Silylation reactions on nanoporous gold via homolytic Si–H activation of silanes†

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Si–H bond activation is an important process implicated in many useful synthetic applications including silylation and transfer hydrogenation reactions. Herein we discovered homolytic activation of Si–H bonds on the surface of nanoporous gold (NPG), forming hydrogen radicals and [Au]–[Si] intermediates. By virtue of this new reactivity, we achieved highly selective mono and sequential alcoholsysis of dihydrosilane. In addition, the amphiphilic nature of the [Au]–[Si] intermediate allows for a new bis-silylation reaction of cyclic ethers. The present work showcased that the surface reactivity of nanocatalysts may provide exciting opportunity for new reaction discovery.

Introduction

Catalytic activation of Si–H bonds provides rapid construction of silicon-containing functional molecules or materials, a process collectively referred to as “silylation reactions”.1–4 This also enables transfer hydrogenation reactions where the silanes serve as the hydrogen gas equivalent.5–8 A plethora of homogeneous metal catalysts including Pd,9 Ir,10–15 Rh,16–18 Ni,19,20 Ru,21–24 Co,25 and Fe26–30 salts had been discovered for these important processes. These homogeneous catalysts often activate silanes through oxidative addition or a σ-bond metathesis pathway, giving rise to metal hydride ([M]–H) intermediates.30,22,23,27 The metal hydride is known to be highly reactive towards unsaturated bonds, thus posing considerable limitations on the substrate scope.22,23,29

There is an intriguing pattern in the past few decades that heterogeneous catalysts can display reactivity inaccessible by known homogeneous catalysts.31–33 For example, Porco and co-workers reported an elegant silver nanoparticle catalysts Diels–Alder cycladdition that was not catalysed by homogeneous Lewis acids.34 Glorius and his colleagues discovered that Pd/C showed completely different regioselectivity in the arylation of benzoazepines as compared to homogeneous Pd catalysts.35 Recently, we reported a catalytic deoxygenation reaction of aromatic epoxides on the Cu surface, affording alkenes with high cis isomer selectivity.36

We thus became curious about Si–H activation on the surface of heterogeneous metal catalysts.37–39 Importantly, the pioneering studies of Yamamoto,40 Jin,41 and Chen42 have shown that silane activation on NPG can lead to the efficient oxidation of silanes to silanols, as well as transfer semi-hydrogenation of alkynes. Herein, we disclose that NPG activates Si–H bonds through a homolytic pathway, forming hydrogen radicals and [Au]–[Si] intermediates. This new activation mode allows efficient monoalcoholsysis of dihydrosilanes in a controlled (only one Si–H is activated) and chemoselective (hydrogen radicals tolerate many functional groups) manner. By tuning the reaction conditions, the second Si–H bond activation/alcoholsysis can proceed to give unsymmetrical silaketals in a one-pot reaction. Moreover, the [Au]–[Si] intermediates could serve as both an electrophile (forming Si–O bonds) and nucleophile (forming C–Si bonds), resulting in the ring opening/bis-silylation reaction of a range of cyclic ethers.

Results and discussion

Monoalcoholsisy of di-tert-butylsilane

Our investigation started with testing a number of heterogeneous catalysts in the monoalcoholsisy of di-tert-butylsilane (A) with benzy alcohol 1a (Table 1). Commercial Pd/C is the most reactive among the tested catalysts, however achieving undesired disiloxane 3a in a high percentage. Au–Pd alloy nanoparticles and Au nanoparticles showed moderate activity but good selectivity. These observations suggested that Au is less reactive but more selective than Pd in the monoalcoholsisy. When NPG was employed, the reaction rate was significantly improved, achieving 70% conversion in 12 h (Table 1, entry 5). A synthetically useful condition was obtained at 50 °C with 92% yield of 2a in 2 h. The turnover number (TON) and the turnover frequency (TOF) of the NPG under these conditions were...
With great curiosity we sought to understand the intrinsic selectivity of NPG by probing how the NPG catalyst activates the dihydrosilanes. To this end, we employed the electron paramagnetic resonance (EPR) spectroscopy using PBN as the spin trap (Fig. 1).42 Under our model reaction conditions, a persistent signal attributed to the PBN-H adduct was observed (Fig. 1a, $\Delta H = 15.26 \text{ G}, a_{\text{H(1)}} = 7.60 \text{ G}$). A similar radical was observed in the absence of the benzyl alcohol under otherwise identical conditions (Fig. 1b, $\Delta H = 15.00 \text{ G}, a_{\text{H(1)}} = 7.28 \text{ G}$). However, no signal was detected in the absence of the di-tert-butyldisilane (Fig. 1c and d). Collectively, these results suggested that the radical originated from the di-tert-butyldisilane but not the benzyl alcohol nor the solvent toluene. Interestingly, EPR of the silane alone in the toluene also gave weak radical signals (Fig. 1e), which is consistent with the spontaneous reaction without a catalyst (Table 1, entry 7), which is similar to the control reaction without any catalyst (Table 1, entry 8), suggesting that the homogenous Au salts did not contribute to the reaction.

### EPR study

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### Scope of the NPG catalysed monoalcoholysis

Based on the observed selective homolytic Si–H activation of dihydrosilanes with NPG, we demonstrated the scope of the monoalcoholysis (Table 2). Primary (1a–1c, 1j–1o) and secondary (1d–1f) alcohols are shown to be compatible with the current method, affording the desired hydridosilyl ethers. Reactions of tertiary alcohols (1g, 1h) with di-tert-butylsilane were slow. However, using diethylsilane instead led to excellent isolated yields of the desired products (2g and 2h). Phenol was also successfully converted into silane 2i, a precursor for ortho-C–H alkenylation.47 Mono silylation of substrates bearing two hydroxyl groups could also be achieved by controlling the stoichiometry of the dihydrosilane. In the presence of 1.0 equiv. of the di-tert-butyldisilane, mono silylation of 3,5-di-tert-butylphenol gave a high yield of the desired product (2m), which might suggest that the absorption of 1m onto the surface of the NPG is faster than that of 2m. Moreover, the primary hydroxyl was selectively silylated over the secondary hydroxyl (2n), and the benzyl alcohol was selectively silylated over the phenol (2o). The two cases indicate that the primary alcohols are more reactive than secondary alcohols and phenols under the current reaction conditions.

The current method showed excellent compatibility with a range of reducible functionalities, since the hydrogen radicals are much less reactive than the metal hydride. The internal 

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**Table 1** Monoalcoholysis of di-tert-butyldisilane

| Entry | Catalyst | Conversion$^b$ | 2$^-$$a$ | 3$^-$$a$ |
|-------|----------|----------------|--------|--------|
| 1     | Pd/C     | 99%            | 60%    | 15%    |
| 2     | AuPd/C$^d$ | 79%            | 30%    | 11%    |
| 3     | AuAl$_2$O$_3$ | 25%         | 21%    | n.d.   |
| 4     | Gold foil | 13%            | 11%    | n.d.   |
| 5     | NPG      | 70%            | 66%    | n.d.   |
| 6$^e$ | NPG      | 99%            | 92%    | n.d.   |
| 7$^e$ | Au salt  | 10%            | 8%     | n.d.   |
| 8$^e$ | —        | 10%            | 8%     | n.d.   |

$^a$ Reaction conditions: 1a (1.0 mmol), t-But$_3$SiH$_2$ (A, 2.1 mmol) and catalyst (1 mol%) in toluene (5 mL) under nitrogen, 25 °C for 12 h.

$^b$ Conversion monitored by NMR on the crude material.

$^c$ Calculated to be as high as 5377 and 2689 h$^{-1}$, respectively (Table 1, entry 6). Such a high TOF and TON highlight the activity enhancement brought by the nanostructure of the NPG. In contrast, homogeneous Au salts, such as 0.5 ppm AuCl or AuCl$_3$, solutions showed a very low conversion even at 50 °C (Table 1, entry 7), which is similar to the control reaction without any catalyst (Table 1, entry 8), suggesting that the homogenous Au salts did not contribute to the reaction.

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However, when two representative hydridosilyl ethers 2a (Fig. 1g) and 2r (Fig. 1h) were submitted to the EPR experiments, no PBN-H adduct was observed. These data suggested that the activation of the Si–H bonds in the hydridosilyl ethers by the NPG was much slower. The steric and electronic differences brought by the alkox substituitions might account for the retarded Si–H activation on the NPG surface. Altogether, it is clear that the selective activation of the first Si–H bond in dihydrosilane by NPG is the origin of the observed monoalcoholysis.
alkene showed tolerance under the reaction conditions \((2\text{j})\). Surprisingly, the terminal alkyne was also intact \((2\text{k})\) despite the known reactivity of alkynes in NPG catalysis.\(^{5,4,49}\) The stability of the \(\pi\) bonds in the solvent acetone (see ESI 3.1\(^\dagger\)), the alkyne \((2\text{j})\), the terminal alkyne \((2\text{k})\), the carbamate \((2\text{p})\) and the aryl halide \((2\text{f})\) towards the NPG is noteworthy.

The reactions between the less hindered diethyilsilane (1.5 eq.) and less hindered alcohols \((1\text{p}, 1\text{q}, \text{ and } 1\text{r})\) also gave exclusive monoalcoholsy products. These observations suggested that the chemoselectivity towards monoalkoholsy is dictated by the intrinsic reactivity of the NPG catalyst, and not by the steric hindrance of the dialkysilanes or alcohols.

The overall selectivity of the NPG catalyst in the controlled monoalkoholsy of dialkysilanes is unprecedented. There are a few reported catalysts for this purpose, for example, \(\text{PdCl}_2\),\(^{50}\) \((\text{PPh})_3\)RhCl\((\text{PPh})_2\)Cl\(_2\),\(^{51}\) \text{tris}[(\text{oza}xolinyl)]boretozinc hydride,\(^{32}\) and \(\text{Ph}_3\text{PCuH}\).\(^{33}\) However, these methods encounter one or more limitations including competitive disiloxane formation, hydrosilylation of \(\pi\) bonds and hydrogenolysis of aryl halide bonds. The efficient synthesis of compound \(2\text{p}\) is a good example showing the advantage of our method, as the Hartwig group reported that the ruthenium complex \(\text{Ru(Ph}_3)_2\)Cl\(_2\) catalysed reaction suffered from hydrosilylation of the amide.\(^{12}\)

### Table 2 Monoalcoholysis of di-\text{\text{-}tert} Butyilsilane and Diethyilsilane

| R\(_2\)SiH \cdot R\(_{-}\)OH | NPG | R\(_2\)SiHOR ' \cdot H\(_2\) |
|----------------|--------|-----------------|
| \(\text{Si} \cdot \text{Bu}\) | 2a, 93% | \(\text{Si} \cdot \text{Bu}\) |
| \(\text{Si} \cdot \text{H} \cdot \text{Bu} \cdot \text{H}\) | 2b, 94% | \(\text{Si} \cdot \text{Bu} \cdot \text{H}\) |
| \(\text{Si} \cdot \text{Bu} \cdot \text{H}\) | 2c, 96% | \(\text{Si} \cdot \text{Bu} \cdot \text{H}\) |
| \(\text{Si} \cdot \text{Bu} \cdot \text{H}\) | 2d, 3h, 93% | \(\text{Si} \cdot \text{Bu} \cdot \text{H}\) |
| \(\text{Si} \cdot \text{Bu} \cdot \text{H}\) | 2e, 96% | \(\text{Si} \cdot \text{Bu} \cdot \text{H}\) |
| \(\text{Si} \cdot \text{Bu} \cdot \text{H}\) | 2f, 24 h, 95% | \(\text{Si} \cdot \text{Bu} \cdot \text{H}\) |
| \(\text{Si} \cdot \text{Bu} \cdot \text{H}\) | 2g, 12 h, 84% | \(\text{Si} \cdot \text{Bu} \cdot \text{H}\) |
| \(\text{Si} \cdot \text{Bu} \cdot \text{H}\) | 2h, 87% | \(\text{Si} \cdot \text{Bu} \cdot \text{H}\) |

\(1\) (1.0 mmol), silane (1.1 mmol) and catalyst (1 mol%) in toluene (5 mL), 50 °C for 2 h (if not indicated) under nitrogen. \(2\) (1.0 mmol) was used. \(3\) Et\(_2\)SiH\(_2\) (1.5 eq.), r.t., in CDCl\(_3\). 99% conversion and only product according to \(^1\)H NMR (see ESI 3.3).

### Syntheses of unsymmetrical silaketals

We further developed efficient one-pot syntheses of unsymmetrical silaketals via sequential Si–H activations (Fig. 2). Unsymmetrical silaketals have been widely used in temporary tethering strategies.\(^{54}\) Their classic syntheses largely depended on multistep, stoichiometric displacement reactions of chlorosilanes. The Wilcox group has reported remarkable one-pot syntheses\(^{55,56}\) through sequential alcoholsy using two different readily available catalysts: a rhodium acetate dimer for the first Si–O formation and Pd/C\(^{55}\) or Mn(CO)\(_5\)Br\(^{56}\) for the second Si–O formation. We discovered that using 1.1 equiv. diethylsilane, the first Si–H activation/alcoholsy could proceed smoothly to give the desired hydrosilyl ethers. Addition of a second alcohol 5 and then increasing the reaction temperature to 50 °C triggered the second Si–H activation/alcoholsy. The desired unsymmetrical silaketals were then obtained in good yields. Our method is complementary to the Wilcox protocol because: (1) a single catalyst is employed; (2) a less hindered, more reactive diethylsilyl linkage is used, while diisopropyl linkage was used in Wilcox reactions; (3) our system is compatible with alkene functionality without compromising the yields. Importantly, in both sets of reactions (Table 2 and Fig. 2), the NPG catalyst can be easily removed from the reaction and reused for a number of cycles (see ESI 3.3). The products in Table 2 could be obtained in analytical purity by simple evaporation, which avoided the problem that these hydrosilyl ethers often decomposed rapidly during silica gel chromatography.

### Ring-opening bis-silylation reaction of cyclic ethers

We uncovered a surprising ring-opening bis-silylation reaction of cyclic ethers (Table 3). When di-\text{\text{-}tert} Butyilsilane was treated with epoxide \(7\text{a}\) in the presence of 3 mol% NPG, an unexpected product was isolated in a 62% yield which was subsequently identified as bis-silicon \(9\text{a}\) (Table 3, entry 1). Such a ring-opening/bis-silylation reaction was successfully extended to cyclic ethers of four, five and six membered rings (entries 2, 3 and 4). We discovered that at higher temperatures, the ratio of bis-silylation/hydrosilylation products could be improved, such that the bi-silicon compounds were obtained at synthetically useful yields in the range of 70–82%. The ring strain seems to dictate the reaction rate: smaller rings react faster and demand lower reaction temperature. Interestingly, the ratio of bis-silylated products increase as the ring size becomes bigger. In the case of \(7\text{d}\), only the bis-silylation products were obtained in a 82% yield.

This represents the first examples of incorporating two silicon atoms into cyclic ethers, affording simultaneously an O-
Si and a C–Si bond. In hindsight, the direct hydrosilylation of cyclic ethers had been reported in four papers, catalysed by a homogeneous iridium complex,[57] dicobalt octacarbonyl,[58] or boron Lewis acid.[59,60] Stratakis and co-workers reported an elegant gold nanoparticle catalysed silaboration of oxetanes and unactivated epoxides. This interesting formal addition of Si–B was proposed to proceed through the oxidative insertion of Au into Si–B bonds, which is different than our radical pathway (vide infra).[61] Others examples were limited to the ring-opening bromo- or iodo-silylation with Br- or I-silanes[62] or its synthetic equivalents.[63] Notably, our NPG catalysis provides a different bis-functionalization method, affording bis-silicons that could serve as useful building blocks in organic syntheses and materials chemistry. For example, the chemoselective O-Si cleavage of the bis-silicon 9c was successfully achieved under TBAF conditions, affording a terminal alcohol synthon (see ESI 5.1†).

The versatile reactivity of the remaining silane bond also provides ample synthetic opportunities, including the preparation of silicone monomers.

To further probe the reaction scope as well as to shed light on the mechanism of the reaction, substituted cyclic ethers were then studied (Table 4). Overall, the reactivity of the substituted cyclic ethers (mainly hydrosilylation) is very different than that of the unsubstituted ones (mainly bis-silylation). Three conspicuous trends could be summarized from the data: (1) the incorporation of a substituent on the carbon next to the oxygen atom shut down the bis-silylation (7e–7l), while a substituent further away resulted in bis-silylation (cf. 7l and 7m). The hydrosilylation of 7g, 7h, and 7i is not what we intended, but the yields are high. The bis-silylation of 7m and 7n is very efficient. (2) The ring opening took place preferentially at the more substituted carbon (cf. 7e, 7f, and 7l), with the only exception in the case of 7n (vide infra); (3) when a quaternary carbon (7k) or a carbon connected with phenyl groups (7f, 7j and 7l) is present in the substrate, elimination products were observed. The elimination products (i.e., 10) alerted us to a possible elimination–hydrosilylation pathway. To this end, an independently synthesized (but-3-enyloxy)di-tert-butylsilane

| Table 3 | Ring-opening/silylation of cyclic ethers<sup>a</sup> |
| --- | --- |
| Cyclic ether + t-Bu2SiH2 → NPG Products <sup>8 & 9</sup> |  |
| **Entry** | **7** | **Condition** | **Products, yields** |
| 1 |  | 0–50 °C, 0.5 h | 7a, 8a, 15% 9a, 70% |
| 2<sup>′</sup> |  | 50 °C, 3 h | 7b, 8b, 10% 9b, 77% |
| 3 |  | 90 °C, 7 h | 7c, 6% 9c, 82% |
| 4 |  | 90 °C, 36 h | n.d. 9d, 82% |

<sup>a</sup> Products and isolated yields of the NPG catalysed ring opening silylation reaction. Conditions: unless otherwise specified, cyclic ether (3.0 mmol), t-Bu2SiH2 (6.6 mmol) and catalyst (3 mol%), under argon, in a sealed tube. † Toluene 1 mL was added as the solvent. n. d. = not detected.

| Table 4 | Ring-opening/silylation of substituted cyclic ethers<sup>a</sup> |
| --- | --- |
| Entry | **7** | **Products, yields** |
| 1 | 7e | 8b, 70% 10e, 23% |
| 2 | 7f | 8f, 55% 10f, 35% |
| 3 | 7g, 7h, 7l | 8g, 82%, R = Me 2e, 87%, R = -(CH3)3 8l, 85%, R = -(CH3)3 |
| 4<sup>′</sup> | 7j | 8j, 44% 10j, 42% |
| 5<sup>′</sup> | 7k | 8k, 60% |
| 6<sup>′</sup> | 7l | 8l, 80% 10l, 10% |
| 7<sup>′</sup> | 7m | 8m, 12% 9m, 77% |
| 8<sup>′</sup> | 7n | 9n, 85% |

<sup>a</sup> Products and isolated yields of the NPG catalysed ring-opening silylation reaction. Conditions: unless otherwise specified, cyclic ether (3.0 mmol), t-Bu2SiH2 (6.6 mmol) and catalyst (3 mol%), under argon, in a sealed tube, 40 °C, 4 h. † Toluene 2 mL as the solvent, 70 °C, 5 h. ‡ 90 °C, 15 h.
failed to produce the desired product 9c under the reaction conditions (see ESI†), which was consistent with the chemoselectivity shown in Table 2 (cf. 2p) and suggested that the C-Si bond formation is unlikely to proceed through an elimination–hydrosilylation sequence.

A plausible mechanism of this cyclic ether ring opening/bis-silylation reaction is proposed (Fig. 3), which could explain our experimental observations. First, the Si–H bond of silane is activated on the surface of NPG, possibly in an oxidative-insertion manner, which gave the intermediate A. The Au–H bond is more susceptible to the PBN trapping than the Au–Si bond, so that only the hydrogen radical was observed. We observed hydrogen gas formation in the reaction, which is associated with the formation of species B from species A. The formations of hydrogen gas and Au–Si species of type A were also documented by the Stratakis group in their Au/TiO2 NP catalysed cis-1,2-disilylation of alkynes.44 We believe that at higher temperature the formation of hydrogen gas was more favourable, thus the ratio of B/A becomes bigger. A radical type C–O bond cleavage of the cyclic ether can be triggered either by the intermediate A or by the intermediate B, leaving the radical on the more substituted carbon (i.e., C). We proposed this radical was instantly trapped by the NPG surface, thus forming the intermediate D. The fate of the intermediate D diverges depending on the substrate structure. When it is a highly stable benzyl radical, it undergoes elimination to give alkene products (e.g. 10f). It also could form the hydrosilylation products by a formal hydrogen shift on the surface of A, or form the bis-silylation product through a formal Si shift on the surface of B. The higher population of B at higher temperature explains the higher yields of bis-silylation products. This mechanism can also explain why most of the substituted cyclic ethers underwent hydrosilylation but not bis-silylation. The reason might lie in the steric reactivity of the catalyst; (2) the [Au]–[Si] species formally serves as both an electrophile and a nucleophile in this bis-silylation ring-opening reaction. Both aspects cannot be achieved by homogeneous catalysis.

Conclusions

In conclusion, by virtue of the NPG catalysed homolytic NPG activation of Si–H bonds, we realized three new silylation reactions: controlled monoalcoholysis of dihydrosilanes, one-pot formation of unsymmetrical silaketals and ring-opening bis-silylation of cyclic ethers. These reactions feature a readily available catalyst, operationally simple protocols and high efficiencies. Moreover, the high chemoselectivity and functional group compatibility achieved by this NPG catalysis are unprecedented in the literature. Together with the exotic amphiphilic reactivity of [Au]–[Si] intermediates, this study showcased the unique potential of heterogeneous catalysts in organic syntheses.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This work is supported by the National Key R&D Program of China (No. 2017YFA0505200) and NSFC (21625104 and 21521091).

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