Silanes: Chemistry and applications

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Silane coupling agents belong to a class of organosilane compounds having at least two reactive groups of different types, bonded to the silicon atom in a molecule. One of the reactive groups (e.g., methoxy, ethoxy and silanolic hydroxy groups) reacts with various inorganic materials such as glass, metals, silica, sand and the like, to form a chemical bond with the surface of the inorganic material, while the other of the reactive groups (e.g., vinyl, epoxy, methacryl, amino and mercapto groups) is reactive with various kinds of organic materials or synthetic resins to form a chemical bond. As a result of possessing these two types of reactive groups, silanes are capable of providing chemical bonding between an organic material and an inorganic material. This unique property of silanes is utilized for the surface treatment of glass fiber products, performance improvement of fiber-reinforced plastics by the direct admixture to the synthetic resin, improvement of paints and other coating materials and adhesives, modification of surface properties of inorganic fillers, surface priming of various substrate materials, etc. Dental materials offer a continuously challenging forum for silanes and silanes will play an essential role in material development. This overview presents a description of silanes, their chemistry, properties, use and some of the main clinical experiences in dentistry. The majority of clinical results pointed to silanes playing a significant role in the adhesion process.

Key words: Organo functional silanes, methyl polysiloxane, coupling agents

INTRODUCTION

Silanes (compounds of silicon and hydrogen, of the formula \( Si_n H_{2n+2} \)) and other monomeric silicon compounds have the ability to bond inorganic materials such as glass, mineral fillers, metals and metallic oxides to organic resins. The adhesion mechanism is due to two groups in the silane structure. The \( Si(OR)_3 \) portion reacts with the inorganic reinforcement, while the organofunctional (vinyl-, amino-, epoxy-, etc.) group reacts with the resin. The coupling agent may be applied to the inorganic materials (e.g., glass fibers) as a pre-treatment and/or added to the resin.

Silanes, hybrid organic-inorganic compounds, can function as mediators and promote adhesion between dissimilar, inorganic and organic matrices through dual reactivity.[1] They are called primers, coupling agents, or sizes, depending on their function and substrates. This article aims at giving an introduction to silanes in dentistry, mainly emphasizing on chemistry and silane applications.

SILANE CHEMISTRY

The basic building block of silicon chemistry, silanes are monomeric silicon compounds with four substituent groups attached to the silicon atom. These substituent groups can be nearly any combination of nonreactive, inorganically reactive, or organically reactive groups. The basic structure of organosilanes is \( RnSi(OR)4-n \) with “R” being an alkyl, aryl or organofunctional group and “OR” being a methoxy, ethoxy, or acetoxy group.

Inorganic reactivity represents the covalent bonds formed through oxygen to the silicon atom, to form a siloxane type of bond. The alkoxy groups (-OR) can react with an inorganic substrate in both cases, forming covalent bonds between the matrices.

Organic reactivity occurs on the organic portion of the molecule and does not directly involve the silicon atom. The organic functional part can polymerize with an organic matrix. The reactivity of organic groups attached to Si is similar to organic analogues in C-chemistry.

Silicon is a major constituent of sand and rocks – very durable inorganic materials. Silicon bonds tenaciously to other inorganics such as glass and steel. Silicon will bond tenaciously to organic polymers as well when an organic group, such as aminopropyl, is attached to the silicon.[2] This is because the reactivity of organic groups attached to silicon is similar to or-
ganic analogs in carbon chemistry.

Silanes may or may not contain reactive groups. Special characteristics can be added to the silicon molecule by adding non-reactive groups, such as methyl, the higher alkyls, phenyl and trifluoropropyl. This large number of possible combinations explains silicon’s versatility and its ability to be used in a variety of ways with carbon-based chemicals. Methylsilanes are the raw materials for manufacturing silicones. They are extremely mobile, colourless, liquids that are soluble in organic solvents and in some cases, in anhydrous alcohol. Silanes have low molecular weights and are thus highly volatile. Most important are the Methylchlorosilanes, which form the basis for all silicone chemistry

Industrial silicone production has its commercial basis in the direct synthesis of methylchlorosilanes from silicon and methylchloride, via a process called the Mueller-Rochow synthesis. This technique, developed independently in 1940/41 by professors R Mueller in Germany and E G Rochow in the United States, takes place in the presence of a copper catalyst at approximately 280°C. Finely ground and well mixed Si and Cu are brought together in a fluid bed reactor with methyl chloride in gaseous form. This produces a silane mixture, from which are derived the most important organochlorosilanes i.e. dimethyl-dichlorosilane (CH₃)₂SiCl₂

280°C-300°C
Si+CH₃Cl → CH₃₂SiCl₃ methyltrichlorosilane
Cu (CH₃)₂SiCl₂ dimethyldichlorosilane
(CH₃)₂SiCl trimethylchlorosilane
+ other silanes

The chlorosilanes are distilled or purified and the primary product - dimethyldichlorosilane, (CH₃)₂SiCl₂ - is reacted with water (hydrolysis) to give poly(dimethylsiloxane) oligomers: [Me₂SiO]ₙ. The whole process is self-contained: the chloride ion is recycled with the regeneration of methyl chloride via the reaction of methanol and HCl. Thus, only readily available raw materials are used and useful products emerge from this cycle. The reaction of the monomer organo-chlorosilanes with water (hydrolysis) or methanol (methanolysis) produces silanols, which, using HCl catalysis, lead directly to further reacted oligomers or polymer siloxanes (‘sil’ represents silicon ‘ox’ stands for oxygen and ‘ane’ describes the saturated nature of the bond). The description ‘silicone’ for the whole class of polysiloxanes is an imitation of the oxygen-carbon bonds of carbon chemistry which are known as ‘ketones’. The latter however, because of their particular characteristics, form double bonds instead of single bonds.

\[
\text{(CH₃)}₂\text{SiCl}_₂ \rightarrow \text{H}_2\text{O} \quad \text{2HCl} \\
\Rightarrow \quad \text{(CH₃)}₂\text{Si(OH)}_2 \rightarrow \text{Polymer monomer}
\]

The process of conversion is called polymerization. It occurs by ring opening and/or condensation in the presence of an acid or base catalyst, to form silicone fluids. At about pH4, rate of condensation to form larger oligomers is at the minimum and silane solutions have highest stability. Acetic acid is often used for pH adjustment. During this reaction, which is commonly known as equilibration, the SiO bonds are constantly cleaved and reformed in a series of competing reactions involving both cyclic and linear species - building up molecular structure, until an equilibrium distribution is reached. Molecular weight control is obtained by the use of ‘end-blockers,’ such as hexamethyldisiloxane derivatives, which act as chain-termination agents. This is most simply represented by the equation:

\[
\text{[Me₂SiO]}ₙ + \text{Me₂SiOSiMe₃} \rightarrow \text{Me₂SiO[Me₂Si]}ₙ \text{SiMe₃ siloxane fluid}
\]

Molecular weight and many of the ultimate properties such as viscosity, are determined by the ratio of trimethyl (Me₃) to dimethyl (Me₂) siloxanes in the reaction

Mono-, di-, tri- or tetrafunctional siloxane units with Si-O bonds, arise from polycondensation according to the number of chlorine atoms of the basic silane molecule.

Silanes may be;

- Monofunctional i.e., there is one Si atom with 3 alkoxy groups in the molecule e.g., vinyltriethoxysilane.
- R-Si-(OR)₃ where OR=alkoxy group.
- Bisfunctional i.e. there are 2 Si atoms, each with 3 alkoxy groups in the molecule e.g., bis(3-trimethoxysilyl)propylene diamine.
- 3(OR)-Si-R-Si-(OR)₃ where OR=alkoxy group.
- Trisfunctional i.e., there are 3 Si atoms, each with 3 alkoxy groups in the molecule e.g., tris(3trimethoxysilylpropyl) isocyanurate.

The diverse halogenated silanes serve as building blocks for the synthesis of the various product types of silicones such as fluids or resins. Dimethyldichlorosilane enables the formation of long Si-O chains. At first, the hydrolysis or methanolysis of dimethyldichlorosilane gives a mixture of short chained, difunctional and therefore linear, siloxanes with OH groups as well as cyclic siloxanes, having normally between three and six chain units. The linear siloxanes show a helix structure with the methyl groups being freely able to rotate. All silicone fluids, emulsions and rubbers, are based on dimethyldichlorosilane. This is therefore the decisive
If the trichloro compounds are used, a cross-linking between the linear chains is produced as a result of the three reactive sites of silicon. A three-dimensional network is the consequence. This process is crucial in the formation of silicone resins. Monochlorosilanes, on the other hand, because of their single reactive site, can only be used for the terminating of the chain growth by polycondensation. They react as a sort of ‘capping agent’ for the growing silicone chain.

Silanes are applied in polar aqueous alcohol solutions (e.g., ethanol, isopropanol) and in ethyl acetate, but non polar solutions (e.g., n-pentane, n-hexane) have also been investigated. 

Silane oligomers react with inorganic matrix (e.g., silica, metal oxide that contain hydroxyl-OH groups) to form –Si–O–M–bonds and above and between them–Si–O–Si-bonds. Thus a film (50 to 100 nm) will be formed, containing a hydrophobic and branched polysiloxane layer, that may also contain free hydrogen-bonded oligomers. Also, free water molecules can occur in the film. If the substrate is silica (quartz, SiO$_2$) or silicate, only a siloxane layer, -Si-O-Si – will be formed. Schematic equation for reaction of silanols with metal surface can be presented as:

$$\text{-R'}–\text{Si}–(\text{OH})_2–\text{O}–\text{Si}–(\text{R'}) (\text{OH})' + 2\text{OH}–\text{M}$$

$$\text{R'} \quad \text{R'}$$

$$\text{M} \quad \text{M}$$

Siloxane polymers occur in a variety of common forms - fluids, gels, elastomers and resins - for a wide variety of applications.

Silicon fluids are usually straight chains of poly(dimethylsiloxane) or PDMS, which are terminated with a trimethylsilyl group (or groups). PDMS fluids come in all viscosities - from water - like liquids to intractable fluids. All of these are essentially water insoluble. PDMS fluids may be further modified with the addition of organofunctional groups at any point in the polymer chain.

Silicone gels are lightly cross-linked PDMS fluids, where the cross-link is introduced either through a trifunctional silane - such as CH$_3$SiCl$_3$ giving a “T-branched” silicone structure, or through a chemical reaction between a Si- vinyl group on one polymer chain, with a hydrogen bonded to silicon in another. This chemical “tying” of siloxane chains produces a three-dimensional network that can be swollen with PDMS fluids, to give a sticky, cohesive mass without form.

Silicone elastomers are cross-linked fluids, whose three-dimensional structure is much more intricate than a gel. In addition, there is very little free fluid in the matrix. Fillers, such as amorphous silica, are frequently added to the matrix to give greater reinforcement to the network and thereby increase the strength of the product.

Silicone resins are more heavily cross-linked polymer networks, that are formed by the introduction of a predominance of tri- and tetra-functional monomers, such as CH$_3$SiCl$_3$. The physical properties of the finished silicone resin can be tailored to suit many applications, by varying the ratio of branched and linear siloxanes and also the functional groups attached to the silicon.

**SILANE APPLICATIONS**

Silanes are versatile materials used in a wide range of applications including adhesion promoters, coupling agents, crosslinking agents, dispersing agents, and surface modifiers.

**Adhesion promoters**

When added to adhesives or used as a primer on substrates, silanes often provide dramatic improvement in adhesion, by reducing moisture attack at the interface. This results in improved:

- Moisture resistance
- Temperature resistance
- Chemicals resistance

**Crosslinking agents**

Silanes can be used to crosslink polymers such as acrylates, polyethers, polyurethanes, and polyesters, to improve:

- Tear resistance
- Elongation at the break
- Tear propagation resistance
- Abrasion resistance

**Water scavengers**

Since silanes react very rapidly with water, they can be used in a formulation to capture excess moisture to:

- Prevent premature cure during compounding
- Enhance uniform curing
- Improve package stability

**Coupling agents**

They are defined as chemical substances capable of reacting with both the reinforcement and the resin matrix of a composite material. It may also bond inorganic fillers or fibers to organic resins, to form or promote...
a stronger bond at the interface. The agent acts as an interface between the resin and glass fiber (or mineral filler), to form a chemical bridge between the two. When used as a coupling agent, silanes bind organic polymers to mineral or siliceous fillers, resulting in:

- Improved mixing
- Better bonding of pigment or fillers to resins
- Increased matrix strength
- Decrease water intake of composite
- Minimize wear

**SILANE APPLICATIONS IN DENTISTRY**

**Ceramic restorations and repairs**

Previously, bonding between gold alloys and porcelain was mainly by micromechanical attachment techniques based on roughening the contact surfaces and applying hot temperatures.[3] Silane coupling agents improve the bond of resin composite to ceramic, by about 25%. Evaluation of silanes establishing bonds to metal or ceramics, showed that silanes do not bond to the metal surface as they do with the ceramic.[6] Siloxane bonds are sensitive to moisture between resin and ceramic matrix.[7] Later, studies confirmed that the use of silane increases the bond strength of resin composite to HF acid etched porcelain.[8] Newer ceramic repair systems consisted of MPS, Ferric chloride (FeCl3) and an adhesive opaque resin.[9] Silanized interfaces appeared to be unstable in humid conditions and the silane bond deteriorated under atmospheric pressure. Since the resins are permeable to water, the bond between silane and resin composite was expected to deteriorate by hydrolysis, over time. Although the silane coupling agents are capable of forming bonds with both inorganic and organic surfaces, silane itself does not help bonding. Therefore it is used in combination with silica coating. Silicatization[10] (i.e., air borne SiO2 particle conditioning, use of acidic primer)[11] (HF acid etching), or heat treatment of silanized substrate. The main contribution to the increased bond strength was not made solely by the mechanical interlocking of the resin composite, but also by the formation of covalent siloxane bonds via silane treatment.

**Silanes on filler materials**

Silanization of the filler particle phase yields a better dispersion and wetting for filler particles and it also lowers the viscosity between a filler and liquid resin system, thereby improving the physical properties of composites.[3] Silanization of E-glass and aramid fibers enhances the adhesion between the fibers and organic acrylic resin system in a denture material.[12]

**Base and noble metal alloys**

Titanium and steel-silane coupling agents are of importance for the adhesion of resin composite to ceramic\metal. For an effective bond of resin to feldspathic porcelain and metal, the use of silane in combination with surface conditioning is required. The use of silanes with different techniques has been advised.

Rocatec system (3M/ESPE), based on SiO2 abrasive particle (e.g., with 50-µm diameter), was meant for roughening the substrate surface. It increases the bond strength effectively, because of the increase in silica content on the substrate surface, that provides a basis for silanes to enhance resin bonding. This is called the Tribochemical method. The temperature at the surface while the particles are hitting, may momentarily rise up to 1,200°C, and the particles thus will be embedded in the surface. Silicoater MD (Heraeus Kulzer)- This method can be used in an extraoral process to attach resin composite onto metal surfaces. It was based on immediate pyrochemical silica layer formation onto the substrate surface. It was also used for cobalt-chromium (Co-Cr) alloy cast clasps coated with resin. Noble alloys roughened by CoJet Sand (3M/ESPE; silica coating), followed by silanization, had significantly higher adherence results to resin composite.[13] For both a gold-palladium-platinum alloy and feldspathic porcelain, a treatment of Rocatec-Preabrasion (with alumina particles), followed by an impact alumina-silica coating (Rocatec-Plus) and finally silanization by a commercial MPS silane is effective, when bonding a composite to alloys.[14]

**Some other silane applications**

- Silanization of orthodontic brackets[15-17] and use of silanized ceramic root canal posts in endodontics,[18] has been advocated.
- Long-range intraoral stain protection has been accomplished in a denture, when its surface was modified with a fluorocarbon chain containing silane.
- Fluoroalkylethyl silane in dental ceramics reduces stress corrosion.
- A nonfunctional silane, (3,3,3-trifluoro-propyl) trimethoxysilane, has been used in filler surface treatment.

**CONCLUSION**

A majority of clinical studies pointed to silanes playing a significant role in the adhesion process. Silane reaction mechanisms were not entirely understood and there exist several theories for bonding mechanism, for silanes and substrates. The essential conclusion for the most effective silanes for adhesion, were that the silane backbone body should be rather short, the organofunctional part should preferably be methacrylic,
the hydrolyzable groups should be methoxy and silane should be applied from ethanol. There is continuous discussion as to whether silanization is needed, or, e.g., whether acid etching by HF acid is adequate, for obtaining clinically relevant attachment of composite to glass ceramics. With ceramics that are not etchable to form a microretentive surface (e.g., alumina, zirconia), optimization of silane-promoted adhesion may play a great role in the future.

REFERENCES