INORGANIC CHEMISTRY







FRONTIERS

RESEARCH ARTICLE

View Article Online
View Journal | View Issue



Cite this: *Inorg. Chem. Front.*, 2024, **11**, 4855

Copper-driven formation of siloxanes *via* dehydrocoupling between hydrosilanes and silanols†

Martyna Markwitz, 📵 a Kacper Łyczek, 📵 a Qingqing Bu 📵 b and Krzysztof Kuciński 📵 *a

In this study, we introduce a method for directly synthesizing various siloxanes from hydrosilanes and silanols under ambient conditions. This process relies on the use of Stryker's reagent ([(PPh₃)CuH]₆) as the first copper catalyst, enabling this specific cross-dehydrogenative coupling. This method stands out for its exceptional chemoselectivity, making it a superb alternative to established catalytic systems. It operates under mild conditions yet maintains high process efficiency. Throughout the investigation, we demonstrated the potential for modifying the resulting hydrosiloxanes through alcoholysis to produce silyl ethers and *via* hydrosilylation to yield functionalized siloxanes.

Received 12th May 2024, Accepted 25th June 2024 DOI: 10.1039/d4qi01184h

rsc.li/frontiers-inorganic

Introduction

Siloxanes represent a category of compounds defined by a sequence of alternating silicon and oxygen atoms. The mentioned Si–O–Si motif is widely encountered in nature, although the synthetically derived combinations revolutionized industry. This was driven by the versatility of siloxanes, which are used, among other things, as sealants, adhesives, coatings, plastics, cosmetics, *etc.* In synthetic chemistry, siloxanes and silyl ethers also serve as very important reagents and building blocks. The mention of the synthetic chemistry, siloxanes and silyl ethers also serve as very important reagents and building blocks.

Regardless of their structure, chain length, or shape, the main method leading to the formation of Si-O-Si sequences is associated with the process of chlorosilane hydrolysis (Fig. 1a).² Despite many significant advantages, mainly related to the wide availability of substrates, these methods are accompanied by numerous limitations. This becomes particularly critical in instances where the synthesis of unsymmetrical compounds or complex structures with particular functional groups is envisaged. This, in turn, led scientists to begin searching for alternative synthetic methods replacing chlorosilanes, and using silanols in combination with an appropriate silylating agent.^{2,23,24} Consequently, a series of methods based on coupling reactions were developed, where instead of a corrosive byproduct such as HCl, dihydrogen (dehydrogenative

coupling),^{25–32} alkene (dealkenative coupling),^{33,34} or alkyne³⁵ (dealkynative coupling) is obtained. Additionally, the possibility of functionalizing existing siloxanes, for example through hydrosilylation of tetramethyldisiloxane, has also been demonstrated.^{36–39} Among the aforementioned methodologies, the approach based on the dehydrocoupling reaction between silanols and hydrosilanes appears to be the most extensively explored, particularly concerning the potential util-

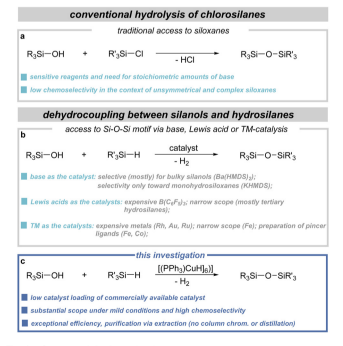


Fig. 1 Context of the investigation.

and spectra. See DOI: https://doi.org/10.1039/d4qi01184h

^aFaculty of Chemistry, Adam Mickiewicz University, Poznan, Uniwersytetu
Poznanskiego St. 8, 61-614 Poznan, Poland. E-mail: kucinski.k@amu.edu.pl

^bSchool of Chemistry and Chemical Engineering/State Key Laboratory Incubation
Base for Green Processing of Chemical Engineering, Shihezi University, North Fourth
Road, 832003 Shihezi, Xinjiang, People's Republic of China

† Electronic supplementary information (ESI) available: Characterization data

ization of the released dihydrogen (Fig. 1b). Among the strategies for selective SiO-H/Si-H dehydrocoupling, approaches involving the utilization of strong bases, 27,28,31 Lewis acids, 40-47 4d/5d metal catalysts, 25,48 and increasingly prevalent 3d metal complexes have emerged. 26,30,32 Each type of catalyst has its own advantages and disadvantages. One of the main limitations of base-catalyzed protocols is the challenge of achieving desired chemoselectivity. When using Lewis acids, we are restricted to the costly tris(pentafluorophenyl) borane and primarily the use of tertiary silanes. Catalysts based on noble metals are very expensive, so the focus is primarily on utilizing 3d metal complexes. Of notable interest is the employment of cobalt complexes featuring NNN-type³⁰ and PNP-type³² pincer ligands. However, the preparation of the ligands themselves can be considered a drawback. Copper compounds have shown effectiveness in synthesizing silyl ethers, ⁴⁹⁻⁵⁴ yet they have not been applied to the production of siloxanes. Within this context, pivotal methodologies, as relevant to this study, have been pioneered by Schubert and Lorenz, leveraging Stryker's reagent ([(PPh3)CuH]6) to obtain diverse alkoxysilanes and silanols. 54,55 Moreover, there exists a sole example where copper compounds have been applied during the synthesis of siloxanes. 29 However, in this scenario, reactions must be performed with copper(II) triflate at 80 °C, where mechanistic studies show that silvl triflate molecules, formed in situ, are the true catalytic species.

Considering the precedent research by Schubert and Lorenz,⁵⁴ as well as our expertise in forging of Si-O bonds, we embarked on an investigation into the application of commercially available (triphenylphosphine)copper hydride hexamer, known as Stryker's reagent, 56-59 within the field of siloxane synthesis (Fig. 1c). It is worth adding here that, until now, this compound and its variations with different phosphine ligands have been used as very useful (pre)catalysts in a range of synthetic processes. 60-64 Our initial hypothesis posited that this easily accessible copper cluster would serve as an exceptionally useful catalyst for the dehydrocoupling between silanols and hydrosilanes, thus circumventing the necessity for intricate pincer ligands commonly utilized in prior instances involving 3d metal complexes. Thus, this approach would yield a catalytic system based on copper, which is much more abundant in nature than cobalt. This endeavor resulted in the development of a highly efficient methodology for siloxane synthesis. Moreover, we substantiated the practical viability of our approach through scaled-up experimentation and exemplification of further product modification via subsequent alcoholysis and hydrosilylation reactions.

Results and discussion

In Table 1, we compared the initial findings from our investigation into the dehydrogenative coupling reaction involving phenylsilane (1a) and *tert*-butyldimethylsilanol (2a). These comparisons were important for optimizing the entire process. This is particularly significant in terms of chemoselectivity.

The use of primary silane may lead, depending on the reagent ratio, to the substitution of one, two, or even three silanol molecules. Theoretically, this may yield three distinct products: disiloxane, trisiloxane, and a branched variant. Hence, it is crucial to establish reaction conditions that ensure selectivity and facilitate straightforward isolation of the desired products. It should be noted that due to the sensitivity of all reagents (substrates and catalyst), they were stored in a glovebox and each reaction was set up there. Subsequently, the sealed vials were removed from the glovebox and mixed for a predetermined duration. We initiated our research by examining how the catalyst affects the progression of the dehydrocoupling reaction. Firstly, two tests were carried out (entries 1 and 2). In both tests, all reagents were mixed for 20 hours without the catalyst present. One test maintained the reaction in a closed vial (entry 1), while the second vial after being set up in the glovebox was opened briefly and exposed to air, then sealed again and mixed for the specified duration (entry 2). Both tests affirmed the necessity of a copper catalyst for the reaction to proceed, prompting us to continue our research with its inclusion. Across all experiments (entries 3-12), it became evident that a catalyst amount as low as 0.125 mol% is sufficient. Any further reduction in catalyst amount began to impact the conversion rate of the substrates. Additionally, these experiments revealed the versatility of conducting the reaction in either an argon atmosphere or in air. The presence of air only shortens the duration of the process compared to carrying out the entire reaction in an argon atmosphere (entries 5 and 6). This observation aligns with the pioneering findings of Lorenz and Schubert, who similarly noted the expedited reaction rate in the presence of air.54

Next, by conducting the reaction in toluene with 0.125 mol% of the catalyst and a reagent ratio of 1a:2a=2:1, we achieved efficient and highly selective synthesis, exclusively yielding the disiloxane product 3a (entry 9). Subsequently, we explored the possibility of substituting two molecules of silanol to obtain the corresponding trisiloxane. Once again, toluene proved to be the most effective solvent (entry 13). Interestingly, despite increasing the amount of silanol in the system (entry 14) and simultaneously increasing the catalyst amount, we did not observe the substitution of a third silanol molecule. Subsequent studies, to be discussed later, indirectly suggest that steric effects likely play a crucial role in preventing the substitution of the third molecule of this bulky silanol. Finally, we investigated the impact of simple copper salts on the progression of this reaction (entries 15 and 16). Both copper(II) chloride and copper(II) acetate demonstrated limited activity in the dehydrocoupling process, yielding only traces of the desired products (less than 3%).

With the optimized conditions in hand, we carried out experiments utilizing different hydrosilanes and silanols to showcase the wide-ranging utility of our protocol. Our reaction conditions demonstrated effectiveness across a diverse array of primary and secondary silanes in the process of mono-dehydrogenative coupling (Fig. 2). It has been shown that various silanols, including those containing bulky groups like isopro-

Table 1 Optimization studies for a copper-catalyzed dehydrogenative silylation of phenylsilane with tert-butyldimethylsilanol

$$\begin{array}{c} H_2 \\ Si-H \\ + \ H-OSiMe_2tBu \end{array} \times \begin{array}{c} x \ mol\% \ [PPh_3CuH]_6 \\ \hline solvent, \ rt, \ time \\ -H_2 \end{array} \\ \begin{array}{c} 3a \end{array} \begin{array}{c} di\text{-substitution} \\ \hline Si-OSiMe_2tBu \\ \hline 4a \\ \hline tri-substitution \\ \hline OSiMe_2tBu \\ \hline Si-OSiMe_2tBu \\ \hline OSiMe_2tBu \\ \hline OSiMe_2$$

Entry	Variation of conditions	Catalyst loading (mol%)	Molar ratio [1a]:[2a]	Conversion of $\mathbf{1a}$ or $\mathbf{2a}^{i}$ [%]; {isolated yield};	Selectivity ^j [%] [3a]:[4a]:[5a]
1	No catalyst ^a	_	1:1	0	
2	No catalyst ^b	_	1:1	0	_
3	In toluene ^c	0.25	1:1	99 (2a)	91:9:0
4	In toluene ^d	0.25	1:1	99 (2a)	60:40:0
5	In toluene ^c	0.125	1:1	99 (2a); { 3a , 80%}	90:10:0
6	In toluene ^d	0.125	1:1	99 (2a); { 3a , 79%}	92:8:0
7	In toluene ^d	0.125	1.5:1	99 (2a); { 3a , 82%}	98:2:0
8	In hexane ^e	0.125	1.5:1	99 (2a); { 3a , 81%}	97:3:0
9	In toluene ^d	0.125	2:1	99 (2a); { 3a , 87%}	99:1:0
10	In hexane ^e	0.125	2:1	99 (2a); { 3a , 84%}	99:1:0
11	In acetonitrile ^f	0.125	2:1	99 (2a); { 3a , 86%}	95:5:0
12	In 2-methyltetrahydrofuran ^g	0.125	2:1	65 (2a)	97:3:0
13	In toluene ^d	0.25	1:2	99 (1a); {4a, 92%}	0:100:0
14	In toluene ^e	0.25	1:3	99 (1a)	0:100:0
15	CuCl ₂ instead of [(PPh ₃)CuH] ₆ ^h	2	1:2	3 (1a)	0:100:0
16	$Cu(OAc)_2$ instead of $[(PPh_3)CuH]_6^h$	2	1:2	2 (1a)	0:100:0

^a All reaction conditions: **1a** (1 eq.), **2a** (1 eq.), toluene (0.25 mL), rt, under argon atmosphere, 20 h. ^b All reaction conditions: **1a** (1 eq.), **2a** (1 eq.), toluene, rt, under air atmosphere, 20 h. ^c Under air atmosphere, 30 min. ^c Under air atmosphere, 1 h. ^f Under air atmosphere, 3 h. ^g Under air atmosphere, 6 h. ^h Under air atmosphere, in toluene, 20 h. ^c Conversion of **1a** or **2a** determined by GC. ^f Selectivity of [mono-sub]: [di-sub]: [tri-sub] products determined by GC.

pyl, are reactive under the studied conditions. All of them easily undergo coupling with phenylsilane or *p*-tolylsilane to give dihydrosiloxanes (3a–3k) in very good yields (76–99%). In the case of trimethylsilanol, a significant excess of 1a was needed (10.0 eq.). Otherwise, we observed the formation of disubstituted product.

At this point, a brief comparison can be made between the previously known methods of synthesizing dihydrosiloxanes and the method described in this article. Because the authors of all discussed studies often used various reagents and reaction conditions, often focusing on different aspects, this comparison is very general in nature. However, the obtained results are highly satisfactory. Firstly, comparable numbers of products were achieved as when using Au catalyst, 25 and more than with cobalt complexes. 30,32 In the protocol where Ba (HMDS)₂ was used as a catalyst,²⁷ only bulky silanols were investigated, resulting in the production of dihydrosiloxanes. However, there is insufficient knowledge about the potential synthesis of dihydrosiloxanes from non-hindered silanols in the presence of a barium base. However, when a similar catalyst like KHMDS was used, no product containing the SiH₂ motif was observed. Even after altering the substrate ratio, the reaction did not exclusively yield dihydrosiloxanes.³¹

Following that, we advanced to examine various secondary hydrosilanes (Fig. 2). In this scenario, irrespective of the substituents' character around the silicon atom – aliphatic or aromatic – the anticipated monohydrosiloxane products (3g–3k) were consistently acquired with good yields, ranging between 80–98%.

This notable outcome sets apart our method from, for instance, the one utilizing KHMDS, which exclusively yields products of double dehydrogenative coupling.³¹ Finally, we wanted to verify whether suitable tertiary hydrosilanes would undergo a single dehydrogenative coupling reaction leading to unsymmetrical disiloxanes. The initial test using a combination of dimethylphenylsilane and trimethylsilanol as reagents resulted in the desired product 31 with a yield of 90%. This sets apart the copper-catalyzed method from those employing cobalt complexes with PNP-type ligands.³² In the latter, it was impossible to carry out the dehydrogenative coupling of any tertiary hydrosilane. However, attempts to expand the scope to include more sterically hindered silanols were unsuccessful. The main reason undoubtedly lies in steric hindrance. This is evidenced by the possibility of only attaching trimethylsilanol - the smallest of the investigated molecules.

Equipped with prior knowledge of the mono-dehydrogenative coupling reaction, we launched investigations into the subResearch Article

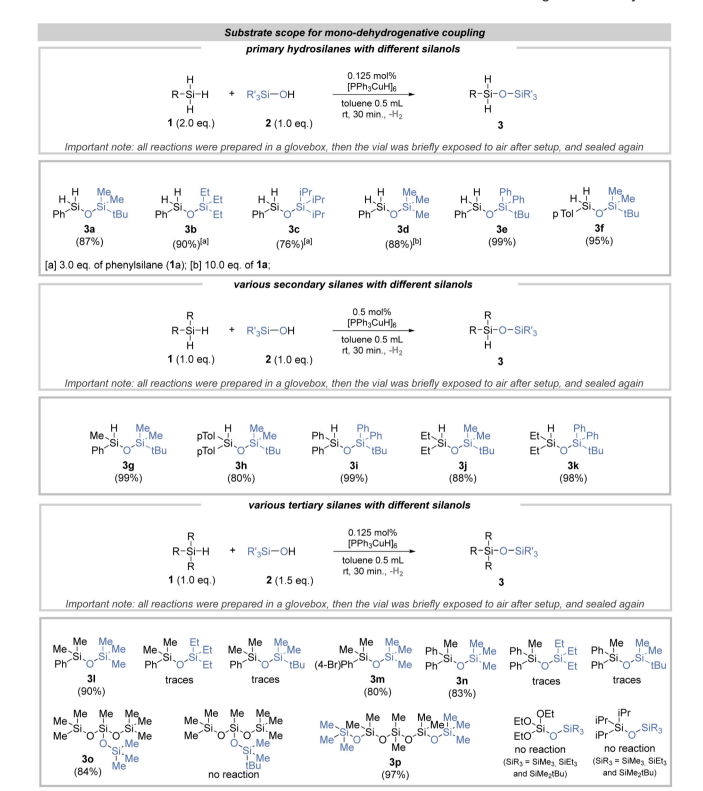


Fig. 2 Substrate scope for mono-dehydrogenative coupling reaction between various silanes and silanols.

strate's compatibility in the double dehydrogenative coupling reaction (Fig. 3). In the case of various primary hydrosilanes, the situation was quite clear, and a series of monohydrosiloxanes **4a-4i** was obtained (with yields up to 99%). Secondary

hydrosilanes displayed a distinct pattern: only reactions employing the least sterically hindered trimethylsilanol resulted in the formation of products *via* double dehydrogenative coupling (products 4j-4l). Conversely, when bulkier sila-

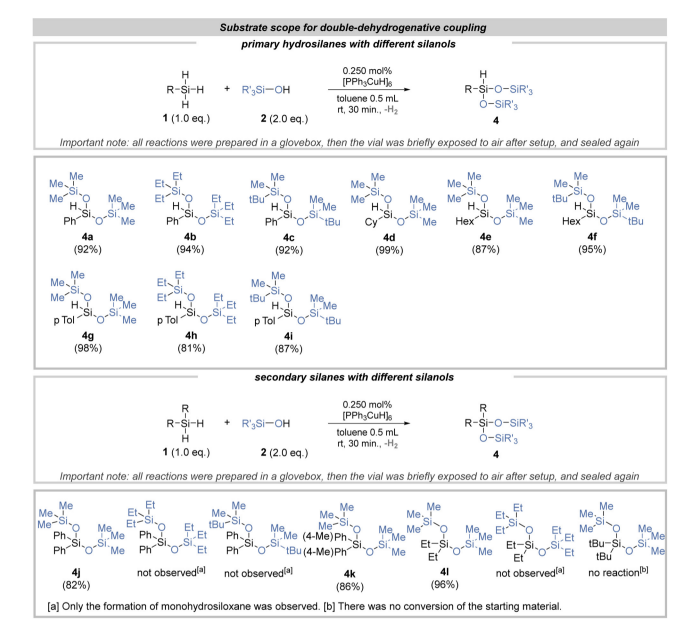


Fig. 3 Substrate scope for double-dehydrogenative coupling reaction between various silanes and silanols.

Fig. 4 Substrate scope for triple-dehydrogenative coupling reaction between aliphatic and aromatic hydrosilane and trimethylsilanol.

nols were utilized, only products of mono-dehydrogenative coupling were detected. Notably, di-tert-butylsilane stood as a unique case, failing to react with any silanol due to its inherent steric hindrance. Finally, we also investigated whether it was possible to obtain a product of triple dehydrogenative coupling in the case of primary hydrosilanes (Fig. 4).

It turned out that in combination with the least sterically hindered trimethylsilanol, branched siloxanes were successfully obtained (Fig. 4). Unfortunately, endeavors to selectively synthesize siloxanes from two distinct silanols yielded no success (Fig. 5). In pursuit of this, previous products 3a and 4c were subjected to reactions with trimethylsilanol. Upon analysis of

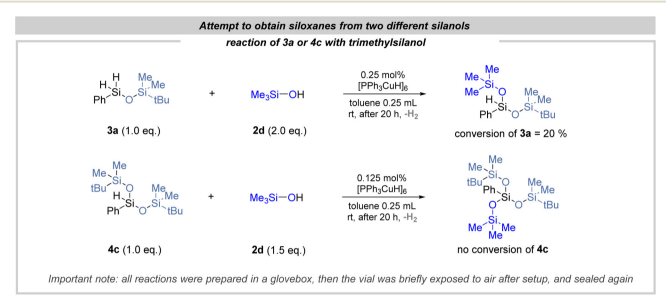


Fig. 5 Unsuccessful attempt to synthesize siloxanes from two different silanols.

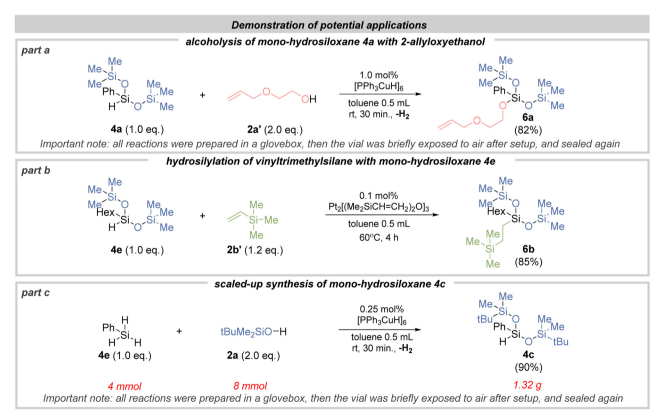


Fig. 6 Potential pathways for derivatization and scale-up of the reaction.

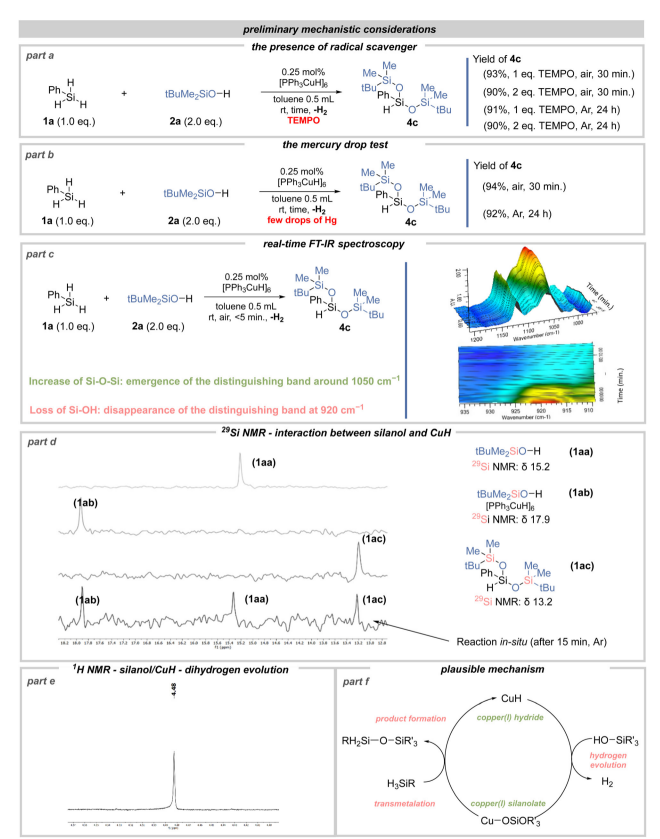


Fig. 7 Mechanistic studies.

the post-reaction mixture, the anticipated product originating from 3a was detected; however, the conversion of the starting substrates was notably low (20%). This is particularly intriguing considering the anticipation of a substantially higher conversion in this scenario.

Research Article

In the subsequent phase, our aim was to illustrate the potential for functionalizing the hydrosiloxanes we had synthesized (Fig. 6). Initially, drawing on research by Schubert and Lorenz,⁵⁴ we performed alcoholysis of the previously obtained product 4a using the same catalyst. This process yielded a silyl ether 6a (Fig. 6, part a). Following this, we focused on employing product 4e in a hydrosilylation process with platinum as the catalyst (Fig. 6, part b). Platinum-based catalysts are widely utilized and studied in this field. 65-67 The reaction of product 4e with vinylsilane resulted in the formation of product 6b with 85% yield. These reactions collectively highlight the significant application potential of the hydrosiloxanes obtained in our study. Furthermore, we successfully demonstrated the scalability of the original siloxane synthesis reaction by conducting it on an 8-fold larger scale (Fig. 6, part c).

Next, to get some mechanistic insights into copper catalysis, we carried out some of preliminary experiments (Fig. 7). Initially, we conducted tests involving the radical scavenger -TEMPO (Fig. 7, part a).32 We observed no significant changes in the siloxane synthesis process, regardless of the atmosphere (air or argon) or the amount of TEMPO used (1.0 or 2.0 eq.). This suggests that the process may not proceed via a radical mechanism. Subsequently, a mercury drop test was performed to determine the nature of the catalysis, whether homogeneous or heterogeneous (Fig. 7, part b).⁶⁸⁻⁷⁰ Once again, irrespective of the atmosphere, the reaction proceeded in the presence of mercury, indicating homogeneous catalysis.

Additionally, Stryker's reagent, characterized by its brickred color and utilized in all reactions, exhibited excellent solubility in solvents like benzene or toluene. Analysis of its solution in benzene-d₆ via ¹H NMR spectroscopy confirmed the existence of a copper-hydrogen bond (broad signal at 3.51 ppm).⁷¹ Then, we employed real-time in situ FT-IR spectroscopy to observe the reaction system. The kinetic profiles derived from the silvlation of phenylsilane with tert-butyldimethylsilane at room temperature substantiated a swift vanishing (within less than 5 minutes) of the distinctive peak at 920 cm⁻¹ attributed to the Si-OH functionalities, accompanied by a concomitant increase in the intensity of the Si-O-Si band at 1050 cm⁻¹ (Fig. 7, part c). Moreover, the instrumentation indicated the exothermic nature of the reaction. Then, an attempt was made to verify the catalytic system using NMR spectroscopy. The behaviour of phenylsilane and tert-butyldimethylsilanol in the presence of Stryker's reagent was monitored using both ¹H and ²⁹Si NMR (Fig. 7, part d and e). The visible change was observed for the silicon atom of the tertbutyldimethylsilanol molecule. Under normal conditions, the signal from this silicon atom appears at 15.2 ppm. Investigating the same molecule but in the presence of a copper complex showed a shift in the signal to 17.9 ppm. Both

of these signals appeared in the spectrum recorded after 15 minutes of the reaction between tert-butyldimethylsilanol and phenylsilane, with an observed signal from the reaction product 4c also at 13.2 ppm (Fig. 7, part d). Furthermore, using ¹H NMR, the evolution of hydrogen was demonstrated (signal at 4.48 ppm; Fig. 7, part e), which was also observed visually in the case of each reaction (formation of gas bubbles, particularly vigorous upon opening the vial to the air). Given that the NMR analysis of the phenylsilane/catalyst mixture revealed no discernible changes in the spectrum, in contrast to the silanol/catalyst system, and drawing from prior literature indicating that copper hydrides typically undergo protonolysis in the presence of alcohols or weak acids to produce H2 gas, 49,72,73 we propose a plausible mechanism outlined in Fig. 7 (part f). In the initial step, Stryker's reagent goes through protonolysis, which forms a copper-silanolate complex and releases hydrogen. Subsequently, the copper complex undergoes transmetalation, resulting in product formation and the regeneration of copper(1) hydride. It has been indirectly demonstrated that the silanolate form of the complex and its size play a crucial role in driving the reaction forward. Hence, when using the smallest silanol, trimethylsilanol, even trisubstitution reactions were achievable. However, even a minor alteration to triethylsilanol promptly stops the reaction at the disubstitution stage.

Finally, the acceleration of the reaction in the presence of oxygen (air), compared to under argon, is not well understood and requires further experimentation. Schubert⁵⁴ and Riant⁷⁴ observed the same effect with copper hydrides, presenting a challenging and interesting problem to solve.

Conclusions

In summary, we have documented the first copper-catalyzed cross-dehydrogenative coupling between hydrosilanes and silanols under remarkably gentle conditions (room temperature), without the need for additional activators or specialized pincer ligands. In this study, commercially available Stryker's reagent proved to be an effective catalyst with just 0.125 mol% loading. This resulted in a catalytic system that is not only straightforward and easily accessible but also significantly more competitive compared to previously used - more expensive gold or cobalt-based systems. Consequently, a range of products including unsymmetrical disiloxanes, symmetrical trisiloxanes, and branched siloxanes were synthesized. This also encompasses hydrosiloxanes containing free Si-H groups, which offer further functionalization potential, as demonstrated through alcoholysis and hydrosilylation processes. Moreover, the employed catalytic system allows for scalability, enhancing the practicality of this approach. Mechanistic investigations highlight the homogeneous nature of the catalysis and the absence of free radicals. Consequently, a proposed mechanism revolves around the formation of a copper-silanolate complex followed by its transmetalation with hydrosilanes.

Data availability

The ESI contains general information, synthetic procedures, characterization data and spectra of the obtained products.

The authors have cited additional references within the ESI.† $^{75\text{--}86}$

Author contributions

Methodology, K. K.; Synthesis of products – K. K., M. M., K. L.; Formal analysis, K. K., M. M., K. Ł.; writing—original draft preparation, K. K. and Q. B.; visualization, K. K. and Q. B.; supervision, K. K.; funding acquisition, K. K. All authors reviewed the manuscript.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This work was supported by a National Science Centre Grant UMO-2021/43/D/ST4/00132.

References

- 1 T. Köhler, A. Gutacker and E. Mejía, Industrial synthesis of reactive silicones: reaction mechanisms and processes, *Org. Chem. Front.*, 2020, 7, 4108–4120.
- 2 K. Kuciński, H. Stachowiak-Dłużyńska and G. Hreczycho, Catalytic silylation of O-nucleophiles via Si-H or Si-C bond cleavage: A route to silyl ethers, silanols and siloxanes, Coord. Chem. Rev., 2022, 459, 214456.
- 3 M. Selvan T. and T. Mondal, Radiation curable polysiloxane: synthesis to applications, *Soft Matter*, 2021, **17**, 6284–6297.
- 4 J. Hou, J. Sun and Q. Fang, Recent Advance in Low-Dielectric-Constant Organosilicon Polymers, *Chin. J. Chem.*, 2023, 41, 2371–2381.
- 5 F. Ganachaud and S. Boileau, *in Handbook of Ring-Opening Polymerization*, John Wiley & Sons, Ltd, 2009, pp. 65–95.
- 6 M. Woźnica, M. Sobiech and P. Luliński, A Fusion of Molecular Imprinting Technology and Siloxane Chemistry: A Way to Advanced Hybrid Nanomaterials, *Nanomaterials*, 2023, 13, 248.
- 7 N. A. Chekina, V. N. Pavlyuchenko, V. F. Danilichev, N. A. Ushakov, S. A. Novikov and S. S. Ivanchev, A new polymeric silicone hydrogel for medical applications: synthesis and properties, *Polym. Adv. Technol.*, 2006, 17, 872–877.
- 8 A. C. Marmo and M. A. Grunlan, Biomedical Silicones: Leveraging Additive Strategies to Propel Modern Utility, *ACS Macro Lett.*, 2023, **12**, 172–182.

- 9 X. Zhang, J. Liu, R. Qu, H. Suo, Z. Xin and Y. Qin, Synthesis and characterization of siloxane functionalized CO₂-based polycarbonate, *Polymer*, 2023, 266, 125623.
- 10 S. Grade, X. Zhang, C.-H. Yang, I. Oduro and C. Wang, Cross-linking lignin and cellulose with polymers using siloxane compounds, Front. Mater., 2024, 11, 1355976.
- 11 L. Lu, F. Dong, X. Chen, T. Guo, J. Qian, X. Xu, Y. Liu, L. Ma, L. Pang, R. Chen, P. Wang and X. Tang, Preparation and properties of an antimicrobial silane-modified polyether sealant, *Polym. Bull.*, 2023, **80**, 13129–13141.
- 12 F. de Buyl, V. Hayez, B. Harkness, J. Kimberlain and N. Shephard, in *Advances in Structural Adhesive Bonding*, ed. D. A. Dillard, Woodhead Publishing, 2nd edn, 2023, pp. 179–219.
- E. V. Ivanova, E. O. Minyaylo, M. N. Temnikov,
 L. G. Mukhtorov and Yu. M. Atroshchenko, Silicones in Cosmetics, *Polym. Sci., Ser. B*, 2023, 65, 578–594.
- 14 A. S. Madsen, H. M. E. Kristensen, G. Lanz and C. A. Olsen, The Effect of Various Zinc Binding Groups on Inhibition of Histone Deacetylases 1–11, *ChemMedChem*, 2014, 9, 614–626.
- 15 P. Nareddy, F. Jordan and M. Szostak, Ruthenium(II)-catalyzed ortho-C–H arylation of diverse N-heterocycles with aryl silanes by exploiting solvent-controlled N-coordination, *Org. Biomol. Chem.*, 2017, **15**, 4783–4788.
- 16 P. Nareddy, F. Jordan and M. Szostak, Ruthenium(II)-Catalyzed Direct C-H Arylation of Indoles with Arylsilanes in Water, *Org. Lett.*, 2018, **20**, 341–344.
- 17 J. Zhang, Y. Hou, Y. Ma and M. Szostak, Synthesis of Amides by Mild Palladium-Catalyzed Aminocarbonylation of Arylsilanes with Amines Enabled by Copper(II) Fluoride, *J. Org. Chem.*, 2019, **84**, 338–345.
- 18 I. K. Goncharova, E. A. Ulianova, R. A. Novikov, A. D. Volodin, A. A. Korlyukov and A. V. Arzumanyan, Siloxane-containing derivatives of benzoic acid: chemical transformation of the carboxyl group, *New J. Chem.*, 2022, 46, 18041–18047.
- 19 T. Sokolnicki, A. Franczyk, R. Kozak and J. Walkowiak, Coupling agents with 2,4,6,8-tetramethylcyclotetrasiloxane core – synthesis and application in styrene-butadiene rubber production, *Inorg. Chem. Front.*, 2023, 10, 5897– 5907.
- 20 A. Shimojima and K. Kuroda, Alkoxy- and Silanol-Functionalized Cage-Type Oligosiloxanes as Molecular Building Blocks to Construct Nanoporous Materials, Molecules, 2020, 25, 524.
- 21 M. Kikuchi, T. Hayashi, T. Matsuno, K. Kuroda and A. Shimojima, Direct cross-linking of silyl-functionalized cage siloxanes via nonhydrolytic siloxane bond formation for preparing nanoporous materials, *Dalton Trans.*, 2024, 53, 6256–6263.
- 22 T. Hayashi, M. Kikuchi, N. Murase, T. Matsuno, N. Sugimura, K. Kuroda and A. Shimojima, Hexagonal Prismatic Siloxanes Functionalized with Organosilyl Groups as Building Blocks of Nanoporous Materials, *Chem. Eur. J.*, 2024, 30, e202304080.

- 23 K. Kuciński and G. Hreczycho, A Highly Effective Route to Si-O-Si Moieties through O-Silylation of Silanols and Polyhedral Oligomeric Silsesquioxane Silanols with Disilazanes, ChemSusChem, 2019, 12, 1043-1048.
- 24 B. Yang, P. Guo, X. He and K.-Y. Ye, Cobalt-catalyzed dehydrative approach for the synthesis of unsymmetric disiloxanes and polysiloxanes, Org. Chem. Front., 2024, DOI: 10.1039/D4QQ00706A.
- 25 Y. Satoh, M. Igarashi, K. Sato and S. Shimada, Highly Selective Synthesis of Hydrosiloxanes by Au-Catalyzed Dehydrogenative Cross-Coupling Reaction of Silanols with Hydrosilanes, ACS Catal., 2017, 7, 1836-1840.
- 26 T. Takeshita, K. Sato and Y. Nakajima, Selective hydrosiloxane synthesis via dehydrogenative coupling of silanols with hydrosilanes catalysed by Fe complexes bearing a tetradentate PNNP ligand, Dalton Trans., 2018, 47, 17004-17010.
- 27 E. L. Coz, S. Kahlal, J.-Y. Saillard, T. Roisnel, V. Dorcet, I.-F. Carpentier and Y. Sarazin, Barium Siloxides and Catalysed Formation of Si-O-Si' Motifs, Chem. - Eur. J., 2019, 25, 13509-13513.
- 28 J. Kaźmierczak and G. Hreczycho, Highly effective functionalization of silsesquioxanes mediated by inexpensive earth-abundant metal catalyst - Potassium tertbutoxide, J. Catal., 2019, 378, 90-96.
- 29 J. Kaźmierczak and G. Hreczycho, Copper(II) triflatemediated synthesis of functionalized silsesquioxanes via dehydrogenative coupling of POSS silanols with hydrosilanes, Dalton Trans., 2019, 48, 6341-6346.
- 30 S. Pattanaik and C. Gunanathan, Cobalt-Catalyzed Selective Synthesis of Disiloxanes and Hydrodisiloxanes, ACS Catal., 2019, 9, 5552-5561.
- 31 K. Kuciński, H. Stachowiak and G. Hreczycho, Silylation of silanols with hydrosilanes via main-group catalysis: the synthesis of unsymmetrical siloxanes and hydrosiloxanes, Inorg. Chem. Front., 2020, 7, 4190-4196.
- 32 E. Szafoni, K. Kuciński and G. Hreczycho, Cobalt-catalyzed dehydrogenative cross-coupling reaction: Selective access to dihydrosiloxanes, hydrosiloxanes and functionalized silsesquioxanes, J. Catal., 2023, 423, 1-9.
- 33 Y.-R. Yeon, Y. J. Park, J.-S. Lee, J.-W. Park, S.-G. Kang and C.-H. Jun, Sc(OTf)₃-Mediated Silylation of Hydroxy Functional Groups on a Solid Surface: A Catalytic Grafting Method Operating at Room Temperature, Angew. Chem., Int. Ed., 2008, 47, 109-112.
- 34 B. Marciniec, P. Pawluć, G. Hreczycho, A. Macina and M. Madalska, Silylation of silanols with vinylsilanes catalyzed by a ruthenium complex, Tetrahedron Lett., 2008, 49, 1310-1313.
- 35 K. Kuciński, H. Stachowiak and G. Hreczycho, Silylation of Alcohols, Phenols, and Silanols with Alkynylsilanes-an Efficient Route to Silyl Ethers and Unsymmetrical Siloxanes, Eur. J. Org. Chem., 2020, 4042-4049.
- 36 R. Januszewski, I. Kownacki, H. Maciejewski and B. Marciniec, An efficient catalytic and solvent-free method for the synthesis of mono-organofunctionalized 1,1,3,3-

- tetramethyldisiloxane derivatives, I. Organomet. Chem., 2017, 846, 263-268.
- 37 R. Januszewski, M. Grzelak, B. Orwat, M. Dutkiewicz and I. Kownacki, Simple catalytic approach to highly regioselective synthesis of monofunctionalized disiloxanes decorated with metalloids, *J. Catal.*, 2020, 390, 103-108.
- 38 J. Szyling, R. Januszewski, K. Jankowska, J. Walkowiak, I. Kownacki and A. Franczyk, Synthesis of bifunctional disiloxanes via subsequent hydrosilylation of alkenes and alkynes, Chem. Commun., 2021, 57, 4504-4507.
- 39 J. Szyling, J. Walkowiak, A. Czapik and A. Franczyk, Synthesis of unsymmetrically and symmetrically functionalized disiloxanes via subsequent hydrosilylation of C≡C bonds, Sci. Rep., 2023, 13, 10244.
- Shinke, T. 40 S. Tsuchimoto and Y. Kawakami, Stereochemistry in Lewis acid-catalyzed silylation of alcohols, silanols, and methoxysilanes with optically active methyl(1-naphthyl)phenylsilane, Silicon Chem., 2007, 3, 243-249.
- 41 K. M. Rabanzo-Castillo, M. Hanif, T. Söhnel and E. M. Leitao, Synthesis, characterisation and electronic properties of naphthalene bridged disilanes, Dalton Trans., 2019, 48, 13971-13980.
- 42 J. Peng, Y. Bai and J. Li, Piers-Rubinsztajn Reaction and the Application in Siloxane/Polysiloxane Chemistry, Lett. Org. Chem., 2019, 16, 525-530.
- 43 K. M. Rabanzo-Castillo, V. B. Kumar, T. Söhnel and E. M. Leitao, Catalytic Synthesis of Oligosiloxanes Mediated by an Air Stable Catalyst, (C₆F₅)₃B(OH₂), Front. Chem., 2020, 8, 477.
- 44 J. Kaźmierczak, D. Lewandowski and G. Hreczycho, B (C₆F₅)₃-Catalyzed Dehydrocoupling of POSS Silanols with Hydrosilanes: A Metal-Free Strategy for Effecting Functionalization of Silsesquioxanes, Inorg. Chem., 2020, 59, 9206-9214.
- 45 A. M. Szawiola, B. H. Lessard, H. Raboui and T. P. Bender, Use of Piers-Rubinsztajn Chemistry to Access Unique and Challenging Silicon Phthalocyanines, ACS Omega, 2021, 6, 26857-26869.
- 46 H. Gao, A. Battley and E. M. Leitao, The ultimate Lewis acid catalyst: using tris(pentafluorophenyl) borane to create bespoke siloxane architectures, Chem. Commun., 2022, 58, 7451-7465.
- 47 S. Rubinsztajn, J. Chojnowski and U. Mizerska, Tris(pentafluorophenyl)borane-catalyzed Hydride Transfer Reactions in Polysiloxane Chemistry-Piers-Rubinsztajn Reaction and Related Processes, Molecules, 2023, 28, 5941.
- 48 Z. M. Michalska, Reactions of organosilicon hydrides with organosilanols catalysed by homogeneous and silica-supported rhodium (L) complexes, Transition Met. Chem., 1980, 5, 125-129.
- 49 H. Ito, A. Watanabe and S. Masaya, ersatile Dehydrogenative Alcohol Silylation Catalyzed by Cu(I)-Phosphine Complex, Org. Lett., 2005, 7, 1869–1871.
- 50 Y. Gunji, Y. Yamashita, T. Ikeno and T. Yamada, Convenient and selective preparation of mono-alkoxyphe-

- nylsilanes from phenylsilane and alcohols, *Chem. Lett.*, 2006, 35, 714–715.
- 51 G.-B. Liu, H.-Y. Zhao and T. Thiemann, Two New Catalysts for the Dehydrogenative Coupling Reaction of Carboxylic Acids with Silanes—Convenient Methods for an Atom-Economical Preparation of Silyl Esters, *Synth. Commun.*, 2007, 37, 2717–2727.
- 52 J. F. Blandez, A. Primo, A. M. Asiri, M. Alvaro and H. García, Copper nanoparticles supported on doped graphenes as catalyst for the dehydrogenative coupling of silanes and alcohols, *Angew. Chem., Int. Ed.*, 2014, 53, 12581–12586.
- 53 X. Wang, Y. Bai, X. Zhai, B. Wu and Y. Zhou, Synthesis of poly(silyl ether)s via copper-catalyzed dehydrocoupling polymerization, *Chin. Chem. Lett.*, 2022, 33, 2639–2642.
- 54 C. Lorenz and U. Schubert, An efficient catalyst for the conversion of hydrosilanes to alkoxysilanes, *Chem. Ber.*, 1995, 128, 1267–1269.
- 55 U. Schubert and C. Lorenz, Conversion of Hydrosilanes to Silanols and Silyl Esters Catalyzed by [Ph₃PCuH]₆, *Inorg. Chem.*, 1997, **36**, 1258–1259.
- 56 W. S. Mahoney, D. M. Brestensky and J. M. Stryker, elective hydride-mediated conjugate reduction of .alpha.,.beta.unsaturated carbonyl compounds using [(Ph₃P)CuH]₆, *J. Am. Chem. Soc.*, 1988, **110**, 291–293.
- 57 B. H. Lipshutz, W. Chrisman and K. Noson, Hydrosilylation of aldehydes and ketones catalyzed by [Ph₃P(CuH)]₆, *J. Organomet. Chem.*, 2001, **624**, 367–371.
- 58 D. Lee and J. Yun, Direct synthesis of Stryker's reagent from a $Cu(\pi)$ salt, *Tetrahedron Lett.*, 2005, **46**, 2037–2039.
- 59 J. F. Daeuble, J. M. Stryker, P. Chiu and W. H. Ng, in *Encyclopedia of Reagents for Organic Synthesis*, American Cancer Society, 2014, pp. 1–9.
- 60 D. H. Appella, Y. Moritani, R. Shintani, E. M. Ferreira and S. L. Buchwald, Asymmetric Conjugate Reduction of α,β-Unsaturated Esters Using a Chiral Phosphine–Copper Catalyst, J. Am. Chem. Soc., 1999, 121, 9473–9474.
- 61 R. Moser, Ž. V. Bošković, C. S. Crowe and B. H. Lipshutz, CuH-Catalyzed Enantioselective 1,2-Reductions of α,β-Unsaturated Ketones, *J. Am. Chem. Soc.*, 2010, **132**, 7852–7853.
- 62 K. Junge, B. Wendt, D. Addis, S. Zhou, S. Das and M. Beller, Copper-Catalyzed Enantioselective Hydrosilylation of Ketones by Using Monodentate Binaphthophosphepine Ligands, Chem. – Eur. J., 2010, 16, 68-73.
- 63 S. Nishino, M. Miura and K. Hirano, An umpolung-enabled copper-catalysed regioselective hydroamination approach to α-amino acids, *Chem. Sci.*, 2021, **12**, 11525–11537.
- 64 N. M. Harmon, N. R. Gehrke and D. F. Wiemer, Conjugate reduction of vinyl bisphosphonates, *Tetrahedron Lett.*, 2022, **106**, 154078.
- 65 H. Stachowiak, K. Kuciński, F. Kallmeier, R. Kempe and G. Hreczycho, Cobalt–Catalyzed Dehydrogenative C– H Silylation of Alkynylsilanes, *Chem. Eur. J.*, 2022, **28**, e202103629.

- 66 A. Walczak, H. Stachowiak, G. Kurpik, J. Kaźmierczak, G. Hreczycho and A. R. Stefankiewicz, High catalytic activity and selectivity in hydrosilylation of new Pt(II) metallosupramolecular complexes based on ambidentate ligands, J. Catal., 2019, 373, 139–146.
- 67 H. Stachowiak-Dłużyńska, M. Gruszczyński, M. Kubicki and G. Hreczycho, Pt(II) complexes bearing P,N-donor ligands as catalysts in chemoselective hydrosilylation, hydrogermylation, and hydroboration of terminal alkenes, *J. Catal.*, 2024, 433, 115494.
- 68 T. Jurado-Vázquez, E. Rosaldo, A. Arévalo and J. J. García, Levulinic Acid Hydrogenation with Homogeneous Cu(1) Catalyst, *ChemCatChem*, 2022, **14**, e202200628.
- 69 T. Tewari, R. Kumar and S. H. Chikkali, Iron–Catalyzed Magnesium–Mediated Formal Hydroformylation of Alkynes and Alkenes, *ChemCatChem*, 2023, **15**, e202201394.
- 70 D. Lewandowski and G. Hreczycho, Cobalt pincer-type complexes demonstrating unique selectivity for the hydroboration reaction of olefins under mild conditions, *Inorg. Chem. Front.*, 2023, 10, 3656–3663.
- 71 E. L. Bennett, P. J. Murphy, S. Imberti and S. F. Parker, Characterization of the Hydrides in Stryker's Reagent: $[HCu\{P(C_6H_5)_3\}]_6$, *Inorg. Chem.*, 2014, 53, 2963–2967.
- 72 C. M. Wyss, B. K. Tate, J. Bacsa, T. G. Gray and J. P. Sadighi, Bonding and reactivity of a μ-hydrido dicopper cation, *Angew. Chem., Int. Ed.*, 2013, 52, 12920–12923.
- 73 R. L. M. Bienenmann, A. J. Schanz, P. L. Ooms, M. Lutz and D. L. J. Broere, A Well–Defined Anionic Dicopper(1) Monohydride Complex that Reacts like a Cluster, *Angew. Chem., Int. Ed.*, 2022, **61**, e202202318.
- 74 N. Mostefaï, S. Sirol, J. Courmarcel and O. Riant, Air-Accelerated Enantioselective Hydrosilylation of Ketones Catalyzed by Copper(i) Fluoride-Diphosphine Complexes: Investigations of the Effects of Temperature and Ligand Structure, *Synthesis*, 2007, 1265–1271.
- 75 P. D. Prince, M. J. Bearpark, G. S. McGrady and J. W. Steed, Hypervalent hydridosilicates: synthesis, structure and hydride bridging, *Dalton Trans.*, 2007, 2, 271–282.
- 76 H. Zhang and D. Zhao, Synthesis of Silicon-Stereogenic Silanols Involving Iridium-Catalyzed Enantioselective C-H Silylation Leading to a New Ligand Scaffold, ACS Catal., 2021, 11, 10748–10753.
- 77 Y. Lin, K.-Z. Jiang, J. Cao, Z.-J. Zheng, Z. Xu, Y.-M. Cui and L.-W. Xu, Iridium–Catalyzed Intramolecular C–H Silylation of Siloxane–Tethered Arene and Hydrosilane: Facile and Catalytic Synthesis of Cyclic Siloxanes, *Adv. Synth. Catal.*, 2017, 359, 2247–2252.
- 78 K. Hirabayashi, J. Ando, J. Kawashima, Y. Nishihara, A. Mori and T. Hiyama, Novel Carbon-Carbon Bond Formation through Mizoroki-Heck Type Reaction of Silanols and Organotin Compounds, *Bull. Chem. Soc. Jpn.*, 2000, 73, 1409–1417.
- 79 K. Kuciński and G. Hreczycho, O-Metalation of silanols and POSS silanols over Amberlyst-15 catalyst: A facile route to unsymmetrical siloxanes, borasiloxanes and germasiloxanes, *Inorg. Chim. Acta*, 2019, 490, 261–266.

80 P. A. Di Giorgio, L. H. Sommer and F. C. Whitmore, Ditbutyl-di-aminoalkyl Silicates, *J. Am. Chem. Soc.*, 1949, 71, 3254–3256.

Research Article

- 81 M. Igarashi, K. Kubo, T. Matsumoto, K. Sato, W. Ando and S. Shimada, Pd/C-catalyzed cross-coupling reaction of benzyloxysilanes with halosilanes for selective synthesis of unsymmetrical siloxanes, *RSC Adv.*, 2014, 4, 19099–19102.
- 82 G. Calzaferri, R. Imhof and K. W. Törnroos, Structural and vibrational properties of the octanuclear silasesquioxane C₆H₁₃(H₇Si₈O₁₂), *J. Chem. Soc., Dalton Trans.*, 1994, 3123–3128.
- 83 Z. Li, J.-G. Liu, W.-P. Zhang and M.-H. Xu, Enantioselective Si– H Insertion of Arylvinyldiazoesters Promoted by Rhodium(i)/ Diene Complexes, Adv. Synth. Catal., 2024, 366, 2514–2518.
- 84 T. Aoki, J. Watanabe, Y. Ishimoto, E. Oikawa, Y. Hayakawa and M. Nishida, Synthesis and polymerization of p-pentamethyldisiloxanyl-α,β,β-trifluorostyrene and the oxygen permeability of the polymer, *J. Fluor. Chem.*, 1992, **59**, 285–288.
- 85 G. Engelhardt, H. Jancke, M. Mägi, T. Pehk and E. Lippmaa, Über die 1H-, 13C- und 29Si-NMR chemischen Verschiebungen einiger linearer, verzweigter und cyclischer Methylsiloxan-Verbindungen, *J. Organomet. Chem.*, 1971, 28, 293–300.
- 86 J. Ohshita, D. Hamamoto, K. Kimura and A. Kunai, Anodic polymerization of dithienosilole and electroluminescent properties of the resulting polymer, *J. Organomet. Chem.*, 2005, **690**, 3027–3032.