LETTERS 2005Vol. 7, No. 13 2523 - 2526

ORGANIC

Boro-Induced ipso-Desilylation. Link to an in situ Suzuki Reaction[†]

Methodology. Halo-, Nitroso-, and

Directed ortho Metalation-Based

Zhongdong Zhao and Victor Snieckus*

Department of Chemistry, Queen's University, Kingston, ON, K7L 3N6, Canada snieckus@chem.queensu.ca

Received March 28, 2005

ABSTRACT



Treatment of DoM-derived silylated aromatics 2-4 under standard electrophilic halogenation conditions cleanly affords ipso-desilyation products 5-7, while nitration of methoxy-substituted analogues 8, 9 leads to non-ipso isomers 10, 12 and 11, 13, controlled by a silicon steric effect. Sequential ipso-borodesilylation of 2a, 3a, and 20 followed by treatment with aryl halides under Pd-catalyzed conditions constitutes an in situ Suzuki-Miyaura cross-coupling protocol to biaryls and heterobiaryls 23.

In the 30 year evolution of modern organosilicon chemistry,¹ during which its rich and broad impact on organic methodology, total synthesis, and industrial applications has been amply demonstrated, the total output of arylsilane chemistry has constituted a minor component^{1a} despite the comprehensive and careful physical organic studies of Eaborn² and others,³ which have potentially set a platform for a synthetic playground. A salient feature is marked by kinetic studies that show ipso-protodesilyation/proton-deuterium exchange

rates⁴ $k_{deSi}/k_{H,D} = 10^4$, which stimulated brief studies of other electrophile-induced desilylation reactions.^{1a,5} Our longstanding interest⁶ and that of others⁷ have stimulated an exploratory study of electrophile-induced desilylation reactions for arylsilanes derived by directed ortho metalation

(7) (a) Krizan, T. D.; Martin, J. C. J. Am. Chem. Soc. 1983, 105, 6155. (b) Taylor, S. L.; Lee, D. Y.; Martin, J. C. J. Org. Chem. 1983, 48, 4156. (c)Yamamoto, Y.; Yanagi, A. Heterocycles 1982, 19, 168. (d) Effenberger, F.; Krebs, A. J. Org. Chem. 1984, 49, 4687. (e) Hari, Y.; Shoji, Y.; Aoyama, T. Synthesis 2004, 8, 1183.

[†] Dedicated to the memory of Colin Eaborn (1923–2004) in appreciation of his insightful mechanistic work in organosilicon chemistry.

^{(1) (}a) Čolvin, E. Silicon in Organic Synthesis; Butterworth: New York, 1981; p 125. (b) Fleming, I.; Dunoguès, J.; Smithers, R. Org. React. 1989, 37, 57. (c) Brook, M. Silicon in Organic, Organometallic, and Polymer Chemistry, Wiley: New York, 2000, p 569. (d) Fleming, I. Science of *Synthesis*; Thieme: Stuttgart, 2002; Vol. 4, p 685. (2) Eaborn, C. J. Organomet. Chem. **1975**, 100, 43.

 ^{(3) (}a) Freiser, H.; Eagle, M. V.; Speier, J. J. Am. Chem. Soc. 1953, 75, 2821.
 (b) Bott, R. W.; Eaborn, C.; Greasley, P. M. J. Chem. Soc. 1964, 4804. (c) Berwin, H. J. Chem. Commun. 1972, 237. (d) Al-Omran, F.; Ridd, J. H. J. Chem. Soc., Perkin Trans. 2 1983, 1185. (e) Herrlich, M.; Hampel, N.; Mayr, H. Org. Lett. 2001, 3, 1629.

⁽⁴⁾ Eaborn, C.; Pande, K. C. J. Chem. Soc. 1960, 1566.

^{(5) (}a) Dey, K.; Eaborn, C.; Walton, D. R. M. Organomet. Chem. Synth. 1970/1971, 1, 151. (b) Felix, G.; Dunoguès, J.; Calas, R. Angew. Chem., Int. Ed. Engl. 1979, 18, 402. (c) Bennetau, B.; Dunoguès, J. Synlett 1993, 171.

^{(6) (}a) Reed, J. N. Ph.D. Thesis, University of Waterloo, Waterloo, Canada, 1984. (b) Mills, R. J.; Horvath, R.; Sibi, M. P.; Snieckus, V. Tetrahedron Lett. 1985, 26, 1145. (c) Bourguignon, M. L.; Snieckus, V. Unpublished results, University of Waterloo, Waterloo, Canada, 1987. (d) Mills, R. J.; Taylor, N. J.; Snieckus, V. J. Org. Chem. 1989, 54, 4372. (e) Beaulieu, F.; Snieckus, V. J. Org. Chem. **1994**, 59, 6508. (f) MacNeil, S. L.; Familoni, O. B.; Snieckus, V. J. Org. Chem. **2001**, 66, 3662. For an intramolecular carbodesilylation, see: Sibi, M. P.; Shankaran, K.; Alo, B. I.; Hahn, W. R.; Snieckus, V. Tetrahedron Lett. 1987, 28, 2933.

(DoM) protocols. Herein, we report preliminary studies on electrophilic halo-, nitroso-, and boro-induced *ipso*-desily-lations (Figure 1, 1, path a), the steric effect of silicon groups





promoting non-*ipso* electrophilic substitution (particularly with $E^+ = NO_2^+$) in the presence of electron-donating groups (EDGs) (**1**, path b), and the conjunction of the borodesilylation procedure (**1**, path a, $E^+ = BX_2^+$)^{8,9} with the Suzuki cross-coupling regimen. Considered in sum, these results provide new extensions and opportunities in DoM-mediated, regioselective construction of aromatics and certain complementarity with the direct usage of arylboronic acids for the synthesis of biaryls and heterobiaryls.¹⁰

Results of halo-induced *ipso*-desilylation studies using I⁺-, Br⁺-, and Cl⁺-generating¹¹ reagents for three prototypical, powerful, and commonly used directed metalation group (DMG)-bearing aromatics 2-4 are summarized in Table 1. All substrates show highly regioselective *ipso* reactivity to give products 5-7. In sharp contrast, all of the corresponding nonsilylated derivatives show no reaction, leading to recovery of starting material. As expected, the para-directing effect of the moderate *O*-carbamate EDG is evident at least in iodination and bromination reactions but at higher temperatures.¹²

To obtain initial appreciation of *ipso*-desilylative NO_2^+ reactivity, a series of unsubstituted (**8a**, **9a**), TMS-substituted (**8b**, **9b**) and TIPS-substituted (**8c**, **9c**) benzamides and *O*-carbamates containing the powerful OMe EDG were tested under standard electrophilic nitration conditions.^{12,13} As gleaned from Table 2, instead of *ipso*-desilylation, electrophilic nitration occurred, assumably dictated by the OMe

(10) For recent reports of electrophile-induced *ipso*-deboronation studies, see:
(a) Salzbrunn, S.; Simon, J.; Prakash, G. K. S.; Petasis, N. A.; Olah, G. A. *Synlett* **2000**, *10*, 1485. (b) Prakash, G. K. S.; Panja, C.; Mathew, V. S.; Petasis, N. A.; Olah, G. A. *Org. Lett.* **2004**, *6*, 2205.

(11) Chlorination using chlorine was not convenient due to an inability to control amounts of chlorine added, resulting in the formation of some dichlorination (15%) and side products.

(12) For details, see General Procedure in Supporting Information.

(13) (a) Eaborn, C.; Salih, Z. S.; Walton, D. R. M. J. Chem. Soc., Perkin Trans. 2 **1972**, 172. (b) Dwyer, C. L.; Holzapfel, W. Tetrahedron, **1998**, 54, 7843. Table 1. ipso-Halodesilylation Reactions of Compounds 2-4

$R \xrightarrow{[i]}{U} \xrightarrow{DMG} \xrightarrow{Hal^{+a}} R \xrightarrow{[i]}{U} \xrightarrow{DMG} \xrightarrow{Hal}$ 2, 3, 4 5, 6, 7										
2, 5:	DMG =	OCONEt ₂ ; 3, 6: DMG =	CONEt ₂ ; 4	, 7: D	MG = SC	yield				
compd	R	hal ⁺ /solvent/temp	product	hal	R	%				
2a	н	ICl/CH ₂ Cl ₂ /rt	5a	Ι	н	86				
2a	н	Br ₂ /CH ₂ Cl ₂ /0 °C-rt	$\mathbf{5b}$	\mathbf{Br}	Н	$92 (82)^b$				
2b	$4-NO_2$	Br ₂ /CH ₂ Cl ₂ /mw ^c	5c	\mathbf{Br}	$4-NO_2$	70				
2a	н	NCS/MeCN/reflux	5d	Cl	Н	70				
3a	н	ICl/CH ₂ Cl ₂ /rt	6a	Ι	Н	71				
3a	н	Br ₂ /CH ₂ Cl ₂ /reflux	6b	\mathbf{Br}	Н	78				
3a	н	$NCS/MeCN/mw^d$	6c	Cl	Н	65				
4a	Н	ICl/CH ₂ Cl ₂ /rt	7a	Ι	Η	66				
4b	4-Me	ICl/CH ₂ Cl ₂ /rt	7b	Ι	4-Me	76				
4a	Н	Br ₂ /CH ₂ Cl ₂ /reflux	7c	\mathbf{Br}	Η	77				
4b	4-Me	Br ₂ /CH ₂ Cl ₂ /reflux	7d	\mathbf{Br}	4-Me	76^e				
4c	5-Me	Br ₂ /CH ₂ Cl ₂ /reflux	7e	\mathbf{Br}	5-Me	53				
4a	Н	$NCS/MeCN/mw^d$	7f	Cl	Н	NR				
^{<i>a</i>} Hal 150 W	^{<i>a</i>} Hal ⁺ = 1.5–5 equiv, $t = 3-15$ h. ^{<i>b</i>} NBS/MeCN/reflux. ^{<i>c</i>} Microwave: 150 W/25' min. ^{<i>d</i>} On SiO ₂ /250 W/10' min. ^{<i>e</i>} See ref 6f.									

group. Synthetically useful trends in nitration selectivity were observed in the series **8b,c**, **9b,c** to **10b,c**, **12b,c** conversions

OMe DMG R	HNO ₃ / 0 °C /	['] Ac ₂ O 1 h	OM - O ₂ N 3	e DMG + R	OMe DMG 5 NO ₂			
8, 9			10, 1	2	11, 13			
DMG	R		C3:C5		yield (%)			
CONEt_2	Н	8a	1:1	10a:11a	76^a			
	TMS	8b	4:1	10b:11b	95^a			
	TIPS	8c	19:1	10c:11c	83^b			
$OCONEt_2$	Н	9a	2:1	12a:13a	76			
	TMS	9b	14:1	12b:13b	82			
	TIPS	9c	15:1	12c:13c	78			
^a See ref 6c. ^b See ref 6d.								

 Table 2.
 Nitration of Benzamides 8 and O-Aryl Carbamates 9

although only a minor steric effect was noted in the change from TMS **9b** to TIPS **9c** in the *O*-carbamate series. However, a change of conditions to ammonium nitrate^{10a} on the ortho-silylated *O*-carbamate **2a** led to the formation of the corresponding 4-, 2-, and *ipso*-nitro-substituted products, **17**, **18**, and **19**, respectively, in a ratio of 7.5:1.5:1 in quantitative yield (Scheme 1). The observed selectivity may be due to the milder nitration conditions. Furthermore, when nitrosating conditions¹⁴ were adapted to the ortho-silylated *O*-carbamates **2a**, the *ipso* nitroso product **15** was obtained in modest yields in addition to recovered starting material.

^{(8) (}a) Haubold, W.; Herdtle, J.; Gollinger, W.; Einholz, W. J. Organomet. Chem. 1986, 315, 1. (b) Sharp, M. J.; Cheng, W.; Snieckus, V. Tetrahedron Lett. 1987, 28, 5093. (c) Kaufmann, D. Chem. Ber. 1987, 120, 853, 901. (d) Gross, U.; Kaufmann, D. Chem. Ber. 1987, 120, 991. (e) Farinola, G. M.; Fiandanese, V.; Mazzone, L.; Naso, F. Chem. Commun. 1995, 2523. (f) Fu, J.-m.; Snieckus, V. Can. J. Chem. 2000, 78, 227. (g) Hupe, E.; Calaza, M. I.; Knochel, P. Chem. Commun. 2002, 1390.

⁽⁹⁾ Recently, a Merck group has reported an in situ metalationelectrophile quench method that adds convenience to the synthesis of silylated and boronated indole derivatives; see: Vazquez, E.; Davies, I. W.; Payack, J. F. J. Org. Chem. **2002**, 67, 7551.



These results illustrate advantages for the synthesis of oxidized (**19**) and reduced (**16**) products as well as 1,2,3-contiguously substituted aromatics (**14** from **12b**), which are difficult to prepare by classical electrophilic chemistry.¹⁵

While attempts to date in electrophilic fluorination, cyanation, amination, and Friedel-Crafts acylation¹⁶ have been relatively unproductive, ipso-borodesilvation has led to useful results (Table 3). Thus, treatment of ortho-silvlated DMG aromatics 2a, 3a, and 20, usually cleanly obtained by DoM chemistry, with either BCl₃ or BBr₃ at ambient temperatures followed by derivatization with pinacol^{17,18} afforded good to excellent yields of the ortho-boronopinacolates 21.12 In the carboxamide series, tertiary (entry 1) and secondary amides, including the TFA-labile cumyl amide¹⁹ (entry 2), are successful reactants. 2-Silylated O-carbamates (entry 3),²⁰ a secondary sulfonamide (entry 5), and an indole (entry 6)⁹ also furnish pure, usually crystalline, boronated derivatives. Interestingly, the bis-TMS substrate (entry 4) provides the monoborodesilvlated product, a result with implications for further ipso-desilylative chemistry.

Although the boronopinacolates described in Table 3 are useful Suzuki cross-coupling partners, the development of an in situ *ipso*-borodesilyation-coupling procedure was undertaken for the potential convenient utility in medicinal chemistry and library generation programs. Thus, under mild conditions using either BCl₃ or BBr₃ interchangeably without significant change in yield, substrates **2a**, **3a**, and **20** were converted over a 2 h period into their intermediate dihaloboranes **22**,^{8c} which, when treated with aryl bromides or iodides under typical Suzuki-Miyaura coupling conditions, afforded the biaryl or heterobiaryl products **23** (Table 4).²¹

 Table 3.
 Ortho-DMG-Substituted Arylboronopinacolates 21 by

 ipso-Borodesilylation
 Image: Comparison of the substituted arylboronopinacolates 21 by



^{*a*} Yields of isolated products. ^{*b*} Contains pinacol; calculated by NMR. ^{*c*} Yields of corresponding arylboronic acids via DoM, which was used directly in the subsequent cross-coupling reactions. ^{*d*} Fu, J.-m.; Sharp, M. J.; Snieckus, V. *Tetrahedron Lett.* **1988**, 29, 5459. ^{*e*} Yields of corresponding boroxazine derivatives. ^{*f*} Sharp, M. J. M.S. Thesis, University of Waterloo, Waterloo, Canada, 1986. ^{*s*} See ref 8b. ^{*h*} Performed with 3 equiv of BBr₃ and microwave conditions (50 W/80 °C/30 min). ^{*i*} Addition of pinacol and NEt₃ at room temperature for 1.5 h instead of evaporation in a vacuum.

Thus, borono tertiary (entries 1 and 2) and secondary benzamides (entries 3 and 4) afforded clean coupling results with bromobenzene as a partner. Phenyl *O*-carbamates underwent coupling with a range of aryl bromides (entries 5 and 6) and 2-bromonaphthalene (entry 7), as well as 3-bromopyridine (entry 8), to afford synthetically useful yields of biaryl products. A secondary sulfonamide (entry 9) and the 2-TMS, *N*-carbamoyl indole (entries 10 and 11), a useful substance for C-7 DoM chemistry,²² give coupled products in good yields.

This *ipso*-borodesilylative methodology offers the following features: (a) the starting arylsilanes are stable, readily

^{(14) (}a) Birkofer, L.; Franz, M. *Chem. Ber.* **1971**, *104*, 3062. (b) Uemura, S.; Toshimitsu, A.; Okano, M. J. Chem. Soc., Perkin Trans. 1 **1978**, 1076.

⁽¹⁵⁾ For example, nitration of 2-methoxyphenol affords a 1:1 mixture of the 4- and 6-nitro derivatives; see ref 12b.

⁽¹⁶⁾ Various Lewis acid (AlCl₃)-catalyzed acylation attempts led to protodesilylation products together with recovery of starting material.

⁽¹⁷⁾ Wong, K.; Chien, Y.; Liao, Y. L.; Lin, C.; Chou, M.; Leung, M. J. Org. Chem. 2002, 67, 1041.

⁽¹⁸⁾ For the preparation of stable diethanolamine adducts of boronic acids, see ref 8b.

⁽¹⁹⁾ Metallinos, C.; Nerdinger, S.; Snieckus, V. Org. Lett. 1999, 1, 1183.(20) For a phenanthrene O-carbamate case, see ref 8e.

⁽²¹⁾ **Typical One-Pot** *ipso***-Borodesilylation**–**Cross-Coupling Procedure.** A solution of BBr₃ (1.0 M, 0.48 mL) in CH₂Cl₂ was added dropwise by syringe to a stirred solution of arylsilane (0.4 mmol) in anhydrous CH₂-Cl₂ (4 mL) at 0 °C under an argon atmosphere. The reaction mixture was stirred at room temperature for 2 h after which time TLC monitoring indicated completion of the reaction. After evaporation to dryness, bromobenzene (0.33 mmol) and Ph(PPh₃)₄ (3 mol %) were added, and the whole was degassed for 15 min. Degassed glyme (4 mL) and 2 M Na₂CO₃ solution (1 mL) were added by syringe. The mixture was then heated to reflux for 5 h, cooled to room temperature, and poured into water (10 mL), and the whole was extracted with Et₂O (3 × 10 mL). The organic layer was washed with saturated NaCl, dried (Na₂SO₄), and subjected to filtration, and the filtrate was evaporated to dryness. The residue was purified by flash column chromatography (EtOAc/hexanes, 1:5) to afford the biaryl compound.

⁽²²⁾ Hartung, C. G.; Fecher, A.; Chapell, B.; Snieckus, V. Org. Lett. 2003, 5, 1899.



^{*a*} Yields of isolated products using ArBr coupling partners. ^{*b*} Yields of isolated products using ArI coupling partners. ^{*c*} Yields obtained by direct coupling with boronic acids. ^{*d*} Fu, J.-m. Ph.D Thesis, University of Waterloo, Waterloo, Canada, 1990. ^{*e*} Performed with 1.5 mol % Pd(PPh₃)₄. ^{*f*} See ref 8b. ^{*s*} See Table 3, footnote f. ^{*h*} BBr₃ was added at -25 °C, and the reaction mixture was stirred at this temperature for 1 h.

purified (chromatography, crystallization), and obtained by clean, high-yielding DoM reactions under conditions (e.g.,

Martin conditions^{7a,b}) not requiring $-78 \, ^{\circ}C^{9}$ and hence amenable to scale-up; (b) since *ipso*-borodesilyations proceed quantitatively, as evidenced by high yields of isolated boronopinacolates, the uncertainty associated with the boron intermediates (boronic acid, half-ester acid, borinic acid, boroxane) obtained by direct treatment with B(OR)₃ electrophiles,²³ as also reflected in product yields, is eliminated; (c) in contrast to the coupling reactions of the boronic acid derivatives, those of the solutions of the intermediate dihaloboranes proceed under homogeneous conditions and involve simple evaporative rather than acidic or aqueous workup to give comparable yields of products (Table 4); and (d) although requiring an extra step for the preparation of the silanes, the *ipso*-borodesilyation reaction constitutes a general and efficient route to stable arylboronates (Table 3).

In summary, preliminary results of electrophile-induced *ipso*-desilylation chemistry have shown that (a) facile haloand nitroso-induced *ipso* reactions proceed on simple DMGbearing substrates (2–4 and 2a, respectively), (b) nitration conditions lead to non-*ipso* but substitution products 10b and 12b, which can be converted to synthetically useful 1,2,3substituted aromatics, e.g., 14, and (c) *ipso*-borodesilylation provides²⁴ a method for the convenient synthesis of pure boronopinacolates 21 (Table 3) and also serves as part of a DoM-initiated method for an in situ Suzuki cross-coupling synthetic protocol for biaryls and heterobiaryls (Table 4). Mechanistic and further preparative studies are in progress.

Acknowledgment. NSERC Canada is warmly acknowledged for consistent support of our synthetic programs. Z.Z.D. is grateful for an R.S. McLaughlin Fellowship. We thank Justin Morin for providing some starting materials.

Supporting Information Available: Experimental procedures and characterization of new compounds. This material is available free of charge via the Internet at http://pubs.acs.org.

OL0506563

⁽²³⁾ For other stable pyridyl boronates that participate in cross-coupling, see: Hodgson, P. B.; Salingue, F. H. *Tetrahedron Lett.* **2004**, *45*, 685. For a review on heterocyclic boronic acids, see: Tyrrell, E.; Brookes, P. Synthesis **2003**, *4*, 469.

⁽²⁴⁾ Fluoride-mediated carbodesilylation is another synthetically useful E⁺-induced FG transfer reaction of arylsilanes; see refs 6b and 7d and: Effenberger, F.; Spiegler, W. *Chem. Ber.* **1985**, *118*, 3900. For *ipso*-cyanodesilylation, see: (a) Bennetau, B.; Dunoguès, J.; Babin, P. *Tetrahedron* **1993**, *49*, 10843. (b) Calle, M.; Cuadrado, P.; Gonzalez-Nogal, A. M.; Valero, R. Synthesis **2001**, 1949.