm, min 245 nm; M.S. m/z 602.1924 $(6.1\%, M^{+}-iPr[\epsilon_{26}H_{36}N_{5}O_{6}Si_{2}S] = 602.1927), 492.2471 (12%,$ M^+ -OCSO ϕ), 491.2473 (7.1%, M^+ -HOCSO ϕ), 449.1984 (8.6%, M⁺-iPr-OCSOφ), 164.0576 (5.4%, BHCHO) 135.0545 (2.0%, B+H).

2'-Deoxy-3',5'- $\underline{0}$ -(1,1,3,3-tetraisopropyldisilox-1,3-diyl)-adenosine ($\underline{127}$)

The crude $(\underline{125})$ was dissolved in 20 ml of distilled toluene and 32 mg (0.2 mmol) of azobisisobutyronitrile

(AIBN) and 400 µl (1.5 mmol) of \underline{n} -Bu $_3$ SnH were added. The solution was decaygenated with oxygen-free nitrogen for 20 min and then heated to 75°C for 3 h. Solvent was evaporated $\underline{in\ vacuo}$ and the residue was chromatographed on a column of silica (10 g, 2 x 10 cm) using chloroform as the eluant. Evaporation of the eluant and crystallization of the residue from ethanol gave 370 mg (75%) of ($\underline{127}$): mp 113-114.5°C; uv (MeOH) max 259 nm (ε 14,000); (0.1 \underline{N} HCl) max 257 nm (ε 13,500); M.S. m/z 493.2538 (18%, M⁺ = 493.2544), 450.1986 (100%, M⁺-iPr), 164.0574 (11%, BHCHO), 162.0778 (1.3%, BHCH=CH $_2$); Anal. Calcd. for C29H39N5O4Si $_2$: C 53.51, H 7.96, N 14.18. Found: C 53F31, H 7.90, N 13.93.

2'-Deoxyadenosine (10)

Deprotection of crude (127) was effected by addition of 2 molar equivalents of <u>n</u>-Bu₄NF (as a 1 <u>M</u> solution in THF) directly to the above reduction mixture. The solution was heated for an additional hour at 75°C. Solvent was evaporated and the residue was partitioned between ether and water. The aqueous phase was applied to a column of Dowex 1 X 2 (0H⁻) resin. Elution of the product with water, evaporation and crystallization of the residue from ethanol/ether gave 195 mg (86%) of (10): mp 191-192°C; M.S. m/z 251.1073 (7.1%, M⁺ [C₁₀H₁₃N₅O₃] = 251.1082), 162.0827 (100%, BHCH=CH₂),

135.0660 (99%, B+H); (lit. 149 mp 187-189°C). This product was identical to naturally occurring 2'-deoxy-adenosine by the usual criteria.

Treatment of 1.068 g (4 mmol) of adenosine ($\underline{6}$) by the four step sequence ($\underline{6} + \underline{97} + \underline{125} + \underline{127} + \underline{10}$) without purification of intermediates gave 780 mg (3.1 mmol) of recrystallized 2'-deoxyadenosine ($\underline{10}$) in 78% overall yield.

3',5'-0-(1,1,3,3-Tetraisopropyldisilox-1,3-diyl)uridine (96)

The conditions of Procedure A were followed using 92 mg (4 mmol) of dried uridine (9), 50 ml of anhydrous pyridine and 1.3 ml (4 mmol) of TPDSC1 (95). After the standard work-up procedure a small portion of the crude residue was chromatographed on silica using chloroform and 2% methanol/chloroform as eluants. Evaporation of the appropriate fractions yielded an oil with uv (MeOH) max 262 nm (ε 9,600); (0.1 N HCl) max 262 (ε 9,600); (0.1 N NaOH) max 262 nm (ε 7,000); M.S. m/z 486.2262 (12%, M⁺[C₂₁H₃₈N₂O₇Si₂] = 486.2268), 443.1718 (100%, M⁺-iPr).

 $2'-\underline{0}$ -Phenoxythiocarbonyl-3',5'- $\underline{0}$ -(1,1,3,3-tetraisopropyldisilox-1,3-diyl)uridine ($\underline{128}$)

The conditions of Procedure B were applied to the residue of crude (96) using 60 ml of anhydrous aceto-

nitrile, 1 g (8 mmol) of DMAP, and 800 μ l (4.4 mmol) of PTC chloride (114). After the standard work-up procedure a small sample of the crude residue was chromatographed on silica using chloroform and 2% methanol/chloroform as eluants. Evaporation of the eluate gave an oil with μ l (MeOH) max 262, 232 nm, min 245 nm; M.S. μ l 469.2206 (0.5%, μ l -QCSO μ [C₂₁ μ l 37N₂O₆Si₂] = 469.2172), 426.1577 (32%, μ l -QCSO μ -iPr), 425.1554 (100%, μ l -QCSO μ -iPr-H).

2'-Déoxyuridine (38)

The crude (128) was subjected to the reduction conditions of Procedure C using 75 ml of toluene, 1.6 ml (6.0, mmol) of \underline{n} -Bu₃SnH, and 130 mg (0.8 mmol) of AIBN. After the reaction was complete, the deblocking step of Procedure D was performed directly. Solvents were evaporated and the residue was partitioned between ether and water. The aqueous phase was stirred with 10 g of carbon. To a glass column (2 x 30 cm) was added 3 g of carbon and the slurry of carbon containing the absorbed nucleos de was layered on top. The column was washed thoroughly with water before a stepwise gradient from 20% ethanol/ H_2 0 to 40% ethanol/ H_2 0 was applied. Evaporation of the appropriate fractions and crystallization of the residue from ethanol/ether gave 620 mg (68% overall from (9) of (38): mp 162-163°C; M.S. m/z 228.0742 (4.1%, $M^{+}[C_{9}\dot{H}_{12}N_{2}O_{5}] = 228.0746$), 139.0726 (2.5%, BHCH=CH₂), 112.0278 (25%, B+H); (1it. 149 mp 163°C).

 $3',5'-\underline{0}$ -(1,1,3,3-Tetraisopropyldisilox-1,3-diyl)-guanosine (130)

To 1.132 g (4 mmol) of dried guanosine ($\underline{7}$) suspended in 60 ml of anhydrous DMF was added 4 ml of anhydrous pyridine and 1.3 ml (4 mmol) of TPDSC1 ($\underline{95}$). The mixture was stirred for 5 h and then added slowly to 1 ℓ of vigorously stirred ice water. The resulting precipitate was collected by filtration and washed thoroughly with water. Crystallization from 95% ethanol gave 1.47 g (70%) of ($\underline{130}$): mp >250°C (decomp.); uv (MeOH) max 256 nm (ε 14,300), (0.1 N HCl) max 256 nm (ε 11,700), (0.1 N NaOH) max 263 nm (ε 11,400); M.S. m/z 525.2443 (24%, M^+ [C22H39N506Si2] = 525.2435), 526.2464 (9%, M^+ +1), 483.1912 (41%, M^+ +1-iPr), 482.1876 (91%, M^+ -iPr); Ana \underline{T} ? Calcd. for C22H39N506Si2: C 50.26, H 7.48, N 13.3 \underline{T} . Found: C 49.90, H 7.37, N 13.54.

 $2'-\underline{0}$ -Phenoxythiocarbonyl-3',5'- $\underline{0}$ -(1,1,3,3-tetraiso-propyldisilox-1,3-diyl) guanosine ($\underline{131}$)

The conditions of Procedure B were applied to 1.05 g (2 mmol) of (130) using 30 ml of anhydrous CH₃CN, 500 mg (4 mmol) of DMAP and 400 μ l (2.2 mmol) of PTC chloride (114). After the standard work-up procedure the residue was applied to a column of silica (20 g, 2 x 20 cm) using chloroform and 2% methanol/chloroform as eluants. Evaporation of the appropriate fractions

and crystallization of the residue from 95% EtOH gave 1.25 g (94%) of ($\underline{131}$): mp 255-258°C; uv (EtOH) max 245 nm (ε 18,800), (0.1 N HC1) max 252 nm (ε 15,100), (0.1 N NaOH) max 257 nm (ε 14,600); M.S. m/z 661.2436 (3.8%, M⁺ = 661.2414), 662.2465 (1.6, M⁺+1), 507.2347 (19%, M⁺-0CSO ϕ), 508.2375 (8.7%, M⁺-HOCSO ϕ), 464.1801 (12%, M⁺-OCSO ϕ -iPr); Anal. Calcd. for C₂₉H₄₃N₅O₇Si₂S: C 52.62, H 6.55, N 10.58, S 4.84. (Found: C 52.33, H 6.52, N 10.61, S 4.86.

2'-Deoxyguanosine (11)

A 1.00 g (1.5 mmol) sample of (131) was subjected to the reduction conditions of Procedure C using 30 ml of toluene, 600 μ l (2.25 mmol) of \underline{n} -Bu $_3$ SnH and 50 mg (0.3 mesol) of AIBN. The deblocking step of Procedure D was performed directly on the reaction solution. were evaporated and the residue was partitioned between ether and water. The aqueous phase was applied to a column of Dowex 1 X 2 (OH $^{-}$) resin (10 ml, 2 x 5 cm) and the resin was washed well with water. Elution was effected with 0.25 M tetraethylammonium bicarbonate (TEAB) buffer (pH = 9.0). After evaporation of the eluate in vacuo the residue was taken up in 10 ml of water and reevaporated. This procedure was repeated four times to remove residual TEAB. Crystallization of the residue from water gave 345 mg (85%) of (11): mp 251-252°C; M.S. m/z 249.0864 (0.76%, $M^{+}[C_{10}H_{13}N_{5}O_{4}]-18 = 249.0853),$

151.0498 (13%, B+H), 117.0553 (41%, sugar ion); (lit. 149 mp 250°C).

4-Amino-7-[3,5-0-(1,1,3,3-tetraisopropyldisilox-1,3-diyl)-8-D-ribofuranosyl]pyrrolo[2,3-d]pyrimidine (132) (3',5'-0-TPDStubercidin)

The conditions of Procedure A were followed using 266 mg (1 mmol) of dried tubercidin (54), 10 ml of anhydrous pyridine and 320 µl (1 mmol) of TPDSCl (95). After the standard work-up procedure, a small portion of the crude residue was chromatographed on a column of silica using chloroform and 2% methanol/chloroform as eluants. Evaporation of the appropriate fractions and crystallization of the residue from CH_3CN gave (132) as colourless platelets with mp 238-242°C; uv (MeOH) max 270 nm (ε 10,800) and 227 nm (ε 20,000), min 245 nm (ε 3,800); (0.1 N HC1) (max 270 nm (ϵ 10,000) and 227 nm (ϵ 21,300), min 245 nm (ϵ 3,400); (0.1 N NaOH) max 273 nm (ϵ 11,300) and 217 nm (ϵ 38,100), min 244 nm (ϵ 3,900); M.S. m/z 508.2547 (8.0%, $M^{+}[C_{23}H_{40}N_{4}O_{5}Si_{2}] = 508.2528), 509.2598 (7.4%, M^{+}+1),$ 466.2031 (55%, M+1-iPr), 465.1980 (60%, M+-iPr), 163.0588 (100%, BHCHO), 134.0589 (38%, B+H).

 $\frac{2'-\underline{0}-\text{Phenoxythiocarbonyl}-3',5'-\underline{0}-(1,1,3,3-\text{tetraiso-propyldisilox-1,3-diyl})\text{tubercidin (133)}$

The conditions of Procedure B were applied to the residue of crude (132) using 15 ml of anhydrous CH3CN,

250 mg (2 mmol) of DMAP and 200 μ l (1.1 mmol) of PTC chloride (114). After the standard work-up procedure a small sample of the crude residue was chromatographed on silica using chloroform and 2% methanol/chloroform as eluants. Evaporation of the eluate gave an oil with uv (MeOH) max 271 nm and 232 nm, min 245 nm; M.S. m/z 491.2460 (12%, M⁺-OCSO ϕ [C₂₃H₃₉N₄O₄Si₂] = 491.2562), 490.2415 (24%, M⁺-HOCSO ϕ), 448.1905 (3.1%, M⁺-OCSO ϕ -iPr), 447.1887 (6.7%, M⁺-HOCSO ϕ -iPr), 163.0620 (75%, BHCHO), 134.0593 (44%, B+H).

2'-Deoxy-3',5'-0-(1,1,3,3-tetraisopropyldisilox-1,3-diyl)tubercidin (134)

The crude (133) was reduced according to Procedure C using 200 ml of distilled toluene, 32 mg (0.2 mmol) of AIBN, and 400 µl (1.5 mmol) of n-Bu₃SnH. An aliquot of the reaction solution was removed for characterization of (134). Solvent was evaporated in vacuo and the residue was cpromatographed on silica using chloroform and 1.5% methanol/chloroform as eluants. Evaporation of the appropriate fractions gave an oil with uv (MeOH) max 271 nm and 227 nm, min 245 nm; 200 MHz nmr (CDCl₃, TMS) δ 1.05 (m, 28, iPr), 2.58 (m, 2, H-2', 2"), 3.88 (m, 1, H-4'), 4.02 (d of d, $J_{4'-5''} = 4.6$ Hz, $J_{5''-5''} = 12.3$ Hz, 1, H-5"), 4.10 (d of d, $J_{4'-5''} = 3.9$ Hz, $J_{5''-5''} = 12.3$ Hz, 1, H-5'), 4.86 ('q', J - 7.5 Hz, 1, H-3'), 5.55 (br s, 2, NH₂-4), 6.40 (d, $J_{5'-6} = 3.7$ Hz, 1, H-5), 6.56 (d of

d, $J_{1'-2'} = 4.5 \text{ Hz}$, $J_{1'-2''} = 6.5 \text{ Hz}$, 1, H-1'), 7.21 (d, $J_{5,6} = 3.7 \text{ Hz}$, 1, H-6), 8.30 (s, 1, H-2); M.S. m/z 492.2488 (13%, $M^{+}[C_{23}H_{40}N_{4}O_{4}Si_{2}] = 492.2467$), 449.1923 (100%, M^{+} -iPr), 161.0820 (1.5%, BHCH=CH), 134.0941 (51%, B+H).

4-Amino-7-(2-deoxy- β -D-erythro-pentofuranosyl)pyrrolo-[2,3-d]pyrimidine (55) (2'-Deoxytubercidin)

The balance of the above reaction solution was subjected to deblocking Procedure D using 2 ml (2 mmol) of the 1 \underline{M} \underline{n} -Bu₄NF solution. Solvents were evaporated and the residue was partitioned between ether and water. The aqueous phase was applied to a column of Dowex 1 X 2 (OH⁻) resin (10 ml, 1.5 \bigstar 10 cm). The resin was washed with water prior to elution of the product with 10% methanol/water. Evaporation of the eluant and crystallization of the residue from ethanol gave (55): 217-218.5°C; $[\alpha]_D^{24}$ -43.8 (<u>c</u> 0.34 MeOH); uv (MeOH) max 271 (ϵ 13,600) and 227 nm (ϵ 24,200), min 239 nm (ϵ 2,700); (0.1 \underline{N} HC1) max 271 nm (ϵ 12,500) and 226 nm (ϵ 25,700), min 244 nm (ϵ 3,800); (0.1 <u>N</u> NaOH) max 271 nm (ϵ 13,100), min 239 nm (ϵ 3,100); M.S. m/z 250.1070 (5.0%, M⁺[C₁₁H₁₄- N_4O_3] = 250.1066), 161.0826 (14%, BHCH=CH₂), 135.0637 (11%, B+2H), 134.0592 (100, B+H). Anal. Carcd. for C₁₁H₁₄N₄O₃: C 52.78, H 5.64, N 22.39. Found: C 52.48, H 5.62, N 22.31; (lit. 76 mp 217-218°C; $\left[\alpha\right]_{D}^{24}$ -43 (\underline{c} 0.58 EtOH)).

Treatment of 1.064 g (4 mmol) of tubercidin (54) by the four step sequence (54+132+134+135+55) without purification of intermediates gave 680 mg (2.72 mmol) of recrystallized 2'-deoxytubercidin (55) in 68% overall yield.

4-Amino-5-cyano-7-[3,5- $\underline{0}$ -(1,1,3,3-tetraisopropyldisilox-1,3-diyl)- β - \underline{D} -ribofuranosyl]pyrrolo[2,3-d]pyrimidine (136) (3',5'- $\underline{0}$ -TPDStoyocamycin)

The conditions of Procedure A were applied to 582 mg (2 mmol) of dried toyocamycin (135) using 30 ml of anhydrous pyridine and 650 ml (2 mmol) of TPDSC1 (95). After the standard work-up procedure the residue was chromatographed on a column of silica (15 g; 2×15 cm) using chioroform and 2% methanol/chloroform as eluants. Evaporation of the appropriate fractions and crystallization of the residue from $\mathrm{CH_3CN}$ gave 945 mg (89%) of mp 171-172°C; uv (MeOH) max 280 nm (ε 17,400) (136):and 232 nm (ε 11,900), min 247 nm (ε 4,700); (0.1 N HC1) max 274 nm (ϵ 13,100) and 235 nm (ϵ 17,900), min 249 nm . (ϵ 5,000); (0.1 NaOH) max 280 nm (ϵ 16,700) and 233 nm (ϵ 11,000), min 247 nm (ϵ 4,600); M.S. m/z 533.2478 (4.1%, $M^{+}[C_{24}H_{39}N_{5}O_{5}Si_{2}] = 533.2492), 534.2514 (1.7%, <math>M^{+}+1),$ 491.1960 (38%, M⁺+1-iPr), 490.1938 (100%, M⁺-iPr), 188.0566 (18%, BHCHO), 159.0507 (10%, B+H); Anal. Calcd. for $C_{24}H_{39}N_50_5Si_2$: C 54.04, H 7.37, N 13.13. Found: C 53.95, H 7.33, N 12.97.

 $2'-\underline{0}$ -Phenoxythiocaraonyl-3',5'- $\underline{0}$ -(1,1,3,3-tetraisopropyl-disilox-1,3-diyl)toyocamycin ($\underline{137}$)

The conditions of Procedure B were applied to 800 mg (1.5 mmol) of (136) using 20 ml of anhydrous CH_3CN , 375 mg (3.0 mmol) of DMAP and 300 µl (1.6 mmol) of PTC chloride (114). After the standard work-up procedure a portion of the residue was chromatographed on a column of silica using chloroform and 1.5% methanol/chloroform as eluants. Evaporation of the appropriate fractions and crystallization of the residue from ethanol gave 136-139°C; uv (MeOH) max 278 nm (ε 16,600) (137):and 230 nm (ε 17,000), min 250 nm (ε 8,400); (0.1 N HC1) max 272 nm (ε /3,200) and 233 nm (ε 21,100), min 250 nm $(\epsilon 7,900)$ (0.1 N NaOM) max 279 nm (ϵ 16,800) and 233 nm (ϵ 21,800), min 253 nm (ϵ 7,400); M.S. m/z 626.1904 $(11\%, M^{+}-iPr = 626.1925), 516.2407 (46\%, M^{+}-0CSO_{\phi}),$ 515.2379 (100%, M+-MOCSOD), 472.1833 (11%, M+-ipr-HOCSOD), 159.0544 (9.6%, B+H); Anal. Calcd. for C31H43N5O6Si2S:. C 55.59, H 6.47, N 10.46, S 4.79. Found: C 55.20, H 6.49, N 10.74, S 4.65.

2'-Deoxy-3',5'-0-(1,1,3,3-tetraisopropyldisilox-1,3-diyl)toyocamycin (138)

The reduction conditions of Procedure C were applied to the crude residue of $(\underline{137})$ using 30 ml of toluene, 600 μ l (2.2 mmol) of \underline{n} -Bu₃SnH and 45 mg (0.3 mmol) of

After the standard work-up procedure the residue was chromatographed on a column of silica (10 g, 2×10 cm) using chloroform and 1.5% methanol/chloroform as eluants. Evaporation of the appropriate fractions and crystallization of the residue from CH3CN gave 675 mg (87%) of (138): mp 174-177°C; uv (MeOH) max 279 nm (ϵ 16,400) and 230 nm (ϵ 11,700), min 246 nm (ϵ 4,000); (0.1 N HC1) max 275 nm (ϵ 13,100) and 232 nm (ϵ 18,100), min 248 nm (ϵ 5,000); (0.1 N NaOH) max 279 nm (ϵ 15,700) and 233 nm (ϵ 10,300), min 246 nm (ϵ 4,400); 200 MHz nmr (CDC1 $_3$, TMS) δ 1.05 (m, 28, iPr), 2.43 (d of d of d, $J_{2"-1}$ ' = 2.0 Hz, $J_{2"-2}$ ' = 13.0 Hz, $J_{2"-3}$ ' = 7.2 Hz, 1, H-2"), 2.02 (d of d of d, $J_{2'-1}$ = 6.8 Hz, $J_{2'-2}$ = 13.0 Hz, $J_{2'-3'}$ = 9.8 Hz, 1, H-2'), 3.84 (d of t, $J_{4'-3'}$ = 8.0 Hz, $J_{4'-5'} = J_{4'-5''} = 3.0$ Hz, 1, H-4'), 4.03 (d of d, $J_{5"-4'} = 3.0 \text{ Hz}, J_{5"-5'} = 12.7 \text{ Hz}, 1, H-5"), 4.15 (J_{5'-4'} = 12.7 \text{ Hz})$ 3.0 Hz, $J_{5'-5''}$ = 12.7 Hz, 1, H-5'), 4.68 (m, 1, H-3'), 5.63 (s, 2, NH_2-4), 6.43 (d of d, $J_{1'-2''} = 2.0 Hz$, $J_{1'-2'} = 6.8 \text{ Hz}, 1, H-1'), 7.94 (s, 1, H-6), 8.34 (s, 1,$ H-2); M.S. m/z 517.2548 (6.4%, $M^{+}[C_{24}H_{39}N_{5}O_{4}Si_{2}] =$ 517.2543), 518.2568 (2.8%, $M^{+}+1$), 474.2001 ($M^{+}-iPr$), 159.0535 (17%, B+H); Anal. Calcd. for $C_{24}H_{39}N_5O_4Si_2$: C 55.69, H 7.59, N 13.53. Found: C 55.76, H 7.55 N 13.78.

4-Amino-5-cyano-7-(2-deoxy-β- \underline{D} -erythro-pentofuranosyl)pyrrolo[2,3-d]pyrimidine ($\underline{139}$).(2'-Deoxytoyocamycin):

Procedure D was applied to 517pomg (1 mmol) of (138)

dissolved in 20 ml of toluene using 2 ml of \underline{n} -Bu $_4$ NF solution (1 \underline{M} in THF) and stirring at 80°C for 2 h. Solvents were evaporated in vacuo and the residue was partitioned between ether and water. The aqueous phase was applied to a column of carbon (2 g, 1 x 5 cm) and the carbon was washed with 50 ml water and then 50 ml ethanol. ing was continued with chloroform and a 20% solution of benzene in chloroform was required for elution of the product. Evaporation of the appropriate fractions and crystallization of the residue from ethanol/ether gave 208-209°C; $[\alpha]_{n}^{24}$ -24.6 (c. 245 mg (89%) of (139): mр 0.28 MeOH); uv (MeOH) max 278 nm (ϵ 15,700) and 229 nm (ϵ 12,700), min 247 nm (ϵ 4,700) and 221 nm (ϵ 11,800); (0.1 \underline{N} HC1) max 273 nm (ε 12,400) and 231 \nm (ε 17,600), min 247 nm (ϵ 5,200); (0.1 NaOH) max 27/8 nm (ϵ 15,100) . and 231 nm (ε 12,000), min 246 (ε 4,800); 200 MHz nmr (DMSO- $\frac{d}{d}$ TMS) δ 2.28 (d of d of d, $J_{2"-2'} = 13.1 \text{ Hz}$, $J_{2"-1}$, = 6.1 Hz, $J_{2"-3}$, = 3.2 Hz, 1, H-2"), 2.48 (d of d of d, $J_{2'-1}$ = 7.2 Hz, $J_{2/2''}$ = 13.1 Hz, $J_{2'-3'}$ = 5.8 Hz, 1, H-2'), 3.58 (m, 2, H-5', H-5"), 3.87 (m, 1, H-4'), 4.38 (m, 1, H-3'), 5.06 (t, J_{5'.5"-OH-5'} = 5.5 Hz, 1, OH-5'), 5.32 (d, $J_{3'-0H-3'} = 4.1$ Hz, 1, 0H-3'), 6.53 (d of d, $J_{1'-2'}$ = 7.2 Hz, $J_{1'-2''}$ = 6.1 Hz, 1, H-1'), 6.88 (br s, 2, NH₂-4), 8.26 (s, 1, H-6), 8.44 (s, 1, H-2); M.S. m/z 275.1020 $(6.6\%, M^{+}[C_{12}H_{13}N_{5}O_{3}] = 275.1012), 245.0933 (2.2\%, M^{+}-CH_{2}O),$ 186.0780 (21%, BHCH*CH₂), 160.0592 (18%, B+2H), 159.0544 (100%, B+H); Anal. Calcd. for $C_{12}H_{13}N_5O_3$: C 52.34,

H 4.76, N 25.45. Found: C 52.42, H 4.83, N 25.16.

4-Amino-5-carboxamido-7-(2-deoxy- β -D-erythro-pentofurano-syl)pyrrolo[2,3-d]pyrimidine (141) (2'-Deoxysangivamycin)

A 550 mg (2 mmol) sample of (139) was applied to a column of Dowex 1 X 2 (OH $^-$) resin (2 x 10 cm). The resin was washed with 200 ml of water and then with a mixture of methanol-water (1:9). The ratio of methanol in water was gradually increased to 2:3 which eluted the product. Evaporation of the eluate and crystallization of the residue from ethanol gave 551 mg (94%) of (141): 272-275°C; $[\alpha]_{0}^{24} = -22.3$ (<u>c</u> 0.26 MeOH); uv (MeOH) max 279 nm (ϵ 14,800) and 231 nm (ϵ 10,200), min 255 nm (ϵ 6,500); (0.1 \underline{N} HC1) max 274 nm (ϵ 12,500) and 230 nm (ϵ 14,400), min 253 nm (ϵ 6,900); (β .1 NaOH) max 278 nm (ϵ 14,800), min 255 nm (ϵ 7,400) $\frac{1}{2}$ 200 MHz nmr (DMS0- \underline{d}_6 , TMS) & 2.27 (d of d of d, $J_{2"-1}$, = 6.1 Hz, $J_{2"-2'} = 12.9 \text{ Hz}, J_{2"-3'} = 3.4 \text{ Hz}, 1, H-2"), 2.42 (d of start of sta$ d of d, $J_{2'-1}$ = 7.5 Hz, $J_{2'-2''}$ = 12.9 Hz, $J_{2'-3'}$ = 5. Hz, 1, H-2'), 3.56 (m, 2, H-5",5"), 3.86 (m, 1, H-4'). 4.39 (m, 1, H-3'), 5.00 (t, $J_{5',5''-0H-5'}$ = 5.7 Hz, 1, OH-5'), 5.31 (d, $J_{3'-OH-3'} = 4.4 \text{ Hz}$, 1, OH-3'), 6.53 (d of d, $J_{1'-2'} = 7.5 \text{ Hz}, J_{1'-2''} = 6.1 \text{ Hz}, 1, H-1'), 7.36 \text{ (br s,}$ 2, NH_{2} -4) 7.94 (br s, 2, $CONH_{2}$), 8.1 (s, 1, H-6), 8.15 (s, 1, H-2); M.S. m/z 293.1128 (5.9%, $M^{+}[C_{12}H_{15}N_{5}O_{4}] =$ 293.1124), 204.0886 (12%, BHCH=CH₂), 177.0649 (100%, B+H), 176.0573 (1.6%, B); Anal. Calcd. for $C_{12}H_{15}N_50_4$:

49.15, H 5.11, N 25.44. Found: C 48.95, H 5.11, N 25.55.

3'.5'-0-(1,1,3,3-Tetraiso disilox-1,3-diyl)- cytidine (142)

The conditions of Procedure A were applied to 972 mg (4 mmol) of dried cytidine (8) using 50 ml of anhydrous pyridine and 1.3 ml (4 mmol) of TPDSC1 (95). After the standard work-up procedure crystallization of the residue from chloroform gave 1.77 g (91%) of (142) mp $267-269^{\circ}\text{C}$; uv (MeOH) max 273 nm (ϵ 8,800) min 250 nm (ϵ 6,100), (0.1 N HCl) max 280 nm (ϵ 13,100) min 240 nm (ϵ 1,400), (0.1 N NaOH) max 271 nm (ϵ 8,800), min 250 nm (ϵ 6,100); M.S. m/z 485.2384 (12%, M⁺[C₂₁H₃₉-N₃0₆Si₂] = 485.2379), 486.2399 (5.6%, M⁺+1), 442.1833 (96%, M⁺-iPr), 443.1853 (28%, M⁺+1-iPr), 140.0460 (10%, BHCHO), 112.0512 (100%, B+2H); Anal. Calcd. for C₂₁H₃₉N₃0₆Si₂: C 51.94, H 8.10, N 8.65. Found: C 52.20, H 8.07, N 8.52.

 $\frac{4-N-Acetyl-3',5'-0-(1,1,3,3-tetraisopropyldisilox-1,3-diyl)cytidine (144)}{\frac{144}{2}}$

The conditions of Procedure A were applied to 570 mg (2 mmol) of 4-N-acetylcytidine 125 (143) using 25 ml of anhydrous pyridine and 650 µl (2 mmol) of TPDSCl (95). After the standard work-up procedure crystalliza-

tion of the residue from ethanol gave 975 mg (92%) of (144); mp 111-113°C; uv (MeOH) max 297 nm (ε 8,200) and 247 nm (ε 15,000), min 270 nm (ε 4,200); (0.1 N HCl) max 305 nm (ε 11,200) and 243 nm (ε 10,300), min 268 nm (ε 3,600); M.S. m/z 527.2524 (0.3%, M⁺[C₂₃H₄₁N₃O₇Si₂] = 527.2533), 484.2006 (7.4%, M⁺-iPr), 153.0535 (1.0%, B+H). $2'-\underline{0}$ -Phenoxythiocarbony1-3',5'- $\underline{0}$ -(1,1,3,3-tetraisopropy1-disilox-1,3-diy1)cytidine (146)

To 970 mg (2 mmol) of (142) suspended in 100 ml of anhydrous CH₃CN was added 1.5 g (12 mmol) of DMAP and 800 µl (3.3 mmol) of PTC chloride (114). The resulting solution was stirred for 1 h. before 2 ml of water was added and the solvent was evaporated in vacuo. After the standard work-up procedure the residue was chromatographed on a column of silica (20 g, 2 x 20 cm) using chloroform and 2% methanol/chloroform as eluants. Evaporation of the appropriate fractions gave 700 mg (55%) of (146) as an oil with uv (MeOH) max 273 and 233 nm, min 250 nm; M.S. m/z 425.1615 (27%, M⁺-iPr-OCSOph), 424.1723 (26%, M⁺-iPr-HOCSOph), 141.0667 (24%, BCHO+2H), 111.0432 (8.2%, B+H).

 $4-\underline{N}$ -Acetyl-2'- $\underline{0}$ -phenoxythiocarbonyl-3',5'- $\underline{0}$ -(1,1,3,3-tetraisopropyldisilox-1,3-diyl)cytidine ($\underline{147}$)

To 970 g (2 mmol) of (142) suspended in 100 ml of

anhydrous CH_3CN was added 1.5 g (12 mmol) of DMAP and 800 µl (4.4 mmol) of PTC chloride (114). The resulting solution was stirred for l h before l ml (10 mmol) of acetic anhydride was added. After 30 min solvents were evaporated and the residue was subjected to the standard work-up procedure and then chromatographed on a column of silica (10 g, 2 x 10 cm) using chloroform and l% chloroform/methanol as eluants. Evaporation of the appropriate fractions gave 645 mg (49%) of (147) as an oil with uv (MeOH) max 298, 234 nm, min 270 nm; M.S. m/z 467.1835 (3.9%, M^+ -iPr-OCSO ϕ), 466.1841 (10%, M^+ -iPr-HOCSO ϕ), 183.5327 (7.9%, BCHO+2H).

4-N-Acetyl-2'-deoxy-3',5'-0-(1,1,3,3-tetraisopropyl-disilox-1,3-diyl)cytidine (148)

The conditions of Procedure C were applied to 600 mg (0.9 mmol) of (147) using 15 ml of toluene, 360 μ l (1.3 mmol) of n-Bu₃SnH and 30 mg (0.2 mmol) of AIBN. Solvents were removed in vacuo and the residue was chromatographed on a column of silica (10 g, 2 x 10 cm) to give 400 mg (87%) of (148) as an oil with uv (MeOH) max 298 nm; 200 MH2 nmr (CDCl₃, TMS) δ 1.05 (m, 28, iPr), 2.36 (d of d, $J_{2"-2'}$ = 13.5 Hz, $J_{2"-3'}$ = 7.5 Hz, l, H-2"), 2.60 (d of d of d, $J_{2'-1'}$ = 6.9 Hz, $J_{2'-2''}$ = 13.5 Hz, $J_{2'-3'}$ = 10.9 Hz, l, H-2'), 3.83 (m, l, H-4'), 4.03 (d of d, $J_{5"-5'}$ = 13.2 Hz, $J_{5"-4'}$ = 2.6 Hz, l, H-5"), 4.22 (d of d,

 $J_{5'-5''} = 13.2 \text{ Hz}, J_{5''-4'} \sim 1.5 \text{ Hz}, 1, H-5'), 4.37 \text{ (m, 1, H-3'), } 6.07 \text{ (d, } J_{1'-2'} = 6.9 \text{ Hz}, 1, H-1'), 7.43 \text{ (d, } J_{5-6} = 7.8 \text{ Hz}, 1, H-5), 8.28 \text{ (d, } J_{5-6} = 7.8 \text{ Hz}, 1, H-6); M.S. m/z 511.2524 (0.25%, M⁺[C₂₃H₄₁N₃O₆Si₂] = 511.2533), 468.2001 (9.2%, M⁺-iPr), 153.0535 (1.0%, B+H).$

2'-Deoxycytidine ($\underline{12}$)

To 350 mg (0.68 mmol) of (148) dissolved in 5 ml of tetrahydrofuran (THF) was added 2 ml (2 mmol) of a 1 M m-Bu4NF solution in THF and the solution was heated to reflux for 1 h. Solvents were removed in vacuo and the residue was treated with 1 N NaOMe in methanol for 0.5 h at 40°C. The reaction mixture was concentrated in vacuo and the residue was partitioned between ether and water. The aqueous phase was applied to a column of Dowex 1 X 2 (0H⁻) resin. Elution of the product with water, evaporation and crystallization of the residue from ethanol gave 130 mg (85%) of (12): mp 200-201°C; M.S. m/z 227.0906 (1.8%, $M^+[C_9H_{13}N_3O_4] = 227.0907$), 138.0668 (18%, $_{\pi}$ BHCH=CH2), 111.0436 (100%, BH); (1it. 149 mp 200-201°C).

Methyl 3,5-0-(1,1,3,3-tetraisopropyldisilox-1,3-diyl)- β - \underline{D} -ribofuranoside ($\underline{150}$)

The conditions of Procedure A were applied to 164 mg (1 mmol) of Methyl $\beta-\underline{D}$ -ribofuranoside (149) using

10 ml of anhydrous pyridine and 320 μ l (1 mmol) of TPDSCl (95). The standard work-up procedure was followed (excluding the acid wash) and the crude residue was chromatographed on ammonia impregnated silicá (10 g, 2 x 10 cm). The column was run quickly (to reduce hydrolysis) using Skelly B and chloroform as eluants. Evaporation of the appropriate fractions gave 325 mg (80%) of (150) as an oil. M.S. m/z 363.1686 (3.5%, M⁺-iPr[C₁₅H₃₁O₆Si₂] = 363.1661), 332.1426 (3.5%, M⁺-iPr-OCH₃), 331.1404 (9.6%, M⁺-iPr-CH₃OH).

Methyl 2-0-phenoxythiocarbonyl-3,5-0-(1,1,3,3-tetraiso-propyldisilox-1,3-diyl)- β -D-ribofuranoside (151)

The conditions of Procedure B were applied 6 300 mg (0.76 mmol) of (150) using 15 ml of anhydrous acetonitrile, 200 mg (1.5 mmol) of DMAP and 150 μl (0.80 mmol) of PTC chloride (114). The standard work-up procedure was followed (excluding the acid wash as in the previous reaction) and the residue was chromatographed on ammonia impregnated silica (10 g, 2 x 10 cm) using Skelly B and 1:1 Skelly B/chloroform as eluants. Evaporation of the appropriate fractions gave 300 mg (75%) of (151) as an oil. M.S. m/z 390.2275 (2.1%, M+-0CSOφ[C₁₈H₃₈O₅Si₂] = 390.2259), 359.2086 (9.2%, M+-OCSOφ-OMe), 316.1431 (26%, M+-OCSOφ-OMe-iPr).

To 270 mg (0.5 mmol) of (151) dissolved in 10 ml of toluene was added 200 μl (0.75 mmol) of $\underline{n}\text{-Bu}_3 SnH$ and 15 mg (0.1 mmol) of AIBN. The reduction was carried out following Procedure C. The reaction was subjected to the deblocking step of Procedure D upon completion. The solvent was evaporated in vacuo and the residue was partitioned between ether/water. The aqueous phase was evaporated to dryness and co-evaporated several times with pyridine before it was dissolved in 15 ml of anhydrous pyridine. A 5-fold molar excess of p-toluyl chloride was added to the stirred solution at room temperature. After 3 h the solvent was evaporated in vacuo and the residue was partitioned between ether and water. organic phase was washed with 2 x 10 ml of H_20 , saturated $NaHCO_3/H_2O$, saturated $NaC1/H_2O$, dried (Na_2SO_4) , filtered and evaporated. The residue was chromatographed on a column of silica (CC-7, 5 g, 1.5 x 7 cm) using Skelly B and 10% ether/Skelly B as eluants. Evaporation of the appropriate fractions and crystallization of the residue from ethanol gave 120 mg (60%) of (152): mp 76-78°C; $[\alpha]_D^{25}$ -9.4 (<u>c</u> 1.0, CHCl₃); $[(1it.^{150} m.p. 76.5-$ 78°C; $[\alpha]_D^{20}$ -8.1° (\underline{c} 2.5, $CH(1_3)$).

Treatment of 328 mg (2 mmol) of Methyl $\beta-\underline{D}$ -ribo-furanoside (149) by the five step sequence (149+150+151+152) without isolation of intermediates gave 445 mg of (152)

in 58% overall yield.

1,2;5,6-di- $\underline{0}$ -isopropylidene-3- $\underline{0}$ -phenoxythiocarbonyl- α - \underline{D} -glucofuranose ($\underline{154}$)

The conditions of Procedure B were applied to 260 mg \dagger 1 mmol) of 1,2;5,6-di-0-isopropylideneglucose (153) using 15 ml of acetomitrile and 200 ul (1.1 mmol) of PTC chloride (114). After the standard work-up procedure the residue was purified on a column of silica (10 g, 2 x 10 cm) using chloroform as the eluant. Evaporation of the appropriate fractions and crystallization of the residue from aqueous methanol gave 340 mg (86%) of (154): mp 108-110°C; 100 MHz nmr (CDC13, TMS) δ1,30-1.50 (4 x s, 12, Me), 4.10 (m, 2, H-6, H-6'), 4.30 (m, 2, H-4, H-5), 4.82 (d, $J_{1-2} = 3.8 \text{ Hz}, 1, H-2), 5.70 (m, 1, H-3), 6.00$ (d, $J_{1,2} = 3.8 \text{ Hz}$, 1, H-1), ~7.3 (m, 5, phenyl H's); M.S. m/z 396.1251 (0.25%, $M^{+}[c_{19}H_{24}O_{7}S] = 396.1230$), 381.1010 (23%, M⁺-CH₃), 243.1241 (20%, M⁺-OCSO_Φ), 185.0816 (21%, M+-OCSO+-CH3COCH3); Anat. Calcd. for C19H24O7S: C 57.56, H 6.10, S 8.09. Found: C 57.71, H 6.00, S 8.06.

3-Deoxy-1,2;5,6-di- $\underline{0}$ -isopropylidene- α - \underline{D} -ribo-hexofuranose (155)

The reducing conditions of Procedure C were applied to 792 mg (2 mmol) of (154) dissolved in 30 ml of toluene using 800 μ l (3 mmol) of \underline{n} -Bu $_3$ SnH and 64 mg (0.4 mmol) of AIBN. Solvents were removed in vacuo and the residue

was purified on a column of silica (10 g, 2 x 10 cm) using Skelly B as the eluant. Concentration of the appropriate fractions gave 425 mg (87%) of (155) as an oil; 100 MHz nmr (CDCl₃, TMS) δ -1.4 (4 x s, 12, CH₃), 1.78 (m, 1, H-3'), 2.20 (d of d, J₃₋₃, = 13.0 Hz, J₂₋₃ = 4.0 Hz, 1, H-3), 3.80 (m, 1, H-5), 4.10 (m, 3, H-4, H-6, H-6'), 4.75 (t, J₁₋₂ = J₂₋₃, = 4.0 Hz, 1, H-2), 5.78 (d, J_{1,2} = 4.0 Hz, 1, H-1); M.S. m/z 229 (47%, M⁺-CH₃), 186 (1.2%, M⁺-CH₃COCH₃); Anal. Calcd. for $C_{12}H_{20}O_5$: C 59.00, H 8.25. Found $C_{12}C_{12}O_5$: C 59.30, H 8.25.

Treatment of 520 mg (2 mmol) of (153) by the two step sequence without isolation of (154) gave 415 mg of (155) in 85% overall yield.

9-[3,5- $\underline{0}$ -(1,1,3,3-Tetraisopropyldisilox-1,3-diy])- $\underline{8}$ - \underline{D} -ribofuranosyl]purin-6-one ($\underline{166}$) (3',5'- $\underline{0}$ -TPDSinosine)

The conditions of Procedure A were applied to 134 mg (0.5 mmol) of dried inosine ($\underline{165}$) using 10 ml of anhydrous pyridine and 160 µl (0.5 mmol) of TPDSCl ($\underline{95}$). After the standard work-up procedure the residue was chromatographed on a column of silica (5 g, 1 x 10 cm) using chloroform and 5% methanol/chloroform as eluants. Evaporation of the appropriate fractions and crystallization of the residue from 1:1 acetonitrile/ethanol gave 220 mg (86%) of ($\underline{166}$): mp 219-220.5°C; uv (MeOH) max 249 nm (ε 12,100); (0.1 N HCl) max 251 nm (ε 10,700);

(0.1 N NaOH) max 253 nm (ϵ 13,000); M.S. m/z 510.2331 (4.0%, M⁺[C₂₂H₃₈N₄O₆Si₂] = 510.2328), 467.1774 (100%, M⁺-iPr), 136.0359 (30%, B+H); Anal. Calcd. for C₂₂H₃₈N₄O₆Si₂: C 51.81, H 7.51, N 10.98. Found: C 51.86, H 7.45, N 10.84.

7-Amino-3-[3,5-0-(1,1,3,3-tetraisopropyldisilox-1,3-diyl)-B-D-ribofuranosyl]pyrazolo[4,3-d]pyrimidine ($\underline{176}$) (3',5'- $\underline{0}$ -TPDSformycin)

The conditions of Procedure A were followed using 267 mg (1 mmol) of dried formycin (175), 15 ml of anhydrous pyridine and 320 μ l (1 mmol) of TPDSC1 (95). After the standard work-up procedure the crude residue was chromatographed on silica using chloroform and 2% methanol/ chloroform as eluants. Evaporation of the appropriate fractions and crystallization of the residue from $\mathrm{CH_{3}CN}$ gave 428 mg of (176): mp 149-151°C; uv (MeOH) max 294 nm (ϵ 10,300), (sh) 304 nm (ϵ 6,900), (sh) 287 nm (ϵ 9,400); (0.1 N HC1) max 295 nm (ϵ 10,700) and 232 nm (ϵ 8,800), min 266 nm (ϵ 4,400) and 222 nm (ϵ 2,900); (0.1 NaOH) max 304 nm (ϵ 7,500) and 234 nm (ϵ 16,400), min 267 nm (ϵ 3,100) and 223 nm (ϵ 10,500); M.S. m/z 509.2479 (14%, $M^{+}[C_{22}H_{39}N_{5}O_{5}Si_{2}] = 509.2480$), 510.2526 (10%, M⁺+1), 466.1936 (100%, M⁺-iPr), 178.0722 (18%, внсн₂сно), 164.0561 (65%, внсно).

Procedure A was applied to the following nucleosides on a 0.1 mmol scale using 2.5 ml of pyridine and 32 µl (0.1 mml) of TPDSC1 (95). The standard work-up procedure was, followed and the residues were chromatographed on silica plates (5 x 20 cm) using 5% methanol/chloroform as eluant. The silica which contained the product was scraped off the plate and eluted with 7% methanol/chloroform. Evaporation of the eluate gave an oil in each case.

2,6-Diamino-9- β - \underline{D} -ribofuranosylpurine $(\underline{167})$

 $5-\beta-\underline{D}$ -Ribofuranosyluracil (169) (pseudouridine)

4-Hydroxy-1- β - \underline{D} -ribofuranosylpyridin-2-one ($\underline{171}$) (3-deazauridine)

4-Amino-1- β - \underline{D} -ribofuranosylpyridin-2-one ($\underline{173}$) (3-deazacytidine)

5(3)-Carboxamido-4-hydroxy-3(5)- β - \underline{D} -ribofuranosylpyrazole (177) (pyrazomycin)

 $9-\alpha-D$ -Riboturanosyladenine (179) (α -adenosine)

 $1-\alpha-\underline{D}$ -Ribofuranosyluracil (181) (α -uridine)

 $1-\alpha-\underline{D}$ -Ribofuranosylcytosine (183) (α -cytidine)

 $9-\beta-\underline{D}$ -Arabinofuranosyladenine (159)

 $1-\beta-\underline{D}$ -Arabinofuranosylcytosine ($\underline{189}$)

9-(2-Chloro-2-deoxý-β-D-arabinofuranosyl)adenine (<u>156</u>)

9-(2-Azido-2-deoxy- β - \underline{D} -arabinofuranosyl)adenine (187)

 $9-\alpha-D_{c}$ Arabinofuranosyladenine (185)

3',5'-0-(1,1,3,3-Tetraisopropyldisilox-1,3-diyl)nucleosides (3',5'-0-TPDSnucleosides)

2,6-Diamino-9-(3,5- $\underline{0}$ -TPDS- β - \underline{D} -ribofuranosyl)purine ($\underline{168}$)

The residue obtained was crystallized from ethanol to give white needles with mp 190-192°C; uv (MeOH) max 280 nm (ε 10,700), 258 nm (ε 9,900) and 217 nm (ε 25,200), min 268 nm (ε 8,400) and 235 nm (ε 4,200); (0.1 \underline{N} HC1) max 292 nm (ε 8,800), 252 nm (ε 10,300) and 214 nm (ε 15,500), min 270 nm (ε 5,200) and 237 nm (ε 6,100), (0.1 \underline{N} NaOH) max 280 nm (ε 10,800) and 256 nm (ε 9,900), min 265 nm (ε 8,100) and 239 nm (ε 6,600°); M.S. m/z 524.2598 (18%, M⁺[C₂₂H₄₀N₆O₅Si₂] = 524.2599), 481.2053 (16%, M⁺-iPr), 150.0654 (53%, B+H).

$3',5'-\underline{0}$ -TPDSpseudouridine ($\underline{170}$)

Uv (MeOH) max 260 nm; M.S. m/z 443.1677 (9.5%, M^{+} -iPr[C₁₈H₃₁N₂O₇Si₂] = 443.1662), 141.0350 (100%, BHCHO).

$3',5'-\underline{0}$ -TPDS-3-deazauridine ($\underline{172}$)

Uv (MeOH) max 280 nm; M.S. m/z 485.2262 (12%, $M^{+}[C_{22}H_{39}NO_{7}Si_{2}] = 485.2265$), 442.1718 (100%, $M^{+}-iPr$), 112.0398 (83%, B+2H), 111.0317 (12%, B+H).

$3',5'-\underline{0}$ -TPDS-3-deazacytidine ($\underline{174}$)

Uv (MeOH) max 262 nm; M.S. m/z 484.2429 (1.8%, $M^{+}[C_{22}H_{40}N_{2}O_{6}Si_{2}] = 484.2424$), 485.2481 (3%, $M^{+}+1$),

441.1870 (10%, M⁺-iPr), 442.1927 (19%, M⁺+1-iPr), 139.0503 (5.4%, BHCHO), 111.0557 (53%, B+2H).

$3',5'-\underline{0}$ -TPDSpyrazomycin ($\underline{178}$)

Uv (MeOH) max 266 nm; M.S. m/z 458.1775 (13%, M^{+} -iPr[C₁₈H₃₂N₃O₇Si₂] = 458.1779), 170.0565 (19%, BHCH₂CHO), 156.0414 (2.3%, BHCHO).

3',5'-0-TPDS- α -adenosine (180)

Uv (MeOH) max 260 nm; M.S. m/z 509.2489 (6.1%, $M^{+}[C_{22}H_{39}N_{5}O_{5}Si_{2}^{N}] = 509.2489$), 466.1942 (100%, $M^{+}-iPr$), 178.0728 (5.0%, BHCH₂CHO), 164.0573 (23%, BHCHO), 136.0623 (17%, B+2H), 135.0546 (4.9%, B+H).

3',5'-0-TPDS- α -uridine ($\frac{1.82}{}$)

Uv (MeOH) max 262 nm; M.S. m/z 443.1677 (67%, $M^{+}-iPr[C_{18}H_{31}N_{2}O_{7}Si_{2}] = 443.1670)$, 444.1728 (100%, $M^{+}+1-iPr$), 155.0415 (3.2%, BHCH₂CHO), 113.0354 (3.7%, B+2H), 112.0278 (1.1%, B+H).

3',5'- $\underline{0}$ -TPDS- α -cytidine ($\underline{184}$)

Uv (MeOH) max 270 nm; M.S. m/z 485.2380 (9.0%, $M^{+}[C_{2}H_{39}N_{3}O_{6}Si_{2}] = 485.2378$), 486.2398 (4.3%, $M^{+}+1$), 442.1832 (100%, $M^{+}-iPr$), 140.0462 (8.7%, BHCHO), 112.0512 (95%, B+2H).

9-(3,5-0-TPDS- β -D-arabinofuranosyl)adenine (160) Uv (MeOH) max 260 nm; M.S. m/z 509.2483 (5.2%, $M^{+}[C_{22}H_{39}N_{5}O_{5}Si_{2}] = 509.2490), 466.1927 (100%, <math>M^{+}-iPr),$ 164.0573 (11%, BHCHO), 136.0621 (19%, B+2H), 135.0545 (3.6%, B+H).

1-(3,5-0-TPDS-B-D-arabinofuranosyl)cytosine (190) Uv (MeOH) Max 270 nm; M.S. m/z 485.2386 (1.0%, M⁺[C₂₁H₃₉N₃O₆Si₂] = 485.2378), 442.1832 (10%, M⁺-iPr), 140.0463 (1.1%, BHCHO), 112.0465 (11%, B+2H), 111.0433 (100%, B+H).

9-(2-Chloro-2-deoxy-3,5- $\underline{0}$ -TPDS- β - \underline{D} -arabinofuranosyl)-adenine ($\underline{164}$)

Uv (MeOH) max 259 nm; M.S. m/z 527.2170 (8.0%, $M^{+}[C_{22}H_{38}N_{5}O_{4}Si_{2}C1] = 527.2167)$, 529.2160 (2.6%, $M^{+}+2$), 484.1612 (100%, $M^{+}-iPr$), 486.1603 (33%, $M^{+}+2-iPr$), 164.0574 (10%, BHCHO), 136.0625 (17%, B+2H).

9-(2-Azido-2-deoxy-3,5- $\underline{0}$ -TPDS- β - \underline{D} -arabinofuranosyl)-adenine ($\underline{188}$)

Uv (MeOH) max 261 nm; M.S. m/z 534.2521 (2.9%, $M^{+}[C_{22}H_{38}N_{8}O_{4}Si_{2}] = 534.2557$), 536.2642 (9.0%, $M^{+}+2$), 535.2590 (9.9%, $M^{+}+1$), 493.2129 (79%, $M^{+}+2-iPr$), 492.2074 (85%, $M^{+}+1-iPr$), 491.2008 (30%, $M^{+}-iPr$), 463.1982 (1.6%, $M^{+}-N_{2}-iPr$), 164.0573 (29%, BHCHO).

9-(3,5-0-TPDS- α -D-arabinofuranosyl)adenine (186) - Uv (MeOH) max 260 nm; M.S. m/z 509.2484 (2.0%, M+[C₂₂H₃₉N₅O₅Si₂] = 509.2490), 467.1974 (42%, M+1-iPr), 466.1945 (100%, M⁺-iPr), 164.0573 (15%, BHCHO), 136.0622 (62%, B+2H), 135.0546 (6.7%, B+H).

 $6-\underline{N}$ -Acetyl-9-(2- $\underline{0}$ -acetyl-3,5- $\underline{0}$ -TPDS-8- \underline{D} -arabinofurano-syl)adenine ($\underline{162}$)

Procedure A was followed using 27 mg (0.1 mmol) of $9-B-\underline{D}$ -Arabinofuranosyladenine ($\underline{159}$), 5.0 ml of pyridine and 32 µl (0.1 mmol) of TPDSC1 ($\underline{95}$). After a 3 h period, 20 µl (2 mmol) of acetic anhydride was: added to the reaction solution and stirring was continued for 4 h. Solvents were evaporated and the residue was chromatographed on a column of silica (6 g, 1.5 x 7 cm) using chloroform as the eluant. Evaporation of the appropriate fractions gave 45 mg (82%) of ($\underline{162}$) as an oil with uv (MeOH) 27l nm; M.S. m/z 593.2704 (2.3%, \underline{M}^{\dagger} [C₂₆H₄₃N₅O₇Si₂] = 593.2701), 550.2150 (100%, \underline{M}^{\dagger} -iPr), 509.2076 (15%, \underline{M}^{\dagger} -iPr-OAc), 508.2065 (38%, \underline{M}^{\dagger} -iPr-HOAc), 261.1710 (§.1%, BHCH=CHOAc), 178.0725 (17%, B+2H), 179.0646 (3.6%, B+H).

6-N-Acetyl-9-(2- $\underline{0}$ -acetyl-3,5- $\underline{0}$ -TPDS- α - \underline{D} -arabinofurano-syl)adenine ($\underline{191}$)

The procedure outlined above was applied to 27 mg (0.1 mmol) of $9-\alpha-D$ -arabinofuranosyladenine (185) using 5.0 ml of pyridine, 32 µl (0.1 mmol) of TPDSCl (95) and 20 µl (2 mmol) of acetic anhydride. A 46 mg (84%)

2-Deutero-2'-deoxyadenosines (157a, 157b)

a) The reduction of a 65 mg (0.1 mmol) sample of (125) was carried out as described for (127) substituting n-Bu₃SnD for n-Bu₃SnH. The product obtained showed the identical chromatographic mobility of (127). The deblocking procedure described for (10) was applied directly to the reduction solution and the analogous work-up was continued to give 22.5 mg (90%) of (157a, 157b) as an oil. 400 MHz nmr (DMSO-d₆, TMS) & 2.30 (d of d, $J_{2''-1}$, = 2.8 Hz, $J_{2''-3}$, = 5.8 Hz, ~0.12, H-2" of 157b), 2.71 (d of d, $J_{2''-1}$, = 7.5 Hz, $J_{2''-3}$, = 5.8 Hz, ~0.88, H-2' of 157a), 3.57 (d of d, $J_{5''-4}$, = 4.2 Hz, $J_{5'''-5}$, = 12.0 Hz, 1, H-5"), 3.68 (d of d, $J_{5''-4}$, = 4.2 Hz, $J_{5'''-5}$, = 12.0 Hz, 1, H-5'), 3.94 (m, 1, H-4'), 4.46 (d

- of d, $J_{3,-4}$ = 2.8 Hz, $J_{3,-2}$ = 5.8 Hz, 1, H-3'), 5.35 (m, 2, 3' and 5' 0H), 6.39 (d, $J_{1,-2}$ = 7.5 Hz, 1, H-1'), 7.29 (s, 2, 6-NH₂), 8.18 (s, 1, H-2), 8.36 (s, 1, H-8); M.S. m/z 252.1075 (6.7%, M [C₁₀H₁₂DN₅O₃] = 252.1082), 163.0835 (67%, BHCH=CHD), 135.0662 (100%, B+H).
- b) The four step sequence outlined in a) for the 2'-deoxygenation of adenosine was applied to 53 mg (0.2 mmol) of $9-\beta-\underline{D}$ -arabinofuranosyladenine (159) without purification of the intermediates. A 34 mg (68%) yield of the 2'-deoxy-2'-deutero-adenosines (157a, 157b) was obtained as an oil.
- c) A 53 mg (0.1 mmol) sample of 9-(2-chloro-2-deoxy-3,5- $\underline{0}$ -TPDS- β - \underline{D} -arabinofuranosyl)adenine ($\underline{164}$) was subjected to the reduction conditions of Procedure C using 10 ml of toluene, 40 µl (0.15 mmol) of \underline{n} -Bu₃SnD, and 5 mg of AIBN. Deblocking via Procedure D was performed directly on the reaction solution using 250 µl of \underline{n} -Bu₄NF (as a l M solution in THF). A 22 mg (88%) yield of ($\underline{157a}$, $\underline{157b}$) was obtained as an oil. The spectral data for the mixture of ($\underline{157a}$, $\underline{157b}$) obtained via b) or c) corresponded exactly to those of the mixture of ($\underline{157a}$, $\underline{157b}$) from procedure a) above.

Methyl 3,5-0-(1,1,3,3-tetraisopropyldisilox-1,3-diyl)- α -D-ribofuranoside (193)

The procedure outlined for the synthesis of (150) was applied to 33 mg (0.2 mmol) of methyl α -D-ribofuranoside (192). An 81% (64 mg) yield of (193) was obtained as an oil. M.S. m/z 363.1686 (33%, M⁺-iPr[C₁₅H₃₁O₆Si₂] = 363.1661), 332.1413 (12%, M⁺-iPr-OCH₃), 331.1365 (48%, M⁺-iPr-CH₃OH).

4-N-Acetyl-1-[2-0-acetyl-3,5-0-(1,1,3,3-tetraisopropyl)-disilox-1,3-diyl)- β -D-ribofuranosyl]cytosine (145)

To 57 mg (0.1 mmol) of $4-\underline{N}$ -acetyl-1-(3,5- $\underline{0}$ -TPDS- β - \underline{D} ribofuranosyl)cytosine ($\underline{144}$) and 3.6 mg (0.3 mmol) of DMAP dissolved in 5 ml of anhydrous CH $_3$ CN was added 20 $\mu 1$ (0.2 mmol) of acetic anhydride. After a 3 h period the reaction solution was concentrated in vacuo. The residue was subjected to the standard work-up procedure and chromatographed on a column of silica (5 g, 1.5 x 6 cm) using 1% methanol in chloroform as the eluant. Evaporation of the appropriate fractions gave 49 mg (86%) of (145) as an oil with uv (MeQH) max 297 min; 400 MHz nmr δ ~1.05 (m, 28, iPr H's), 2.10 (2 x s, 6, $4-\underline{N}-COCH_3$, $2'-\underline{O}-COCH_3$), 4.00 (m, 3, H-4', H-5', H-5"), 4.43 (d of d, $J_{3'-2'} = 5.4$ Hz, $J_{3'-4'} = 8.6$ Hz, 1, H-3'), 5.47 (d, $J_{2'-3'} = 5.4$ Hz, 1, H-2'), 5.72 (s, 1, H-1'), 7.22 (d, $J_{5-6} = 7.6 \text{ Hz}$, 1, H-5), 8.04 (d, J_{5-6} = 7.6 Hz, 1, H-6); M.S. m/z 526.2034 (13)%, M^+ -iPr[C₂₂H₃₆- $N_3 O_8 Si_2$] = 526.2041), 235.1185 (36%, BHCH=CHOCOCH₃), 154.0619 (24%, B+2H), 153.0541 (3.9%, B+H).

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