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Modular synthesis of 1-α- and 1-β-(indol-2-yl)-2'-deoxyribose C-nucleosides[†]

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A simple two-step method for the selective preparation of anomerically pure 1α - and 1β -(indol-2-yl)deoxyribose derivatives was developed. The synthesis was based on the Sonogashira reaction of 1α - and 1β -ethynyldeoxyribose and 2haloanilines followed by a Pd-complex catalyzed cyclization to the corresponding indolyldeoxyribosides.

C-Nucleosides¹ are an important class of compounds with potent biological activities and applications in chemical biology. Development of modular methodologies² for their synthesis is of continuing interest in our laboratories. Nitroindoles (in Nnucleosides) are used³ as universal nucleobases that do not discriminate the opposite base in DNA or for studying basestacking interactions.⁴ Several 3-⁵ and 2-linked^{6,7} indole Cribo and -deoxyribonucleosides have been prepared, usually by addition of highly reactive lithiated protected indoles to sugar lactones or hemiacetals followed by reduction or Mitsunobu cyclization. Indol-3-yl C-deoxyribonucleoside has been used5a as a base substitution in DNA, whereas unsubstituted indol-2-yl-2'-deoxyribonucleoside has been utilized⁷ as an artificial base in extension of the genetic alphabet, taking advantage of the minorgroove interaction of its NH. Difficult syntheses and limited access to series of highly substituted derivatives (especially those bearing reactive functional groups incompatible with organolithium chemistry) by the previously known methods⁵⁻⁷ prevent wider studies of potential applications of indole C-nucleosides in chemical biology.

In the past decade this laboratory and others have shown that 1α - and 1β -alkynyl deoxyribose derivatives could be used as suitable synthetic building blocks for the preparation of various *C*-1-substituted derivatives.⁸ As typical examples may serve syntheses of *C*-aryl derivatives by using Rh-⁹ or Ru-catalyzed¹⁰ cyclotrimerization, various substituted alkynyl derivatives by using the Sonogashira coupling or other procedures,¹¹⁻¹³ and triazoles by click-chemistry.^{14,15} The main advantage of these procedures stemmed from the fact that the corresponding 1α - and 1β -ethynyl derivatives can be prepared in anomerically pure forms.

It has been shown in numerous examples that terminal alkynes could serve as building blocks for the synthesis of variously substituted indoles by coupling them with suitably substituted 2-haloanilines.^{16,17} On the other hand it is fair to note that most of the alkyne couplings were carried out with phenylacetylenes and not with alkyl-substituted acetylenes. Nonetheless, we envisioned that the desired indolyldeoxyribosides could be also approached by the same strategy, *i.e.* by coupling of ethynyldeoxyribosides with 2-haloanilines.

The first task was the development of the high yielding Sonogashira reaction of 1-ethynyl-2'-deoxyribose with haloanilines, because undesirable homocoupling to give 1,3-diynes was observed by us¹ as well as by others.¹¹ The coupling of 1α -ethynyldeoxyribose 1α with *N*,*N*-dimethyl-2-iodoaniline under various conditions (Scheme 1) was chosen as a model reaction to tune the reaction conditions.



Scheme 1 Sonogashira reactions of 4a-4d with 1 and Pd-catalyzed cyclizations.

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С	Tol Me;		conditions TolO 3a				
	1α	2a		Ĺ			
Entry	Catalyst (5 mol%)	Cocatalyst	Base	Solvent	Yield (%) ^a		
1	$PdCl_2(PPh_3)_2$ $PdCl_2(PPh_3)_2$	CuI	Et ₃ N	MeCN	25		
3	$PdCl_2(PPII_3)_2$ $PdCl_2(MeCN)_2^{b}$	—	Cs_2CO_3	MeCN	0 4		
4	$PdCl_2(PPh_3)_2$ $Pd(PPh_3)_2$	—	piperidine	—	27		
6	$Pd(PPh_3)_4$ Pd(PPh_3)_4	_	piperidine	THF	26		
7 8	$Pd(PPh_3)_4$ $Pd(PPh_3)_4$	_	piperidine piperidine	DMF MeCN	44 52		
" Isolat triisopi	ed yields. ropylbiphenyl (15 m	^{<i>b</i>} Ligand: ol.%)	2-dicycloh	exylphosp	ohino-2,4,6-		

The coupling (Table 1) under classical conditions (PdCl₂(PPh₃)₂, CuI, Et₃N) proceeded in MeCN to yield **3a** in only 25% yield (Entry 1). When the reaction was carried out in THF, it did not proceed (Entry 2). An attempt to apply the previously used¹³ copper-free conditions in the presence of a sterically hindered phosphine and caesium carbonate¹⁸ gave the title compound only in a marginal yield of 4% (Entry 3). The change of a base to piperidine gave rise to the title compound again in low yield of 27% (Entry 4). The use of Pd(PPh₃)₄ either in pure piperidine or in a mixture with THF gave **3a** in 26% yields (Entries 5 and 6). A slight increase in yield (44%) was observed in DMF (Entry 7). Finally, the best yield of the title compound (52%) was achieved in MeCN with piperidine as a base (Entry 8).

With the required product 3α on hand the intramolecular cyclization under Larock's conditions (I₂, CH₂Cl₂) was attempted.¹⁹ Unfortunately, the cyclization did not proceed and the starting material was recovered intact.

In order to achieve our goal, we decided to change the haloaniline **2** to the protected 2-iodophenyltrifluoroacetamides **4a–4d**, prepared from the corresponding anilines and (CF₃CO)₂O, and use them in a similar reaction.^{16b} However, the above used conditions proved not to be suitable for Sonogashira couplings of **1** α and **1** β with **4a–4d** and the best yields were obtained with a combination of Pd(PPh₃)₄/Et₃N/MeCN/70 °C (Table 2).

The Sonogashira reaction of 1α and 1β (Table 1) with a simple acetamide 4a, two chloro derivatives 4b and 4c, carboxyacetamide 4d furnished the corresponding alkynyldeoxyribose derivatives $5\alpha a - 5\alpha d$ (Entries 1-4) and $5\beta a - 5\beta c$ (Entries 5-8) in reasonable isolated yields.

Although it was reported that spontaneous addition of the N–H bond to the triple bond in 2-alkynylaniline derivatives followed the Sonogashira coupling forming the indole scaffold, we did not observe such a process in any case. Thus it was necessary to find suitable conditions for the cyclization. After a number of experiments it was found that in the presence of $Pd_2(dba)_3 \cdot CHCl_3$ (5 mol%)/(2-biphenyl)di-*t*-butylphosphine (Johnphos, 20 mol%)/120 °C/5 h and then rt (usually 18–20 °C) overnight the cyclization took place.

Ensuing workup of the reaction mixtures proceeded with spontaneous removal of the trifluoroacetyl group yielding the desired indolyl derivatives 6.

The cyclization of $5\alpha a$ to $6\alpha a$ proceeded with a very nice yield of 81% (Entry 1). Cyclizations of $5\alpha b$ – $5\alpha d$ gave indoles $6\alpha b$ – $6\alpha d$ in 57, 48, and 40% isolated yields, respectively (Entries 2–4). Cyclizations of the beta anomeric substrates $5\beta a$ – $5\beta c$ gave indoles $6\beta a$ – $6\beta c$ in 51, 42, and 43% isolated yields, respectively (Entries 5–7). Only in the case of $5\beta d$, was $6\beta d$ obtained in a low yield of 19% (Entry 8)

The last step was the deprotection of the toluoylated indolyldeoxyribosides. Since the removal of the toluoyl protective group under basic conditions is a well established procedure, we decided to use the same method: samples were treated with a solution of NaOMe in MeOH 12 h at rt. The deprotection proceeded uneventfully in all cases with both anomers, 6α and 6β , yielding the corresponding free indolyldeoxyriboses 7α and 7β (Fig. 1). The isolation had to be carried out by using preparative TLC and the yields were in the range of 42–76%, which was typical for this method.



Fig. 1 The prepared free indolyldeoxyriboses 7α and 7β .

In summary we have developed a novel methodology suitable for the selective synthesis of α - and β -anomeric indolyldeoxyribosides.²⁰ This method is quite general and nodular with respect to the substitution pattern of the aniline leading to diverse types of indolyl-*C*-nucleosides. Moreover, it is expected that this method could be extended to other *C*-ethynylsaccharides leading to analogues of naturally occurring indole glycoside antibiotics²¹ and indolyl *C*-mannosides.²²

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Table 2Cross-coupling of 1 with 4 to 5 and cyclization to 6

Entry	Iodoaniline 4	Product 5		Yield (%) ^{<i>a</i>}	Product 6		Yield (%) ^a
1	4a		5αa	77		6αa	81
2	4b		5ab	55		6αb	57
3	4c		5ας	47		6ac	48
4	4d	TolO TolO NHCOCF ₃	5ad	28		6ad	40
5	4a	TolO TolO TolO	5βa	77	TolO N TolO	бβа	51
6	4b	TolO TolO TolO	5βb	62		бβЬ	42
7	4c	TolO TolO TolO	5βc	57		бβс	43
8	4d	TolO TolO TolO	5βd	33	TolO TolO	6βd	19
^a Isolated	1 yields.						

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- 20 A typical procedure: synthesis of 6ba. To a solution of 5ba (262 mg, 0.46 mmol) in MeCN (3 mL) was added Pd₂(dba)₃·CHCl₃ (17.3 mg, 0.016 mmol), (2-biphenyl)di-t-butylphosphine(41 mg, 0.14 mmol) and K₂CO₃ (200 mg) and the reaction mixture was heated to 120 °C for 5 h. Then it was stirred at rt (usually 18-20 °C) overnight. Work-up followed by column chromatography on silica gel yielded 111 mg (51%) of the title compound as a yellowish syrup: ¹H NMR (600 MHz, C₆D₆) δ 8.57 (s, 1H), 8.08 (dd, J = 30.9, 8.1 Hz, 4H), 7.66 (d, J = 7.6 Hz, 1H), 7.25-7.18 (m, 3H), 6.97 (d, J = 7.9 Hz, 2H), 6.80 (d, J = 8.0 Hz, 2H), 6.33 (bs, 1H), 5.35 (d, J = 5.7 Hz, 1H), 5.30 (dd, J = 10.4, 5.5 Hz, 1H), 4.54 (dd, J = 11.6, 5.7 Hz, 1H), 4.43–4.34 (m, 2H), 2.19 (dd, J = 13.5, 5.1 Hz, 1H), 2.13-2.05 (m, 1H), 2.02 (s, 3H), 1.91 (s, 3H); ¹³C NMR (151 MHz, C₆D₆) δ 167.27, 166.28, 144.47, 144.27, 138.31, 137.24, 130.50 (2C), 130.45 (2C), 129.84 (2C), 129.82 (2C), 129.40, 128.31, 128.05, 122.63, 121.27, 120.57, 111.88, 100.28, 83.97, 77.58, 76.30, 65.31, 40.44, 21.80, 21.68; IR (KBr) 3352, 2949, 1715, 1610, 1456, 1271, 1178, 1105, 752 cm⁻¹; HRMS calcd for C₂₉H₂₇NO₅(M + H) 470.1962, found 470.1964.
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