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Concurrent Ring Opening and Halogenation of Spiroketals¹

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Abstract: Ring opening of various spiroketals with triphenylphosphine dihalides under neutral conditions produced ω halo-enolethers in good to excellent yield. The method transformed even the very stable spiroketal of hecogenin acetate at temperatures below any previously reported for such isomerative opening. © 1999 Elsevier Science Ltd. All rights reserved.

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The structural challenges and extreme antitumor activities displayed by the cephalostatins² and ritterazines³ render them of interest.⁴ Recently, we reported the first syntheses of cephalostatin 1 (GI_{50} 1.2 nM)² and a very potent "interphylal" hybrid analogue of these marine products, ritterostatin $G_N I_N I$ (GI_{50} 14 nM),⁵ which incorporated the North G (2) subunit of ritterazine G.³ Derivation of all subunits of these antineoplastics from commercially available hecogenin acetate 3 included an acid catalyzed isomerative spiroketal opening^{5,6} with a carboxylic anhydride (ex. $3\rightarrow4a$, Fig. 1, wherein esterification traps the fleeting intermediate pseudosapogenin 4b). The otherwise efficient synthesis of 2 (13 steps, 15% overall) suffered somewhat in the $3\rightarrow5$ sequence. Conversion of 3 to 4a proved particularly capricious, difficult to scale, required a demanding chromatographic separation and involved subsequent conversion of the sensitive alcohol 4b to obtain 4c, 4d and finally 5.

Use of dichloroacetic anhydride^{6a,b} originally offered an improvement for steroid spiroketal opening in that the C3 acetate is retained, whereas traditional anhydride methods⁷ result in C3 transesterification and identical C3/C26 functionality in the derived pseudosapogenins. These positions show surprisingly similar reactivity, and differentiation of C3 and C26 of 5,6-saturated sapogenins has therefore proven historically nontrivial.^{8,9} Herein we report a general method that affords *one-step* access to useful **4d/e** type intermediates from spiroketals in good to excellent yields by application of a phosphine dihalide under *neutral conditions, at temperatures below any previously reported for isomerative openings*, followed by filtration through silica.



(a) (Cl₂HCCO)₂O, PPTs, 35-70%; (b) KHCO₃, 80-99%; (c) TsCl, 70-91%; (d) Nal, 99%; (e) DBU, 91%; (f) aq. AcOH, 98%



The new method sprang from attempts to bypass tosylate 4c by direct conversion of alcohol 4b to iodide 4d (Ph₃P•I₂, imidazole, 2-12 h).¹⁰ These reactions returned mainly spiroketals 3 (thermodynamic product) and 6 (kinetic product, Fig. 2),¹¹ by acid-catalyzed cyclization. The lability of alcohols such as 4b to even trace acid was also evident during tosylation, which gave 4c (70-90%, 40% if run in CH₂Cl₂ instead of pyridine solvent) accompanied by 3 and 6. However, a Ph₃P•I₂ run allowed to stand 4 days delivered 4d (40%) and 3 (55%). Although 6 was found to slowly afford mainly 4d at 25 °C with this reagent system, spiroketal 3 was inert.

Persistent experimentation revealed that transformation of 3 to halides 4 occurred under these essentially neutral conditions at temperatures (130-140 °C) still well below those required by anhydride-mediated openings (180-240 °C),^{6,7} without loss of the 3-acetate or C25 epimerization¹² (Table 1). Further transformation of 4d to 7, obtained as a single diastereomer (confirmed by X-ray of hydrate 8, Fig. 2)¹³ was accomplished by merely extending the reaction time or by resubjecting 4d to the reaction conditions. Heating 4d (alone or with base, TCE, 150 °C, 2 h) returned only starting iodide with no trace of 7.

Table 1. Conditions^a for isomerative opening of spiroketal 3 with concurrent C26-halogenation.

entry	reagent	base ^b	temp.	time	4 ^c (halide ratio) ¹³	7 °
1	1.1 eq Ph ₃ P•I ₂	2.2 eq imidazole	140 °C	2 h	trace	-
2	2.2 eq Ph ₃ P•I ₂	4.4 eq imidazole	140 °C	2 h	60% (1:6 d/e)	30%
3	2.2 eq Ph ₃ P•I ₂	4.4 eq DBMP	140 °C	2 h	62% (1:6 d/e)	29%
4	2.2 eq Ph ₃ P•I ₂	2.2 eq imidazole	140 °C	1 h	67% (5:1 d/e)	25%
5	2.2 eq Ph ₃ P•I ₂	2.2 eq imidazole	125 °C	2.5 h	30% (2:1 d/e)	62%
6	2.2 eq Ph ₃ P•I ₂	2.2 eq 2,6-lutidine	140 °C	1 h	69% (5:1 d/e)	24%
7	2.2 eq Ph ₃ P•I ₂	0.8 eq 2,6-lutidine	140 °C	0.5 h	75% (8:1 d/e)	18%
8	2.2 eq Ph ₃ P•I ₂	0.8 eq 2,6-lutidine	140 °C	20 min ^d	60% (25:1 d/e) ^d	5%
9	2.2 eq Ph ₃ P	2.2 eq imidazole	140 °C	2 h	85% (e only)	9%
10	2.2 eq Ph ₃ P•Cl ₂	0.8 eq imidazole	140 °C	1 h	90% (e only)	5%
11	2.2 eq Ph ₃ P•I ₂	0.8 eq 2,6-lutidine	140 °C	3 h	8% (e only)	87%

^a 2 mmol 3, 0.1 M in TCE. ^b DBMP = 2,6-di-t-butyl-4-methylpyridine. ^c Isolated yields. ^d 25% unreacted 3.

Since aliphatic chlorinated solvents, which dissolve the reagent,^{10c} were by far superior for the transformation, some chloride **4e** accompanied iodide **4d**. Tetrachloroethane (TCE) permitted sufficient temperature for smooth conversion, but no reaction occurred below 125 °C. Inferior reaction or decomposition was observed in other solvents, including bromo and iodoalkanes, DMF, aliphatic and aromatic hydrocarbons, ethers, nitriles, and aromatic halides. Every soluble "non-nucleophilic" amine base examined served equally well, regardless of steric demand or base strength. Insoluble bases (PVP, alkali carbonates and hydrides) proved unsatisfactory.¹⁴ Minimizing the amount of base decreased reaction time and reduced formation of chloride **4e**. The coproduct Ph₃PO may sequester some HI as Ph₃P(OH)I¹⁵ but use of less than 0.8 eq amine base resulted in HI-catalyzed reactions.¹⁴ Chloride formation may arise from dissociation of the phosphine diiodide (free I₂ was noted upon heating) and reaction of liberated Ph₃P with solvent to give chlorophosphonium salts. Indeed, heating **3** with Ph₃P in TCE afforded **4e** in high yield, and use of Ph₃P•Cl₂ delivered an excellent yield of **4e**. Cyclization of **4e** to give **7** was very slow. Amine bases may promote Ph₃P•I₂ dissociation by forming unrproductive amine diiodides (unlike Ph₃P•Br₂, pyridine•Br₂ does not convert acetals to RBr).^{17a} Bulky base-induced solvent elimination, releasing chloride ion for S_N2 reaction with **4d**, seems less probable.



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	spiroketal	eq Ph ₃ P•I ₂ ; eq. base	solvent, temp, time ^a	yields (I / Cl ratio)b	
1	6 - 5α,9α,20β; Y=Z=O, R=H, R'=Me	1.1; 1.1 imidazole	CH ₂ Cl ₂ , 42 °C, 36 h	82% 4 (17:1 a/b)	12% 3
2	6	1.1; 1.1 imidazole	CH2Cl2/CH3CN, 70 °C, 5 h	86% 4 (14:1 a/b)	7% 3
3	9 - 5 β ,9 α ,20 α ; Y=Z=H, R=Me, R'=H	1.1; 1.1 imidazole	CH2Cl2/CH3CN, 70 °C, 5 h	89% 13 (15:1 a/b)	-
4	10 - 5α,9α,20α; Y=Z=R≈H, R'=Me	2.2; 0.8 lutidine	TCE, 140 °C, 0.5 h	82% 14 (8:1 a/b)	15% 17
5	11 - 5 α , $\Delta^{9(11)}$,20 α ; Y=Z=O, R=H, R'=Me	2.2; 0.8 lutidine	TCE, 140 °C, 0.5 h	76% 15 (8:1 a/b)	1 4% 18
6	12 · 5α,9α,20α; Y=OAc, Z=R=H, R'=Me	2.2; 0.8 lutidine	TCE, 140 °C, 0.5 h	83% 16 (8:1 a/b)	15% 19

^a All reactions run at 0.1 M. ^b Isolated yield; halide ratio determined by integration of the HNMR signals for the 26-CH₂ protons.¹³

The scope of the reaction was briefly examined (Fig. 3 and Table 2). Less stable spiroketal types (e.g. 6 or 9)¹¹ reacted smoothly at much lower temperatures (25-70 °C). Compatible functionality was found to include ester, ketone, and enone (and, of course, enolether and halide) moieties. Isolated alkenes are evidently also tolerated, as resubjection of compounds 7 and 18 returned starting materials unchanged. The absence of α -halogenated ketone products indicates that R₃NHX₃ or R₄PX₃ type reagents¹⁶ are probably not produced.

Plausible mechanistic rationales for formation of ω -halo-enolethers **4**, **13-16** under these neutral conditions are postulated in Fig. 4. Path "a" differs from that of the anhydride methods and the (acidic) Ph₃P•Br₂ reaction of THP-protected alcohols and other acetals.¹⁷ In the latter protocol, initial formation of an enolether, acid (HBr) and an oxophosphonium salt is observed. In a spiroketal system, this scenario (path b) would furnish a tethered C26-oxophosphonium salt prior to iodide incorporation at C26. The increased reactivity of substrates **6** and **9** is consistent with the alternative (path a) sequence conjectured in Fig. 4. Complexation by Ph₃P+I of the equatorial lone pair¹⁸ in **6** (20β-Me) to activate the O-C26 bond, and backside attack by iodide at C26 of the activated spiroketal from **9** (axial 25-Me), should each be more facile than for **3** based on steric considerations. Unimolecular elimination of the elements of Ph₃PO and HI from an intermediate phosphorane **20** is consistent with the lack of effect of steric demand or strength of bases employed.

Intramolecular alkylation at C20 of 4, 13-16 provided C22 ketones 7, 17-19 as single diastereomers. Thus, alkylation likely occurs from the convex side of the *intact* D/E [3.3.0] system. Electrophilic assistance for the alkylation is implied by the failure to produce 7 from 4d in the absence of the $Ph_3P^+X-X^-$ reagent. No products of the type seen when free mineral acid is present¹⁴ were noted even though production of 7 entails net elimination of a second equivalent of HI. Rupture of the E-ring may proceed after alkylation via *anti*



elimination of the oxacarbenium ion, as iodide attack at C16 followed by *syn* HI elimination seems inconsistent with the observed complete regioselectivity of alkene formation.

Immediate benefits of this new method include a more efficient, reliable and scaleable route via 4d to 5 (3 steps, 70% vs. 6 steps, 35-45%) as well the potential for rapid access to other antitumor steroids. North G (2) is now available in 10 steps (25% overall yield) from 3.

Typical Procedure. To a solution of **3** (10.00 g, 21.15 mmol), Ph₃P (11.1 g, 42.3 mmol, 2.0 eq), and 2,6-lutidine (1.9 mL, 16.3 mmol, 0.8 eq) in TCE (200 mL) was added iodine (10.75 g, 42.3 mmol, 2.0 eq) with vigorous stirring.¹⁹ After 5 min the orange solution was heated to 145 °C for 30 min and then rapidly cooled. The deep red mixture was diluted with 1:3 toluene/hexane (400 mL), washed with ice-cold sat. NaHSO₃ and then with sat. NaHCO₃, dried (Na₂SO₄), and suction filtered through a short column of silica gel. The silica plug was washed with hexane to remove residual TCE, then rapidly eluted with 3-5 volumes of 75% hexane/EtOAc. The eluent was concentrated to afford 9.21 g (76%) of **4** as offwhite low-melting solids (6:1 ratio of **4d/e** by NMR).¹³ Further elution with 65% hexane/EtOAc provided 1.60 g (17%) of diketone **7**.

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