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Introduction

Organosilanes, prominently with their physical and chemical properties, have significant applications in organic syntheses,¹ drug discovery,² bioactive compound preparations,³ and advanced material developments.⁴ Alkyl silanes, as one of the principal research areas among organosilanes, are used as isosteres of quaternary carbons, which have found significant applications in pharmaceutical research (Fig. 1).⁵ Therefore, it has attracted much attention on their synthesis. Hydrosilvlations of olefins using Pt, Rh, Pd, or Ru catalysts are pioneering works.6 Recently, instead of noble metal catalysts, several kinds of base metal catalysts such as Fe, Co, Ni, and Mn have been developed for this transformation, generally with elegant catalyst design.7 Other means including carbene insertion into silanes,⁸ cross-coupling with organosilicon reagents,⁹ or direct C(sp³)-H silylation¹⁰ all contribute to provide alkyl silanes.

Due to the significant progress made by transition metal catalyzed carbon-heteroatom bond formation *via* unactivated



Fig. 1 Pharmaceutical active molecules with tetraalkyl silane skeletons.

Nickel(II)-catalyzed reductive silylation of alkenyl methyl ethers for the synthesis of alkyl silanes†

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A new one pot protocol has been developed for the reductive silvlation of alkenvl methyl ethers using Et_3Si -BPin and $HSiEt_3$ with nickel(II) catalyst. Styrene type methyl ethers, multi-substituted vinyl methyl ethers, heterocycles and unconjugated vinyl ethers are all tolerated to form alkyl silanes. Mechanistic study reveals that it is a cascade of a C-O bond silvlation and vinyl double bond hydrogenation process. Internal nucleophilic substitution or oxidative addition pathways were both acceptable for C-O bond cleavage. The acquired intermediate alkenvl silanes then proceeded through an unconventional reduction process thus providing alkyl silanes.

> bond cleavage in synthetic chemistry, especially for C-O bond activation,11 deoxygenative silvlation has been applied for acquiring of multiple silicon compounds. Martin et al. reported the vital work in 2014 that Ni/Cu catalyzed C-O bond silvlation of aryl or benzyl pivalates with electron rich backbones in mild conditions.12 This methodology was also introduced to aryl esters which underwent decarbonylative silvlation via acyl C-O bond activation by Rueping's and Shi's group.13 Carbamates as appropriate substrates were also discovered using nickel or iron catalysts.¹⁴ Different from esters, cut off the C-O bond of aryl ethers are more difficult because of the high bond energy. Martin et al. developed the first nickel catalyzed silvlation of aryl methyl ether compounds with silvl borate.15 Montgomery et al. then employed aryl silvl ethers as the source of C-O bond silvlation.¹⁶ In addition, pyridine assisted silvlation of aryl 2-pyridyl ethers with silvl zinc reagent were realised to generate aryl silanes.¹⁷ Alkyl ethers like benzyl methyl ethers or allyl methyl ethers were succeed as well in nickel catalyzed C-O bond silylation with silvl borate15 or silvl magnesium reagent18. However, vinyl C-O bond silvlation was rarely reported, only two styrene type methyl ethers were tested affording alkenyl silicon compounds as the supplementary for silylation of conjugated ethers in Martin's work.¹⁵ In 2017, Studer et al. reported ring opening silvlation of benzofurans which was similar to alkenyl ethers mediated by a silvl lithium reagent (Scheme 1a).¹⁹ Whereafter, Yorimitsu et al. reported CuCl catalyzed ring opening silvlation of benzofurans with disilane that made the reaction more tolerable with functional groups (Scheme 1b). Also, an addition-elimination mechanism was proposed for C-O bond cleavage.20

> Recently, our group has been interested in the valuable conversion of alkenyl ethers employing inexpensive transition metal catalysts.²¹ Combined with the high reactivity of nickel species in C–O bond activation, Ni(acac)₂ catalyzed one pot reaction of demethoxylative silylation and olefin reduction were

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(b) Copper-catalyzed ring opening silylation with disilane (Yorimitsu's work)

$$\begin{array}{c} & & & \\ &$$

(c) Nickel-catalyzed reductive silvlation with Si-B reagent (this work)



Scheme 1 C-O bonds silvlation of benzofurans and alkenyl methyl ethers.

performed with silyl borate and hydrosilane (Scheme 1c), which offers an alternative process for late-stage functionalization of alkenyl ethers.

Results and discussion

Our investigation was started by testing the reaction of styrene methyl ether 1a with 2, a silvl borate usually applied for silvlation reactions and easily accessible in bulk quantities.²² After careful optimization, we found a cocktail of cheap and air-stable catalyst Ni(acac)₂ (10 mol%), N-heterocyclic carbene ligand IMes·HCl (10 mol%), catalytic amount of zinc powder (20 mol%), HSiEt₃ (3 equiv.) and K₃PO₄ (1.5 equiv.) in toluene at 110 °C gave the best results, affording phenethyl silane 3a as the main product in 76% isolated yield. The ratio of 3a and doublebond reserved byproduct 4a was 97:3 (Table 1, entry 1). NiI₂ instead of $Ni(acac)_2$ gave a low yield of 3a and increased the amount of 4a (Table 1, entry 2). When using other nickel catalyst like Ni(OTf)₂, obvious decrease of **3a** took place (Table 1, entry 3). IMes · HCl was crucial for this transformation, other NHC or phosphine ligands showed little or no reaction of 1a (Table 1, entries 4 and 5). The reaction could not happen without nickel catalyst (Table 1, entry 6), which negated zinc's catalytic activity. While no Zn had little effect to this transformation which could be largely owing to the same reducing property of HSiEt₃ (Table 1, entry 7). No reaction occurred when silyl borate was absent indicated that direct hydrosilylation followed by β-O elimination pathway was not possible (Table 1, entry 8). The additional base K₃PO₄ was essential and irreplaceable for this reaction. As shown in entries 9-12, no reaction was found with other sylvite, phosphate or other base indicating that a subtle balance was required for nucleophilicity and steric bulk of the base. Without HSiEt₃, the yield of 3a dropped dramatically and 4a was obtained as the main product (Table 1, entry 13). A possible reason might be concluded from this result that 4a was an intermediate product and was hydrogenated by HSiEt₃, and 4a could be partly reduced to 3a with little amount of H₂O in reaction mixture when HSiEt₃ was absent. To prove this, extra hydrogen sources such as H2O, MeOH and PhMe2SiH were added instead

Table 1 Optimization of reaction conditions^a

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	10 mol% Ni(acac)₂ 10 mol% IMes-HCl 20 mol% Zn HSiEt ₃ , KyPO₄	
	1a 2 ^{toluene, 110 °C, 24h} 3a standard conditions	4a
Entry	Variation from standard conditions	Yield of $3a^{b}$ (%)
1	None	76 (97 : 3)
2	NiI_2 instead of $Ni(acac)_2$	47(54:46)
3	$Ni(OTf)_2$ instead of $Ni(acac)_2$	23 (78:22)
4	IPr·HCl instead of IMes·HCl	n.r.
5	PCy ₃ or XantPhos instead of IMes · HCl	Trace
6	No Ni $(acac)_2$	n.r.
7	No Zn	62 (97:3)
8	No 2	n.r.
9	No K ₃ PO ₄ , 20 mol% NaO ^t Bu added	n.r.
10	K ₂ CO ₃ or KO ^t Bu instead of K ₃ PO ₄	n.r.
11	K ₂ HPO ₄ or Na ₃ PO ₄ instead of K ₃ PO ₄	n.r.
12	CsF, Na ₂ CO ₃ or Cs ₂ CO ₃ instead of K ₃ PO ₄	n.r.
13	No HSiEt ₃	26 (35:65)
14	H ₂ O instead of HSiEt ₃	35 (100 : 0)
15	MeOH instead of HSiEt ₃	48(100:0)
16	PhMe ₂ SiH instead of HSiEt ₃	56 (90 : 10)

^a Reaction conditions: **1a** (0.3 mmol), **2** (0.6 mmol), HSiEt₃ (0.9 mmol) and base (0.45 mmol) were reacted in toluene (1.5 mL) at 110 °C for 24 h under nitrogen atmosphere. ^b Isolated yield, the ratios of 3a and 4a in parentheses were determined by GC-MS. n.r. equals to no reaction.

of HSiEt₃, 4a was obtained discrepantly with relatively lower vields (Table 1, entries 14-16).

With robust conditions in hand, we turned to exam the feasible scope and limitations for our Ni(acac)₂ catalyzed reductive silvlation of alkenyl methyl ethers. Styrene type methyl ethers were first carried out for alkyl silanes preparation (Table 2). Electron-donating group like methyl, tetra-butyl and phenyl at para- or meta-position were all tolerated providing up to 93% yield of products (3b-3e). When 2-pyridyl, a strong orthodirecting group, was fixed at para-position, 85% yield of 3f acquired under standard conditions without any C-H activation products detected. We are excited to find that aryl methyl ether which was readily for silvlation through nickel catalyst in mild conditions¹⁵ showed no competition against vinyl methyl ether, yet affording 78% yield of 3g. Substituents with stronger electron donor property seemed to enhance the reactivity in this process. Some disubstituted or trisubstituted styrene methyl ethers all turned out to provide products with higher yields (3h-3k). Alkenyl methyl ether bearing benzodihydrofuran skeleton (11) also gave a good result. Fluorine-containing groups such as F(3m) or $CF_3(3n)$ were proved having no impact on this reaction. The chemical selectivity of this transformation were also demonstrated by employing substrates installing with ester or amide group (30, 3p), which were deeply explored in nickel catalyzed C-O bond13 or C-N bond23 activations. Substrates with sterically hindered o-Me and o-OMe substituents were compatible enough to afford 3q and 3r in 47% and 71% yield respectively. Several π -extended alkenyl methyl ethers underwent this process with high reactivity thus providing 90% yield of



^{*a*} Reaction conditions: 1 (0.3 mmol), 2 (0.6 mmol), HSiEt₃ (0.9 mmol) and K_3PO_4 (0.45 mmol) were reacted in toluene (1.5 mL) at 110 °C for 24 h under nitrogen atmosphere. Yields were obtained after purification through column chromatography on silica gel. ^{*b*} React for 48 h. ^{*c*} 20 mol% Ni(acac)₂, 20 mol% IMes·HCl, 40 mol% Zn, 1 (0.3 mmol), 2 (1.2 mmol), HSiEt₃ (1.8 mmol) and K_3PO_4 (0.9 mmol) were reacted in toluene (3 mL) at 110 °C for 24 h under nitrogen atmosphere.

(naphthalen-2-yl)ethyl silane (**3s**) and 85% yield of (naphthalen-1-yl)ethyl silane (**3t**). Even bulky 9-anthryl ethyl silane (**3u**) was obtained with 60% yield. To our delight, this reaction was not limited to simple disubstituted alkenyl methyl ethers, trisubstituted alkenyl methyl ethers could result in structurally diverse alkyl silanes in moderate yields (**3v**, **3w**). Besides, aryl dienyl methyl ether (**1x**) reacted as well providing **3x** through dual reductive silylation process with high reactivity.

In addition of the styrene type methyl ethers, we are interested in the transformation containing heteromatic skeletons. Because of the striking impact exhibited by heteroatoms in hydrosilylation process, alkyl silicon compounds containing heteroatoms were really hard to obtain. In our conditions, several type of heteroaryl vinyl methyl ethers were tested. As shown in Table 3, furan was tolerated leading to the preparation of **6a** in 64% yield. While **6b** was obtained in a low yield probably because the competitive side reaction took place on the C–O bond of the benzofuran ring. N-heterocycles were more reactive and corresponding silanes containing heterocyclic ring
 Table 3
 Scope of heterocycle conjugated vinyl methyl ethers^a



^{*a*} Reaction conditions: 5 (0.3 mmol), 2 (0.6 mmol), HSiEt₃ (0.9 mmol) and K_3PO_4 (0.45 mmol) were reacted in toluene (1.5 mL) at 110 °C for 24 h under nitrogen atmosphere. Yields were obtained after purification through column chromatography on silica gel. ^{*b*} React for 48 h.

such as pyridine, imidazole, pyrimidine, quinoline, indole, pyrrole, and carbazole were all acquired with good results (**6c**-**6h**, **6k**, **6l**). It was worth noting that unprotected indole products **6i** and **6j** were obtained without observably decline in yields implying the efficiency of this C–Si bond forming strategy.

In order to excavate the applicability of this transformation, some unconjugated alkenyl methyl ethers were put into reaction (Scheme 2). Disubstituted alkenyl ethers were suitable for this reaction affording **8a** and **8b** with good results. Yet **8c** and **8d** was obtained in a lower yield presumably due to the steric effect caused by trisubstituted alkenyl methyl ethers.

Subsequently, some mechanistic studies were carried out to give a closer understanding of this process. Compounds *E*-4a and *Z*-4a, acquired from known report,²⁴ were investigated under standard conditions and 3a was obtained in our



Scheme 2 Reductive silylation with alkyl substituted vinyl methyl ethers. Conditions: 7 (0.3 mmol), 2 (0.6 mmol), HSiEt₃ (0.9 mmol) and K₃PO₄ (0.45 mmol) were reacted in toluene (1.5 mL) at 110 °C for 48 h under nitrogen atmosphere.



Scheme 3 Mechanistic studies.



Fig. 2 Probable reaction pathways.

prospection with favourable results despite the original configuration of **4a**. Nevertheless, styrene failed to generate **3a** when employed under the same conditions (Scheme 3a). These all suggested that compound **4a** was probably an intermediate product and a cascade of C–O bond silylation followed with olefin reduction was the possible reaction pathways rather than the way of hydrosilylation to the potential alkenes. All the time it has been difficult to distinguish the oxidative addition mechanism from addition–elimination mechanism in transition metal catalyzed inert bond activation. We tried two substrates **7e** and **9**, derived from which intramolecular competition might exist if Ni–X elimination process would happen when reacting

with 2 under standard conditions, to test the feasibility of addition–elimination pathway for C–O bond silylation (Scheme 3b). Actually, after the reaction of 7e, no structurally dominant addition–(Ni–H elimination) product 8e' nor further reductive product 8e'' were detected. Specifically, when 9 was conducted under the same reaction conditions as 7e, only few amount of compounds 10 and 11 were detected by GC-MS while anticipated product 12 was undiscovered, which suggested that allylic C–O bond scission along with vinyl C–O bond scission occurred in this reaction. No addition–(Ni–OMe elimination) product 13 nor 14 were detected further illuminating the improbability of the Ni–O elimination pathway.

According to Martin's report,¹⁵ an internal nucleophilic substitution mechanism, supported by DFT calculations,²⁵ was possible for C–O bond silvlation. As in our process, K₃PO₄ was supposed to play an important role in the convention of 2 to either Et₃SiK or $[(Et_3Si-Bpin)_n(PO_4)]K_3$ complex that might be regarded as a silvl anion surrogate. A tendency of a discrete [Ni(IMes)-SiEt₃]K complex might be acquired when in situ generated Ni(0) species exposed to IMes·HCl, 2, and K₃PO₄. Unfortunately, isolation of the nickel complex or spectroscopic evidence for this suppositive intermediate was unavailable. Therefore, two feasible pathways were proposed in the C-O bond silvlation procedure (Fig. 2). Classic oxidative addition mechanism was shown in path a, in which Ni(0) inserted into C-O bond followed by transmetalation with 2, reductive elimination then occurred forming the key intermediate 4a. Path b exhibited a constant Ni(0) mediated nucleophilic substitution mechanism, where Ni-Si complex C acted as a nucleophile, and methoxy from 1a was substituted by SiEt₃ to provide intermediate 4a. Unconventional hydrogenation of 4a with HSiEt₃ proceeded to generate 3a in this strategy, which was commonly seen as a side reaction appeared in transition metal catalyzed hydrosilylation of alkenes.26 The byproduct siloxane instead of disilane was isolated (see ESI[†] for details) which not only excluded the continues oxidative addition path of forming SiEt₃-Ni-SiEt₃ complex but also suggested trace amount of moisture in the reaction mixture maybe have some effect for this transformation.27

Conclusions

In summary, we have provided a Ni(π) catalyzed one pot reaction of alkenyl methyl ethers with silyl borate through C–O bond silylation and olefin reduction process. This strategy is successfully implemented not only for electron-rich styrene methyl ethers and π -extended alkenyl methyl ethers, but also succeeded in generating alkyl silanes containing electrondeficient aryls, heterocycles, and unconjugated alkyl groups, that provides alternative approaches for carbon–hetero bond formation and extends the scope of unreactive C–O bond activations. Besides, a possible reaction mechanism including oxidative addition or internal nucleophilic substitution for C–O bond scission and nickel mediated double bond reduction by HSiEt₃ is fully presented. Further mechanistic studies as well as functionalization of alkenyl or alkyl C–O bond are in progress in our lab.

Author contributions

X. Q. conceived the project and analysed the data. L. Z. and H. W. performed the experiments and analysed the data. L. L. helped with the substrates synthesis and analysed experimental data. Y. L. and Y. Z. participated in the mechanism discussion. The manuscript was written by X. Q. and Y. Z. contributed to the editing.

Conflicts of interest

There are no conflicts to declare.

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