

US 20200115239A1

(19) United States (12) Patent Application Publication (10) Pub. No.: US 2020/0115239 A1
Nikiforov et al. (43) Pub. Date: Apr. 16, 2020

Apr. 16, 2020

(54) PROCESS FOR PRODUCING LIQUID POLYSILANES AND ISOMER ENRICHED HIGHER SILANES

- (71) Applicants : L'Air Liquide , Société Anonyme pour l'Etude et l'Exploitation des Procédés Georges Claude, Paris (FR); American Air Liquide, Inc., Fremont, CA (US)
- (72) Inventors: Grigory Nikiforov, Bridgewater, NJ
(US); Guillaume Husson, Newark, DE (US); Gennadiy Itov, Flemington, NJ (US); Yang Wang, Garnet Valley, PA (US)
- (21) Appl. No.: 16/158,089
- (22) Filed: Oct. 11, 2018

Publication Classification

(51) Int. Cl.
 $C0IB\ 33/04$ B01J 31/02 CO8G 77/60 (2006.01)
 (2006.01) (2006.01)

B01J 31/14 (2006.01)
B01J 23/02 (2006.01) *B01J 23/02*
U.S. Cl.

(52) U.S. CI . CPC COIB 33/04 (2013.01) ; B01J 31/0274 $(2013.01);$ $C08G$ 77/60 $(2013.01);$ $B01J$ 31/143 (2013.01); B01J 2523/31 (2013.01); B01J 2231/76 (2013.01); B01J 2523/12 (2013.01); *B01J 2523/41* (2013.01); **B01J** 23/02 (2013.01)

(57) ABSTRACT

Synthesis of silanes with more than three silicon atoms are disclosed (i.e., $(Si_nH_{(2n+2)}$ with n=4–100). More particularly, the disclosed synthesis methods tune and optimize the isomer ratio by selection of process parameters such as temperature, residence time, and the relative amount of starting compounds, as well as selection of proper catalyst.
The disclosed synthesis methods allow facile preparation of silanes containing more than three silicon atoms and particularly, the silanes containing preferably one major isomer.
The pure isomers and isomer enriched mixtures are prepared
by catalytic transformation of silane (SiH_a) , disilane (Si_2H_6) , trisilane (Si_3H_8) , and mixtures thereof.

Abundance

PROCESS FOR PRODUCING LIQUID POLYSILANES AND ISOMER ENRICHED HIGHER SILANES

TECHNICAL FIELD

[0001] Methods of synthesizing higher silanes are disclosed (i.e., Si_nH_{2n+2} with n=4–100). More particularly, the disclosed synthesis methods tune and optimize the isomer ratio of higher silanes. The isomer ratio may be optimized by selection of process parameters, such as temperature, residence time, and the relative amount of starting compounds. as well as selection of proper catalyst. The disclosed syn-
thesis methods allow facile preparation of higher silanes and
particularly silanes containing one major isomer. The pure
isomers and isomer enriched mixtures are

BACKGROUND

[0002] Polysilanes have been used in a variety of industries .

[0003] Vapor deposition of silicon-containing films using polysilanes is disclosed by JP Pat No 3,185,817 to Seiko Epson Corp.; Kanoh et al., Japanese Journal of Applied Physics, Part 1: Regular Papers, Short Notes & Review Papers 1993, 32(6A), 2613-2619; JP Pat No 3,484,815 to Showa Denko KK; and JP Pat App Pub No 2000/031066 to Showa Denko KK, among others.

[0004] US Pat App Pub No 2010/0184268 A1 claims a method for producing a semiconductor device comprising: coating the coating composition for forming an oxide film comprising: a polysilazane and a polysilane on a substrate and forming the oxide film inside the groove by heat treatment in an oxidizing atmosphere. The formulas of polysilazane $(SiH_2NH)_n$ (n—positive integer) and polysilane Si_nR_{2n+2} and Si_nR_2n (n ≥ 3 , R-hydrogen) are mentioned only in embodiment.

[0005] Epitaxial Si-containing films, such as Si, SiGe, SiC, SiN, and SiO, have been grown using polysilanes as disclosed by Hazbun et al., *Journal of Crystal Growth* 2016, 444, 21-27; US Pat App Pub No 2017/018427 to Yi-Chiau Huang et al.; US Pat App Pub No 2016/126093 to Dube et al .; and Hart et al., *Thin Solid Films* 2016, 604, 23-27]; among others.

[0006] Polysilanes have been used as inks for printed electronics as disclosed by US Pat App Pub No 2009/ 0269559 to Lee et al.; PCT Pub No WO 2015/085980 to Forschungszentrum Juelich GmbH; US Pat App Pub No 2010/197102 to Akao et al.; and JP Pat No $\overline{6,191,821}$ to Showa Denko K K; among others.

[0007] Polysilanes have also been used as high specific energy fuels as disclosed by Simone et al., Journal of Propulsion and Power 2006, 22, 1006-1011; and Hidding et al., Journal of Propulsion and Power 2006, 22, 786-789], among others.

[0008] Conversion of lower silanes into higher silanes has
been studied extensively both for research and for commercial purposes. Catalytic reactions have been studied. See, e.g., U.S. Pat. No. 5,047,569 to Berris; Corey et al., Organometallics, 1991, 10, 924-930; Boudjouk et al., J. Chem. Soc. Chem. Comm. 1991 245-246; U.S. Pat. No. 5 7047-7055; Ohshita et al., Organometallics 1994 13, 50025012; Bourg et al., Organometallics, 1995, 14, 564-566; Bourg et al., Organometallics 1995, 14, 564-566; U.S. Pat. No. 5,700,400 to Ikai et al.; Woo et al., Mol. Cryst. Liq. Cryst. Sci. Technol., Sect. A, 2000, 349, 87; Rosenberg et al., J. Am. Chem. Soc. 2001, 123, 5120-5121; Fontaine et al., Organometallics 2002, 21, 401-408; Kim et 48, 3313-3316; PCT Pub No W02010/003729 to Evonik Degussa GMBH; Smith et al., Organometallics 2010, 29, 6527-6533; PCT Pub No WO2012/001180 to SPAWNT PRIVAT S.A.R.L; PCT Pub No WO2013/019208 to Kovio, Inc.; Feigl et al., Chem. Eur. J. 2013, 19, 12526-12536; Tanabe et al., Organometallics 2013, 32, 1037-1043; U.S. Pat. No. 8,709,369 to Brausch et al.; Schmidt et al., Dalton Trans. 2014, 43, 10816-10827; and U.S. Pat. No. 9,567,228 to Matsushita et al.

[0009] All these disclosures notwithstanding, commercial use of polysilanes remains elusive .

SUMMARY

[0010] Methods of producing $Si_nH_{(2n+2)}$, wherein n=4-100 are disclosed. A liquid $Si_aH_{(2a+2)}$ reactant, wherein a=1-4, is transformed in the presence of a catalyst to produce Si_nH_(2n+2), wherein n>a. The catalyst may be a) a heterogeneous catalyst selected from a Group I, II or III element or oxides, alkyls, hydrides, silanides, or silyl amides thereof; or b) a catalyst comprising a Fe halide combined with an alkyl lithium LiR or lithium amide LiNR₂, with each R independently a C1-C4 alkyl group. Alternatively, $Si_nH_(2n+2)$, wherein n=4-100, may be produced by catalytically converting a $\text{Si}_a\text{H}_{(2a+2)}$ reactant, wherein a=1-4 and n>a. In another alternative, a $\text{Si}_a\text{H}_{(2a+2)}$ reactant, wherein a=1-4, is reacted with a heterogeneous catalyst to produce $\text{Si}_{n}H_{(2n+2)}$, wherein n=4-100 and n>a. In yet another alternative, a $\text{Si}_{a}H_{(2a+2)}$ reactant, wherein a=1-4, is contacted with a heterogeneous catalyst to produce $\text{Si}_{n}H_{(2n+2)}$, wherein n=4-100 and n $>$ a.

[0011] Methods of selectively synthesizing isomerically enriched polysilanes having a formula $Si_nH_{(2n+2)}$, wherein n=5-8, are also disclosed. A liquid $Si_nH_{(2n+2)}$ reactant, wherein $n=1-4$, is catalytically converted to the isomerically
enriched polysilane having a ratio of one isomer to another
isomer ranging from approximately 2:1 to approximately $15:1$

[0012] Any of these disclosed methods may include one or more of the following aspects :

- $[0013]$ n=4-10;
- [0014] the catalyst being sodium bis (2-methoxyethoxy) aluminum hydride;
- [0015] the catalyst being $KN(SiMe₃)$ ₂;
- [0016] the catalyst being NaN(SiMe_3)₂;
- [0017] the catalyst being K, Na, $Na₂O$, K₂O, or combinations thereof;
- [0018] the catalyst being K, K_2O , or combinations thereof:
- [0019] the catalyst being Na, $Na₂O$, or combinations thereof;
- $[0020]$ the catalyst being K;
- [0021] the catalyst being Na;
- [0022] the catalyst being Na₂O;
[0023] the catalyst being K₂O;
- [0023] the catalyst being K_2O ;
[0024] $n=10-30$;
- $[0024]$ n=10-30;
[0025] the cataly
- 025] the catalyst comprising a Fe halide combined with an alkyl lithium LiR or lithium amide LiNR₂, with each R independently a C1-C4 alkyl group;
- [0026] the catalyst comprising 5% w/w FeCl₃ combined with BuLi;
- [0027] the catalyst comprising 5% w/w FeCl₃ combined with $LiNMe₂$;
- $[0028]$ n=30-50;
- [0029] the method not utilizing H_2 ;
- [0030] the $\text{Si}_aH_{(2a+2)}$ reactant being a liquid;
- [0031] the $\text{Si}_{\alpha} \text{H}_{(2\alpha+2)}$ reactant being a mixture of a liquid and a gas;
[0032] the $\text{Si}_{\alpha} \text{H}_{(2\alpha+2)}$ reactant being $\text{Si}_{3} \text{H}_{8}$;
-
- [0033] the $\text{Si}_a H_{(2a+2)}$ reactant being liquid $\text{Si}_3 H_8$;
- [0034] the $\text{Si}_2\text{H}_{(2a+2)}$ reactant being a mixture of Si_2H_6 and Si_3H_8 ;
- [0035] the $\text{Si}_a\text{H}_{(2a+2)}$ reactant being a liquid mixture of $Si₂H₆$ and $Si₃H₈$;
- [0036] the $\text{Si}_2\text{H}_{(2a+2)}$ reactant being a mixture of gasedus Si_2H_6 and liquid Si_3H_8 ;
- [0037] the mixture comprising between approximately 0.1% w/w to approximately 60% w/w Si₃H_s and between approximately 40% w/w and 99.9% w/w
- $Si₂H₆$;
[0038] the mixture comprising between approximately 0.1% w/w to approximately 25% w/w $Si₃H₈$ and between approximately 75% w/w and 99.9% w/w Si_2H_6 ;
- [0039] the mixture comprising between approximately 0.1% w/w to approximately 10% w/w $Si₃H₈$ and between approximately 90% w/w and 99.9% w/w $Si₂H₆;$
- [0040] the $\text{Si}_a\text{H}_{(2a+2)}$ reactant being a mixture of Si_3H_8 and Si_4H_{10} ;
- [0041] the $Si_aH_{(2a+2)}$ reactant being a liquid mixture of Si_3H_8 and Si_4H_{10} ;
- [0042] the Si_aH_(2a+2) reactant being a mixture of gas-
eous Si₃H_s and liquid Si₄H₁₀;
[0043] the mixture comprising between approximately
- 0.1% w/w to approximately 60% w/w Si_4H_{10} and between approximately 40% w/w and 99.9% w/w $Si₃H₈$;
[0044] the mixture comprising between approximately
- 0.1% w/w to approximately 25% w/w Si_4H_{10} and between approximately 75% w/w and 99.9% w/w $Si₂H₈;$
- [0045] the mixture comprising between approximately 0.1% w/w to approximately 10% w/w $Si_{4H₁₀}}$ and between approximately 90% w/w and 99.9% w/w Si_3H_8 ;
- [0046] converting approximately 20% w/w to approximately 60% w/w of the $Si_aH_{(2a+2)}$ reactant; [0047] heating the $Si_aH_{(2a+2)}$ reactant prior to mixing with the catalyst;
-
- [0048] mixing the $\text{Si}_a\text{H}_{(2a+2)}$ reactant and catalyst to form a reactant-catalyst mixture;
- [0049] mixing the $Si_aH_(2a+2)$ reactant and catalyst to form a reactant-catalyst mixture for a time period ranging from approximately 1 hour to approximately 24 hours;
- [0050] heating the reactant-catalyst mixture to a temperature ranging from approximately 30° C. to approximately 55° C.;
- [0051] mixing the reactant-catalyst mixture at a temperature ranging from approximately room temperature to approximately 53° C.;
- [0052] mixing the reactant-catalyst mixture at a temperature ranging from approximately 15° C. to approximately 50° C.;
- [0053] mixing the reactant-catalyst mixture at a temperature ranging from approximately 15° C. to approximately 30° C.;
- [0054] filtering the reactant-catalyst mixture to separate any solids from the resulting $Si_nH_{(2n+2)}$ mixture;
- [0055] heating the $Si_aH_{(2a+2)}$ reactant prior to flowing through a reactor containing the catalyst;
- [0056] heating the $\text{Si}_n\text{H}_{(2n+2)}$ reactant prior to flowing through the catalyst;
- [0057] flowing the $\text{Si}_{a}H_{(2a+2)}$ reactant through a reactor containing the catalyst;
- [0058] flowing the Si_aH_(2a+2) reactant through a reactor containing the catalyst on glass wool;
- [0059] flowing the $Si_aH_{(2a+2)}$ reactant through a reactor containing the catalyst pellets;
- [0060] flowing the Si_aH_(2a+2) reactant through a reactor containing the catalyst to produce a Si_nH_(2n+2) mixture; [0061] the Si_aH_(2a+2) reactant having a residence time in
- the reactor ranging from approximately 200 seconds to
approximately 600 seconds;
 $[0062]$ heating the reactor to a temperature ranging from
-
- approximately 15 \degree C. to approximately 170 \degree C.;
[0063] heating the reactor to a temperature ranging from
- approximately 15 \degree C. to approximately 150 \degree C.;
[0064] heating the reactor to a temperature ranging from approximately 15° C. to approximately 100° C.;
- [0065] heating the reactor to a temperature ranging from approximately 15° C. to approximately 50° C.;
[0066] heating the reactor to a temperature ranging from
- approximately 20° C. to approximately 150 $^{\circ}$ C.;
- [0067] heating the reactor to a temperature ranging from approximately 50 \degree C. to approximately 100 \degree C.;
- [0068] heating the reactor to a temperature ranging from approximately 40° C. to approximately 150° C.;
[0069] maintaining the reactor at a pressure ranging
- from approximately 10 psig (69 kPa) to approximately 50 psig (345 kPa);
- [0070] the method not requiring a quenching agent;
[0071] recycling unreacted $Si_aH_{(2a+2)}$ reactant;
-
- [0072] the heterogeneous catalyst comprising an element selected from Group I of the Periodic Table;
- [0073] the heterogeneous catalyst comprising K;
[0074] the heterogeneous catalyst comprising Na
- [0074] the heterogeneous catalyst comprising Na;
[0075] the heterogeneous catalyst comprising an
- 075] the heterogeneous catalyst comprising an element selected from Group III of the Periodic Table;
- $[0076]$ the heterogeneous catalyst comprising Al;
- [0077] the heterogeneous catalyst comprising both an element selected from Group I and an element from Group III of the Periodic Table;
- [0078] the heterogeneous catalyst comprising Na and Al;
100791
- the heterogeneous catalyst comprising Li and Al;
- [0080] the heterogeneous catalyst being selected from a Group I, II or III element or oxides, alkyls, hydrides, silanides, or silyl amides thereof;
- [0081] the heterogeneous catalyst being an oxide catalyst:
- [0082] the heterogeneous catalyst being a Group I metal oxide catalyst;
- [0083] the heterogeneous catalyst being Na₂O;
[0084] the heterogeneous catalyst being K₂O;
- [0084] the heterogeneous catalyst being K_2O ;
[0085] the heterogeneous catalyst being K,
- the heterogeneous catalyst being K, $K₂O$, or combinations thereof;
- [0086] the heterogeneous catalyst being Na, $Na₂O$, or combinations thereof;
- [0087] the hydride catalyst comprising two elements from Group I, II, or III of the Periodic Table;
- [0088] the hydride catalyst being selected from the group consisting of LiAlH₄, LiAlH_nR_{4-n}, NaAlH_nR_{4-n}, $KAIH_nR_{4-n}$, $RbAIH_nR_{4-n}$, $CsAIH_nR_{4-n}$, and combinations thereof, wherein n=1, 2, or 3 and each R is independently C_mH_{2m+1} with m=1-10 or an aliphatic group having an oxygen or nitrogen atom;
- [0089] the hydride catalyst being $LiAlH₄$;
- [0090] the hydride catalyst being $LiAlH_nR_{4-n}$, wherein n=1, 2, or 3 and each R is independently C_mH_{2m+1} with $m=1-10$ or an aliphatic group having an oxygen or nitrogen atom:
- [0091] the hydride catalyst being selected from the group consisting of NaAlH_nR_{4-n}, wherein n=1, 2, or 3 and each R is independently C_mH_{2m+1} with m=1-10 or an aliphatic group having an oxygen or nitrogen atom;
- [0092] the hydride catalyst being selected from the group consisting of $KAlH_nR_{4-n}$, wherein n=1, 2, or 3 and each R is independently C_mH_{2m+1} with m=1-10 or an aliphatic group having an oxygen or nitrogen atom;
- [0093] the hydride catalyst being selected from the group consisting of $RbAlH_nR_{4-n}$, wherein n=1, 2, or 3 and each R is independently C_mH_{2m+1} with m=1-10 or an aliphatic group having an oxygen or nitrogen atom;
- [0094] the hydride catalyst being selected from the group consisting of CsAlH_nR_{4-n}, wherein n=1, 2, or 3 and each R is independently $\mathrm{C}_m\mathrm{H}_{2m+1}$ with m=1-10 or an aliphatic group having an oxygen or nitrogen atom;
- [0095] the aliphatic group having an oxygen or nitrogen atom being selected from the group consisting of $-CH_2OMe$, $-CH_2CH_2OMe$, $-OCH_2CH_2CH_2CH_2OMe$, $-\text{CH}_2\text{CH}_2\text{NMe}_2$, an aromatic group, and combinations thereof;
- $[0096]$ the heterogeneous catalyst being sodium bis $(2$ methoxyethoxy) aluminum hydride $[Na(-O(Me)-C_2H_4-O_2A]H_2]$;
- [0097] the aromatic group being phenyl or substituted phenyl;
- [0098] the heterogeneous catalyst being a metal silyl-
amide catalyst;
- [0099] the metal silylamide catalyst having the formula M[N(SiR₃)₂]_x, wherein x=1 when M is Li, Na, K, Rb, or Cs; $x=2$ when M is Leg and Sr, or Ba; $x=3$ when M = Al or Ga; and each R is independently C_mH_{2m+1}
- with $m=1-10$ or an aromatic group;
[0100] the metal silylamide catalyst having the formula Li [N(SiR₃)₂], wherein each R is independently C_mH_{2m+1} with m=1-10 or an aromatic group;
- [0101] the metal silylamide catalyst having the formula $Na[N(SiR_3)_2]$, wherein each R is independently $C_m H_{2m+1}$ with m=1-10 or an aromatic group;
- [0102] the metal silylamide catalyst having the formula
Na[N(SiR₃)₂], wherein each R is independently $C_m H_{2m+1}$ with m=1-4;
[0103] the metal silylamide catalyst being sodium bis
- (trimethylsilyl) amide;
- [0104] the metal silylamide catalyst having the formula $K[N(SiR₃)₂]$, wherein each R is independently C_mH_{2m+1} with m=1-10 or an aromatic group;
[0105] the metal silylamide catalyst having the formula
- $\mathrm{C}_m\mathrm{H}_{2m+1}$ with m=1-4; $K[N(SiR₃)₂]$, wherein each R is independently
- [0106] the metal silylamide catalyst being potassium bis (trimethylsilyl) amide;
- [0107] the metal silylamide catalyst having the formula $Mg[N(SiR₃)₂]$, wherein each R is independently C_mH_{2m+1} with m=1-10 or an aromatic group;
- [0108] the metal silylamide catalyst having the formula $Ca[N(SiR₃)₂]$, wherein each R is independently $C_m H_{2m+1}$ with m=1-10 or an aromatic group;
- [0109] the metal silylamide catalyst having the formula
Sr[N(SiR₃)₂]₂, wherein each R is independently
- C_mH_{2m+1} with m=1-10 or an aromatic group;
[0110] the metal silylamide catalyst having the formula
Ba[N(SiR₃)₂]₂, wherein each R is independently
- C_mH_{2m+1} with m=1-10 or an aromatic group;
[0111] the metal silylamide catalyst having the formula
Al[N(SiR₃)₂]₃, wherein each R is independently C_mH_{2m+1} with m=1-10 or an aromatic group;
[0112] the metal silylamide catalyst having the formula
- Ga[N(SiR_3)₂]₃, wherein each R is independently C_mH_{2m+1} with m=1-10 or an aromatic group;
- [0113] $\;$ the heterogeneous catalyst being a metal silanide catalyst:
- [0114] the metal silanide catalyst having the formula $Na[Al(SiH₃)(Si₂H₅)(OCH₂CHp₂OMe)₂];$
- [0115] the metal silanide catalyst having the formula $M(Si_nH_{2n+1})_x$ or $M(SiR_3)_x$, wherein n=1-4, x=1 when M is Li, Na, K, Rb, or Cs; x=2 when M — Mg, Ca, Sr, or Ba; $x=3$ when M \equiv Al or Ga; and each R is independently C_mH_{2m+1} with m=1-10 or an aromatic group;
- [0116] the metal silanide catalyst having the formula $M(Si_nH_{2n+1})$, wherein n=1-4 and x=1 when M is Li, Na, K, Rb, or Cs; $x=2$ when M M_{max} Ca, Sr, or Ba; and $x=3$ when M $=$ Al or Ga;
- [0117] the metal silanide catalyst having the formula $LiSiH₃$;
- [0118] the metal silanide catalyst having the formula NaSiH₃;
- [0119] the metal silanide catalyst having the formula $KSiH_3$;
- [0120] the metal silanide catalyst having the formula $RbSiH₃;$
- [0121] the metal silanide catalyst having the formula $CsSiH₃$;
- [0122] the metal silanide catalyst having the formula $Lisiph$ ₂;
- [0123] the metal silanide catalyst having the formula NaSiPh₃;
- [0124] the metal silanide catalyst having the formula $KSiPh_3$;
- [0125] the metal silanide catalyst having the formula $RbSiPh₃$:
- [0126] the metal silanide catalyst having the formula $CsSiPh₃$;
- [0127] the metal silanide catalyst having the formula $LiSi₂H₅;$
- [0128] the metal silanide catalyst having the formula NaSi₃H₅;
- $[0129]$ the metal silanide catalyst having the formula $KSi₂H₅;$
- [0130] the metal silanide catalyst having the formula $RbSi₂H₅;$
- [0131] the metal silanide catalyst having the formula CSSi_2H_5 ,
- [0132] the metal silanide catalyst having the formula $Mg(SiH₃)₂$,
- [0133] the metal silanide catalyst having the formula $Ca(SiH_3)$,
- [0134] the metal silanide catalyst having the formula $Sr(SiH₃)$ ₂,
- [0135] the metal silanide catalyst having the formula $Ba(SiH_3)$,
- $[0136]$ the metal silanide catalyst having the formula $AI(SiH₃)₃;$
- [0137] the metal silanide catalyst having the formula $Ga(SiH_3)_3;$
- [0138] the metal silanide catalyst having the formula $M(SiR_3)$, wherein x=1 when M=Li, Na, K, Rb, or Cs; $x=2$ when M M_{max} , Ca, Sr, or Ba; $x=3$ when M max or Ga; and each R is independently C_mH_{2m+1} with m=1-10 or an aromatic group;
[0139] the metal silanide catalyst having the formula
- Li(SiR₃), wherein each R is independently $C_m H_{2m+1}$ with $m=1-10$ or an aromatic group;
- [0140] the metal silanide catalyst having the formula $Na(SiR₃)$, wherein each R is independently C_mH_{2m+1} with $m=1-10$ or an aromatic group;
- [0141] the metal silanide catalyst having the formula K(SiR₃), wherein each R is independently C_mH_{2m+1}
with m=1-10 or an aromatic group: with $m=1-10$ or an aromatic group;
[0142] the metal silanide catalyst having the formula
- $Rb(SiR_2)$, wherein each R is independently C_mH_{2m+1} with $m=1-10$ or an aromatic group;
- [0143] the metal silanide catalyst having the formula $Cs(SiR_3)$, wherein each R is independently C_mH_{2m+1} with $m=1-10$ or an aromatic group;
- [0144] the metal silanide catalyst having the formula $Mg(SiR_3)_2$, wherein each R is independently C_mH_{2m+1} with $m=1-10$ or an aromatic group;
[0145] the metal silanide catalyst having the formula
- $[0145]$ the metal shamue catalyst having the formula α (SiR₃)₂, wherein each R is independently $C_m \Pi_{2m+1}$ with $m=1-10$ or an aromatic group;
- [0146] the metal silanide catalyst having the formula $Sr(SiR₃)₂$, wherein each R is independently C_mH_{2m+1} with $m=1-10$ or an aromatic group;
- [0147] the metal silanide catalyst having the formula $Ba(SiR_3)_2$, wherein each R is independently C_mH_{2m+1} with $m=1-10$ or an aromatic group;
[0148] the metal silanide catalyst having the formula
- $\text{Al(SiR}_3)$ ₃, wherein each R is independently C_mH_{2m+1} with $m=1-10$ or an aromatic group;
- [0149] the metal silanide catalyst having the formula $Ga(SiR_3)_3$, wherein each R is independently C_mH_{2m+1} with $m=1-10$ or an aromatic group;
- [0150] the heterogeneous catalyst being on a support;
- [0151] the catalyst being physically bound to a support;
-
- [0152] the catalyst being chemically bound to a support;
[0153] the catalyst being both physically and chemithe catalyst being both physically and chemi-
- cally bound to a support;
- [0154] the support being alumina (Al_2O_3) , silica (SiO_2) , or combinations thereof;
- [0155] the support being alumina (Al_2O_3) ;
- [0156] the support being silica ($SiO₂$);
- [0157] the heterogeneous catalyst being in pellet form;
[0158] the heterogeneous catalyst comprising approxi-
-
- mately 0.1% w/w to approximately 70% w/w of the
heterogeneous catalyst and support combination;
[0159] the heterogeneous catalyst comprising approxi-
mately 1% w/w to approximately 50% w/w of the
heterogeneous catalyst and
- [0160] the heterogeneous catalyst comprising approximately 1% w/w to approximately 5% w/w of the
- heterogeneous catalyst and support combination;
[0161] fractionally distilling $Si_nH_{(2n+2)}$ to produce a Si-containing film forming composition comprising approximately 95% w/w to approximately 100% w/w
- n-Si_sH₁₂;
[0162] fractionally distilling $\text{Si}_{n}H_{(2n+2)}$ to produce a Si-containing film forming composition comprising approximately 95% w/w to approximately 100% w/w n-Si₆H₁₄;
- [0163] fractionally distilling $Si_nH_(2n+2)$ to produce a Si-containing film forming composition comprising approximately 95% w/w to approximately 100% w/w
- n-Si₇H₁₆; and/or
[0164] fractionally distilling $Si_nH_{(2n+2)}$ to produce a Si-containing film forming composition comprising approximately 95% w/w to approximately 100% w/w n-Si_sH₁₈.

[0165] Si-containing film forming composition produced by any of the methods disclosed above are also disclosed. The disclosed compositions may further include one or more of the following aspects: [0166] the Si-containing

- comprising approximately 95% w/w to approximately
- 100% w/w n-Si_sH₁₂;
 [0167] the Si-containing film forming composition comprising approximately 95% w/w to approximately 100% w/w n-Si₆H₁₄;
- [0168] the Si-containing film forming composition comprising approximately 95% w/w to approximately
- 100% w/w n-Si₇H₁₆;
 [0169] the Si-containing film forming composition comprising approximately 95% w/w to approximately
- 100% w/w n-Si₈H₁₈;
 [0170] the Si-containing film forming composition comprising approximately 0 ppmw to approximately 100 ppmw halide contaminants;
- [0171] the Si-containing film forming composition comprising approximately 0 ppmw to approximately 25 ppmw halide contaminants; and/or
- [0172] the Si-containing film forming composition comprising approximately 0 ppmw to approximately 5 ppmw halide contaminants.

[0173] Methods of maintaining the vapor pressure of a volatile polysilane during vapor deposition processes are also disclosed. The vapor deposition processes use any of the Si-containing film forming composition disclosed above.
The Si-containing film forming composition is maintained at a vaporizing temperature.

- [0174] The disclosed methods may further include one or more of the following aspects:
[0175] the Si-containing film forming composition
	-
	- comprising $Si_nH_{(2n+2)}$, wherein n=4-10;
[0176] the Si-containing film forming composition comprising approximately 90% w/w to approximately
	- 100% w/w Si_5H_{12} ;
[0177] the Si-containing film forming composition comprising approximately 90% w/w to approximately 100% w/w Si_6H_{14} ;
	- [0178] the Si-containing film forming composition comprising approximately 90% w/w to approximately 100% w/w Si_7H_{16} ;
[0179] the Si-containing film forming composition
	- comprising approximately 90% w/w to approximately 100% w/w Si_sH_{1s} ;
	- [0180] the Si-containing film forming composition having an initial vapor pressure at the vaporizing tempera ture;
[0181] the vaporizing temperature ranging from
	-
	- approximately 0 $^{\circ}$ C. to approximately 50 $^{\circ}$ C.;
[0182] maintaining approximately 80% of the initial vapor pressure of the Si-containing film forming composition at the vaporizing temperature until approximately 75% w/w of Si-containing film forming composition is consumed; and/or
	- [0183] maintaining approximately 90% of the initial vapor pressure of the Si-containing film forming composition at the vaporizing temperature until approximately 75% w/w of Si-containing film forming composition is consumed;
	- [0184] maintaining approximately 95% of the initial vapor pressure of the Si-containing film forming composition at the vaporizing temperature until approximately 75% w/w of Si-containing film forming composition is consumed.

[0185] Methods of reducing the formation of branched polysilanes during polymerization are also disclosed. The polymerization processes use any of the Si-containing film forming compositions disclosed above . The disclosed meth ods may further include one or more of the following aspects :

- [0186] the Si-containing film forming composition comprising approximately 90% w/w to approximately 100% w/w Si_sH₁₂;
- [0187] the Si-containing film forming composition comprising approximately 90% w/w to approximately 100% w/w Si₆H₁₄;
- [0188] the Si-containing film forming composition comprising approximately 90% w/w to approximately 100% w/w Si₂H₁₆; and/or
- [0189] the Si-containing film forming composition comprising approximately 90% w/w to approximately 100% w/w \overline{Si}_8H_{18} .
[0190] Also disclosed are coating methods of forming

Si-containing films on substrates. The Si-containing film forming compositions disclosed above are contacted with the substrate and the Si-containing film formed via a spin coating, spray coating, dip coating, or slit coating technique.
The disclosed methods may include the following aspects:

[0191] the Si-containing film forming composition fur-
ther comprising between approximately 0.5% w/w to approximately 99.5% w/w of perhydropolysilazane;

- [0192] the Si-containing film forming composition fur-
ther comprising between approximately 10% w/w to approximately 90% w/w of perhydropolysilazane;
- [0193] forming the Si-containing film via a spin coating technique;
- [0194] forming the Si-containing film via a spray coating technique;
- [0195] forming the Si-containing film via a dip coating technique;
- [0196] forming the Si-containing film via a slit coating technique:
- [0197] thermal curing the Si-containing film;
[0198] photon curing the Si-containing film;
-
- [0199] annealing the Si-containing film;
- [0200] laser treating the Si-containing film;
- [0201] the Si-containing film being Si;
- [0202] the Si-containing film being $SiO₂$;
[0203] the SiO₂ film having a wet etch rate ranging from
- approximately 1 to approximately 5 as compared to thermal oxide grown at 1100° C.;
[0204] the SiO₂ film having a wet etch rate ranging from
- approximately 1 to approximately 3 as compared to thermal oxide grown at 1100° C.;
- [0205] the Si-containing film being SiN;
- $[0206]$ the Si-containing film being SiC;
-
- [0207] the Si-containing film being SiON;
[0208] the substrate comprising trenches having an aspect ratio ranging from approximately 1:1 to approximately 1:100; and/or
[0209] the trenches having a critical dimension ranging
- from approximately 10 nm to approximately 1 micron.

Notation and Nomenclature

[0210] Certain abbreviations , symbols , and terms are used throughout the following description and claims , and include:

[0211] As used herein, the indefinite article "a" or "an" means one or more.

[0212] As used herein, the terms "approximately" or " about" mean $\pm 10\%$ of the value stated.

[0213] As used herein, the term "comprising" is inclusive or open-ended and does not exclude additional, unrecited materials or method steps; the term " consisting essentially of" limits the scope of a claim to the specified materials or steps and additional materials or steps that do not materially affect the basic and novel characteristics of the claimed invention; and the term "consisting of" excludes any additional materials or method steps not specified in the claim. $[0214]$ As used herein, the term " higher silanes" means Si_nH_{2n+2} , wherein n=4-100 and the term "lower silanes" means S_aH_{2a+2} with a=1-4. The higher silanes may be linear or branched.

[0215] As used herein, the term "catalyst" means a substance that increases the rate of a reaction without modifying the overall standard Gibbs energy change in the reaction. As used herein, the term "catalyst" includes substances that do not undergo any permanent chemical change as well as those that do (the latter sometimes referred to as "pre-catalysts"). [0216] As used herein, the term " heterogeneous catalyst" means a catalyst which is present in a different phase from

the reactants $(e.g., a solid catalyst versus a liquid reactant;$ or a liquid catalyst that is not capable of being mixed with

a liquid reactant). The heterogeneous catalyst may be blended with a support, which is intrinsically inert or less active than the catalyst.

 $[0217]$ As used herein, the term "quenching agent" means a substance that deactivates a reaction.

[0218] As used herein, the term "residence time" means the amount of time the lower silane reactant spends in the

[0219] As used herein, the terms "perhydropolysilazane" or "PHPS" mean a molecule, oligomer, or polymer containing only Si, H, and N characterized by repeating —SiH_x— NH- units, with $x=0-2$, and the fact that the silicon atom is only bonded to N or H atoms.

 $[0220]$ As used herein, the abbreviation " RT " means room temperature, which is a temperature ranging from approximately 18° C. to approximately 25° C.

[0221] As used herein, the term "hydrocarbyl group" refers to a functional group containing carbon and hydrogen; the term "alkyl group" refers to saturated functional groups containing exclusively carbon and hydrogen atoms. The hydrocarbyl group may be saturated or unsaturated. Either term refers to linear, branched, or cyclic groups. Examples of linear alkyl groups include without limitation, methyl groups, ethyl groups, propyl groups, butyl groups, etc. Examples of branched alkyls groups include without limi without limitation, cyclopropyl groups, cyclopentyl groups, cyclohexyl groups, etc.

[0222] As used herein, the abbreviation "Me" refers to a methyl group; the abbreviation "Et" refers to an ethyl group; the abbreviation "Pr" refers to a propyl group; the abbreviation "nPr" refers to a "normal" or linear propyl group; the abbreviation "iPr" refers to an isopropyl group; the abbreviation "Bu" refers to a butyl group; the abbreviation "nBu" refers to a "normal" or linear butyl group; the abbreviation "tBu" refers to a tert-butyl group, also known as 1,1dimethylethyl; the abbreviation "sBu" refers to a sec-butyl group, also known as 1-methylpropyl; the abbreviation " i Bu" refers to an iso-butyl group, also known as 2-meth-ylpropyl; the term "halide" refers to the halogen anions F -Cl-, Br-, I-, and mixtures thereof; and the abbreviation "TMS" refers to trimethylsilyl or \sim SiMe₃.

[0223] As used herein, the term " aromatic group" refers to cyclic, planar molecules with a ring of resonance bonds that exhibit more stability than other geometric or connective arrangements with the same set of atoms . Exemplary aro matic groups include substituted or unsubstituted phenyl groups (i.e., C_6R_5 , wherein each R is independently H or a hydrocarbyl group).

[0224] As used herein, the term "independently" when used in the context of describing R groups should be understood to denote that the subject R group is not only independently selected relative to other R groups bearing same or different subscripts or superscripts, but is also independently selected relative to any additional species of that same R group. For example in the formula $MR_{x}^{1}(NR^{2}R^{3})_{(4-x)}$, where x is 2 or 3, the two or three R¹ groups may, but need not be identical to each other or to R^2 or to R^3 .
Further, it should be understood that unless specifically stated otherwise, values of R groups are independent of each other when used in different formulas .

[0225] As used herein, the abbreviation M_n , stands for the number averaged molecular weight or the total weight of all of the polymer molecules in a sample divided by the total number of polymer molecules in the sample (i.e., $M_n = \Sigma N_i M_i / \Sigma N_i$, wherein N_i is the number of molecules of weight M_i); the abbreviation M_w stands for weight averaged molecular weight or the sum of the weight fracti of molecule (i.e., $M_w = \Sigma[(N_x M / 2N_x M_x)^* N_x M_x]$; and the term "Poly Dispersity Index" or PDI means the ratio of $M_w M_x$. [0226] The standard abbreviations of the elements from the Periodic Table of elements are used herein. It should be understood that elements may be referred to by these abbre viations (e.g., Si refers to silicon, C refers to carbon, H refers to hydrogen, etc.).

[0227] As used herein, the Periodic Table refers to the tabular arrangement of chemical elements; Group I of the Periodic Table refers to H, Li, Na, K, Rb, Cs, and Fr. Group II of the Periodic Table refers to Be, Mg, Ca, Sr, Ba, and Ra. Group III of the Periodic Table refers to B, Al, Ga, In, TI, and Nh.

[0228] Any and all ranges recited herein are inclusive of their endpoints (i.e., $x=1$ to 4 or x ranges from 1 to 4 includes $x=1$, $x=4$, and $x=$ any number in between), irrespective of whether the term "inclusively" is used.

BRIEF DESCRIPTION OF THE DRAWINGS

[0229] For a further understanding of the nature and objects of the present invention , reference should be made to the following detailed description , taken in conjunction with the accompanying drawing wherein:

[0230] For a further understanding of the nature and objects of the present invention, reference should be made to the following detailed description, taken in conjunction with the accompanying drawing in which reference numbers are

used throughout uniformly, wherein:
[0231] FIG. 1 is a schematic diagram of a batch apparatus in which the disclosed synthesis methods may be performed; [0232] FIG. 2 is a schematic diagram of a flow-through apparatus in which the disclosed synthesis methods may be performed;

[0233] FIG. 3 is a schematic diagram of one embodiment of the flow-through apparatus of FIG. 2;

[0234] FIG. 4 is a schematic diagram of one embodiment of the reactor of FIG. 3;

[0235] FIG. 5 is a flow chart diagraming exemplary processes for the preparation of the Si-containing film forming compositions, preparation of the silicon substrate, and the steps of the spin-coating process;

[0236] FIG. 6 is a GPC of the undistilled liquid of Example 2;

[0237] FIG. 7 is a gas chromatogram of the volatile liquid products from the dry ice trap after one pass of liquid $Si₃H₈$ through the VitrideTM on Silica catalyst (Example 5);

[0238] FIG. 8 is a gas chromatogram of the non-distilled liquid of Example 5.

DESCRIPTION OF PREFERRED EMBODIMENTS

[0239] Methods of synthesizing higher silanes are disclosed (i.e., Si_nH_{2n+2} with n=4-100). Methods of selectively synthesizing isomerically enriched polysilanes having the formula $Si_nH_{(2n+2)}$, wherein n=5-8, are also disclosed.

[0240] Higher silanes exist in various isomers with slight differences in vapor pressure. For example, the boiling point of 80-90% n-Si₄H₁₀ is 107° C. according to the online

catalog from Gelest. In contrast, the boiling point for i-Si₄H₁₀ is 101.7° C. Fehér et al., Inorg. Nucl. Chem. Lett.,

1973, 9, 931.
[0241] In addition to different vapor pressures, the isomers may also have different evaporation behaviour, and thermal stability, due at least to different steric geometries. These differences may create a drift in any vapor processes if one isomer enriches over time. This effect has been demonstrated with other types of isomers (see, e.g., Mehwash Zia
and Muhammad Zia-ul-Haq, Journal of Contemporary Research in Chemistry (2016) 1(1): 34-41). As a result, supplying a higher silane precursor consisting essentially of one isomer, enriched with one isomer, or having a fixed isomer ratio is important for having a reproducible rate of film growth per cycle in vapor deposition processes.

[0242] Similarly, polymerization using the different isomers may produce different polymerization products. In other words, iso-tetrasilane may produce a polymer having more branching than one produced by n-tetrasilane.

[0243] Applicants have discovered methods of tuning and optimizing the tetrasilane isomer ratio as well as selective preparation of polysilanes with a low amount of silicon atoms (6-30). The pure isomers or isomer enriched mixtures are prepared by catalytic transformation of silane (SiH₄), disilane (Si_2H_6), trisilane (Si_3H_8), tetrasilane (Si_4H_{10}), or mixtures thereof. The lower silane reactants (i.e., Si_4H_{2a+2} , with a=1-4) provide an attractive starting material due to commercial availability. Various process parameters may be adjusted to produce the desired polysilane and/or isomer quantity. Exemplary process parameters include the relative amount of the starting compounds and catalyst selection. Temperature and reaction time for a batch process or residence time in a flow through process may also impact isomer yield. The resulting higher silane products are isomer content specific and high purity. One of ordinary will recognize that safety protocols are required when work-

ing with these reactants and products.
[0244] The higher silanes are synthesized by reacting a $Si_2H_{(2a+2)}$ reactant, wherein a=1-4, with a heterogeneous catalyst or a catalyst comprising a Fe halide combined with an alkyl lithium LiR or lithium amide LiNR₂, with each R independently a C1-C4 alkyl group. The $Si_aH_{(2a+2)}$ reactant may be SiH_4 , Si_2H_6 , Si_3H_8 , Si_4H_{10} , or combinations thereof. These reactants are available commercially. These reactants may be used in the disclosed processes in gaseous or liquid form or, for a mixture, as a combination. For example, the reactant may be gaseous $Si₃H₈$ and liquid $Si₄H₁₀$.

[0245] In the examples that follow, the Si_aH_(2a+2) reactant is gaseous or liquid Si₃H₈ or a mixture of liquid Si₃H₈ with liquid Si_2H_6 or Si_4H_{10} . The examples demonstrate that the use of liquid Si_3H_8 produces better n- $Si_4H_{10}/i-Si_4H_{10}$ selectivity as compared to the use of gaseous Si_3H_8 . Liquid Si_3H_8 also produces larger heavier polysilanes ($Si \ge 6$) than those produced by gaseous $Si₃H₈$. Examples 6 and 9 demonstrate that a Si_2H_6/Si_3H_8 reaction mixture reduces the number of heavier polysilanes ($Si \ge 6$) while a Si_3H_8/Si_4H_{10} reaction mixture produces more heavier polysilanes (Siz5). As a result, synthesis of the desired polysilane may be optimized by selecting the appropriate $Si_aH_{(2a+2)}$ reactant. Some optional reactant combinations that will reduce of heavier polysilanes include between approximately 0.1% w/w to approximately 60% w/w $Si₃H₈$ and between approximately 40% w/w and 99.9% w/w Si_2H_6 ; between approximately 0.1% w/w to approximately 25% w/w Si_3H_8 and between approximately 75% w/w and 99.9% w/w Si_2H_6 ; or between approximately 0.1% w/w to approximately 10% w/w Si₃H₈ and between approximately 90% w/w and 99.9% w/w $\rm Si_2H_6$. Some optional reactant combinations that will produce a larger number of heavier polysilanes include between approximately 0.1% w/w to approximately 60% w/w Si_4H_{10} and between approximately 40% w/w and 99.9% w/w Si_3H_8 ; between approximately 0.1% w/w to approximately 25% w/w Si_4H_{10} and between approximately 75% w/w and 99.9% w/w Si_3H_8 ; or between approximately 0.1% w/w to approximately 10% w/w Si_4H_{10} and between approximately 90% w/w and 99.9% w/w Si_3H_8 . [0

alkyls, hydrides, silanides, or silyl amides thereof. The disclosed catalysis reactions do not require the use of a quenching agent because the heterogeneous catalysts may be separated from the reactants and products using filtration in
a batch process. Alternatively, the flow through reactor may be designed having suitable residence time to permit cata-lytic conversion of the $Si_aH_(2a+2)$ reactant.

[0247] The heterogeneous catalyst may be a Group I, II, or III element. Exemplary catalysts include K or Na. These catalysts are commercially available or may be synthesized by methods known in the art. For example, 30-40% $Na/Silica$ is sold under the trademark Active GelTM by SiGNa Chemistry, Inc.

[0248] The heterogeneous catalyst may be a metal oxide catalyst, preferably a Group I oxide catalyst. Exemplary oxide catalyst include $Na₂O$ or K₂O. These catalysts are commercially available or may be synthesized by methods

known in the art.
[0249] The heterogeneous catalyst may be a metal alkyl catalyst MR, with R=C1-C4 hydrocarbon. Exemplary metal alkyl catalysts include BuLi. These catalysts are commercially available or may be synthesized by methods known in the art.

[0250] The metal hydride catalyst may be a mixed metal
hydride catalyst. Exemplary mixed metal hydride catalysts
include LiAlH₄, LiAlH_nR_{4-n}, NaAlH_nR_{4-n}, KAlH_nR_{4-n},
RbAlH_nR_{4-n}, CsAlH_nR_{4-n}, and combina nitrogen atom. Exemplary aliphatic groups having an oxygen or nitrogen atom include —CH₂OMe, —CH₂CH₂OMe, $\overline{}$ -OCH₂CH₂CH₂OMe, -CH₂CH₂NMe₂, an aromatic group, and combinations thereof. A particularly preferred mixed metal halide catalysts is sodium bis(2-methoxy-
ethoxy)aluminum hydride [Na($-O(Me) - C_2H_4-O$)
 $_2AlH_2$]. These catalysts are commercially available or may
be synthesized by methods known in the art.
[0251] The heterog

formula $M[N(SiR_3)_2]_x$, wherein x=1 when M is Li, Na, or K; x=2 when M—Mg, Ca, Sr, or Ba; x=3 when M—Al or Ga;
and each R is independently C_mH_{2m+1} with m=1-10 or an aromatic group, preferably with $m=1-4$. Particularly pre-
ferred metal silylamide catalysts include sodium bis(trim-
ethylsilyl)amide and potassium bis(trimethylsilyl)amide.
These catalysts are commercially available or

catalyst . The metal silanide catalyst may have the formula $Na[A(SiH₃)(Si₂H₅)(OCH₂CH₂OMe)₂], M(Si_nH_{2n+1})_x, or$ $M(SiR_3)_x$, wherein n=1-4 and x=1 when M is Li, Na, K, Rb, or Cs; x=2 when M—Mg, Ca, Sr, or Ba; x=3 when M—A1 or Ga; and each R is independently C_mH_{2m+1} with m=1-10 or an aromatic group. Currently, these precursors are not commercially available and are difficult to synthesize because they are pyrophoric solids. However, based on GC and FT-IR data, Applicants believe that the metal silylamide catalysts, metal alkyl catalysts, or $Na[AH_2(OCH_2CH_2OMe)_{21}$ sold under the trademark VitrideTM by Vertellus Holdings LLC or Red-Al® by Sigma-Aldrich
Biotechnology LP, may act as pre-catalysts and convert to
metal silanide catalysts. More particularly, Applicants believe that the following reactions may occur: or

 $Si_3H_8+KN(SiMe_3)_2 \rightarrow KSi_2H_5+H_3SiN(SiMe_3)_2;$

 $Si₃H₈+LiBu\rightarrow LiSi₂H₅+H₃SiBu; or$

$\begin{array}{l} \displaystyle 2\mathrm{Si}_3\mathrm{H}_8 + \mathrm{Na} [\mathrm{AlH}_2(\mathrm{OCH}_2\mathrm{CH}_2\mathrm{OMe})_2]\text{=} \mathrm{Si}_2\mathrm{H}_6 + \mathrm{SiH}_4 + \mathrm{Na} \\ \displaystyle \left[\mathrm{Al}(\mathrm{SiH}_3)(\mathrm{Si}_2\mathrm{H}_5)(\mathrm{OCH}_2\mathrm{CH}_2\mathrm{OMe})_2 \right] \end{array}$

As a result, the solid KSi₂H₅, LiSi₂H₅, or Na[Al(SiH₃) (Si₂H₅)(OCH₂CH₂OMe)₂]catalyst catalytically transforms the Si_aH_{2a+2}, with a=1-4, reactant to produce a Si_nH_(2n+2) mixture, wherein $n=1-100$.
[0253] The heterogeneous catalyst may be a mixture of

two or more catalysts. Exemplary catalyst mixtures include K and K_2O or Na and Na₂O. These catalysts are commercially available or may be synthesized by methods known in the art. For example, 12-15% Na₂O/0-3% Na/85-90% $A1_2O_3$ is sold under the trademark Activelsom™ by SiGNa Chemistry, Inc.

[0254] Any of the heterogeneous catalysts disclosed above may be located on a support. Exemplary supports include alumina (A_2O_3) , silica (SiO_2) , or combinations thereof. One of ordinary skill in the art will recognize that the catalyst may be physically and/or chemically bound to the support. For example, the catalyst may chemically react with the $-$ OH groups on the silica or alumina supports. Alternatively, the catalyst and support may simply be physically mixed together with no chemical reaction occurring. In another alternative, physically mixing the catalyst and support may result in both physical and chemical bonding. The heterogeneous catalyst may comprise between approximately 0.1% w/w to approximately 70% w/w of the total heterogeneous catalyst and support combination. Alternatively, the heterogeneous catalyst may comprise from approximately 1% w/w to approximately 50% w/w of the total heterogeneous catalyst and support combination.

[0255] In another alternative, the heterogeneous catalyst may be commercially supplied in pellet form.

[0256] As shown in the examples that follow, the claimed catalysts permit more control of the polymerization process than the prior art transition metal catalysts of Group IV (Ti, Zr , Hf), VIII (Ru), IX (Co, Rh, Ir), an homogeneous catalysts, the claimed heterogeneous catalysts may also be easily separated from the reaction mixture . As a result, the reaction is easier to scale to commercial volumes and the reaction product is easier to purify to semiconductor acceptable levels (i.e., very pure).

[0257] Catalysis of the lower silane reactants (i.e., $\text{Si}_a\text{H}_{2a+2}$, with a=1-4) occurs when the lower silane reactant contacts the heterogeneous catalyst. The reaction may occur in a batch reactor or flow through reactor. The lower silane reactant and catalyst may be mixed in a batch reactor to form a mixture. Depending on the reactant and catalyst, the catalyst-reactant mixture may be mixed for a time period ranging from approximately 1 hour to approximately 24 hours.

[0258] The batch reaction may be performed at a temperature ranging from approximately room temperature to approximately 53° C. Alternatively, the reaction may be performed at a temperature ranging from approximately 15°
C. to approximately 50° C. In another alternative, the reaction may be performed at a temperature ranging from
approximately 15° C. to approximately 30° C. One of
ordinary skill in the art will recognize that the reaction temperature will vary depending upon the selected catalyst
as well as the desired $Si_nH_(2n+2)$ product. As shown in Table 1 of Example 1, higher temperatures tend to produce heavier polysilanes (Si ≥ 6). The Si_nH_(2n+2) product may be filtered to remove solids, such as the catalyst and/or any solid Si_nH

 $(2n+2)$ product.
[0259] In a flow reactor, the $Si_aH_{(2a+2)}$ reactant may flow through a reactor containing catalyst pellets or catalyst supported on glass wool. The $Si_aH_{(2a+2)}$ reactant may have a residence time in the reactor ranging from approximately 200 second to approximately 600 seconds. The pressure in the reactor may range from approximately 10 psig (69 kPa)

to approximately 50 psig (345 kPa).
[0260] The flow reaction may be performed at a tempera-
ture ranging from approximately 15° C. to approximately
 170° C. Alternatively, the reaction may be performed at a
tempe performed at a temperature ranging from approximately 15° C. to approximately 100° C. In another alternative, the reaction may be performed at a temperature ranging from approximately 15° C. to approximately 50° C. In another alternative, the reaction may be performed at a temperature ranging from approximately 20° C. to approximately 150 $^{\circ}$ C. In another alternative, the reaction may be performed at a temperature ranging from approximately 50° C. to approximately 100° C. One of ordinary skill in the art will recognize that the reaction temperature will vary depending
upon the selected catalyst as well as the desired $Si_nH_{(2n+2)}$ product. As shown in Table 1 of Example 1, higher tem-
peratures tend to produce heavier polysilanes $(Siz6)$.

[0261] The catalyst transforms the lower silane reactant to a $\text{Si}_n\text{H}_{(2n+2)}$ mixture, wherein n=1-100. Preferably, the catalyst converts approximately 20% w/w to approximately 60% w/w of the lower silane reactant. The desired polysilane is isolated from the $Si_nH_{(2n+2)}$ mixture. When n=5-8, an isomerically enriched polysilane having a ratio of one isomer to another isomer ranging from approximately 2:1 to approximately 15:1 may be fractionally distilled from the composition comprising approximately 95% w/w to approximately 100% w/w n-Si₅H₁₂, n-Si₆H₁₄, n-Si₇H₁₆, or n- Si_8H_{18} , and preferably from approximately 98% w/w to approximately 100% w/w n-Si_sH₁₂, n-Si₆H₁₄, n-Si₇H₁₆, or n-Si₈H₁₈. $\text{Si}_n\text{H}_{(2n+2)}$ mixture to produce a Si-containing film forming

[0262] One of ordinary skill in the art will recognize that the reaction rate and product yield will vary depending on whether the lower silane reactant is substituted or not. The reaction products produced by the claimed unsubstituted lower silanes (i.e., $Si_aH_{(2a+2)}$, with a=1-4) will differ from those produced by a substituted silane containing one or more organic groups (i.e., $\text{Si}_n \text{R}_{x} \text{H}_{(2n+2-x)}$, with R is an

organic group and x is 1 or more). See Comparative Examples 1 and 2, which demonstrate that Ru/C and Rh/C are not active for transformation of non-substituted liquid or gaseous trisilane, respectively, even though U.S. Pat. No. 5,700,400 to Nippon Oil Co, Ltd., discloses the use of Ru and Rh catalyst.

[0263] The catalysis reaction may be performed in the presence or absence of unreactive gases, such as H_2 , N_2 , Ar or He . The unreactive gases may be used to maintain an inert atmosphere. The unreactive gases may also be used to dilute the reaction mixture. The unreactive gases may also be used to help maintain the flow of the reaction mixture within a desired range, for example from approximately 0.1 to approximately $1,000 \text{ mL/min}$, alternatively from approximately 1 to approximately 10 mL/min. Of course, the addition of these unreactive gases further requires their removal from the reaction product. Therefore, in another alternative and as demonstrated in the examples that follow, the catalysis reaction may be performed under the vapor pressure of the reactants.

[0264] FIG. 1 is a diagram of an exemplary batch process system for catalytic conversion of the lower silane reactants to the $\text{Si}_n\text{H}_{(2n+2)}$, wherein n=4-100, mixture. In FIG. 1, trisilane 10 and optionally disilane or tetrasilane 11 are used
as the lower silane reactants. The catalysis may be performed under an inert atmosphere, such as N_2 , a noble gas (i.e., He, Ne, Ar, Kr, Xe), or combinations thereof. Any and all air must be removed from various parts of the system (e.g., reactor 20, distillation unit 40, distillation unit 50, etc.) by applying a vacuum and/or inert gas cycles. The inert gas may also serve to pressurize the trisilane 10 and optional disilane or tetrasilane 11 to assist in delivery of the reactants to the reactor 20. Liquid nitrogen, refrigerated ethanol, an acetone/dry ice mixture, or heat transfer agents such as monoethylene glycol (MEG) or the heat transfer fluid sold under the trademark SYLTHERMTM by Dow Corning Corp.
may be used to cool various parts of the system (e.g.,

distillation set up 40, distillation set up 50).
[0265] The Si_2H_8 reactant 10 and optional Si_2H_6 or Si_4H_{10} reactant 11 are added to reactor 20 via lines 12 and 13, respectively. The reactor 20 contains the catal shown). The reactor 20 also includes a stirring mechanism (not shown), such as a paddle mixer or homogenizer. The reactor 20 may also be equipped with multiple "injection" ports," pressure gauges, diaphragm valves (not shown).
[0266] The reactor 20 and any and all components that

come into contact with the trisilane 10 and optional disilane
or tetrasilane 11 reactants and any products and by-products ("contact components") must be clean and air- and moisturefree to prevent unintended reactions and/or contamination of the polysilane product 45. The reactor 20 and other contact components must be free of any impurities that may react with or contaminate the silanes. The reactor 20 and other contact components must also be compatible with the trisilane 10 and optional disilane or tetrasilane 11 reactants and

products and by-products.

10267] Exemplary reactors 20 include stainless steel can-

isters having low surface roughness and mirror finish may be obtained by

low surface roughness and mirror finish may be obtained by mechanical polishing and/or by electropolishing. High purity may be obtained by treatments that include, but are not limited to, (a) cleaning steps using dilute acids (HF, $HNO₃$) or bases (KOH, NaOH); followed by (b) rinsing with high purity de-ionized water to ensure the complete removal of traces of the acid or base; followed by (c) drying the reactor 20. Completion of the deionized water (DIW) rinse (step b) may be indicated when the conductivity of the rinse water reaches $100 \mu s/cm$, and preferably below $25 \mu s/cm$. [0268] The drying step may include purging with an inert gas such as He, N_2 , Ar (preferably N_2 or Ar); reducing the pressure in the reactor 20 or other contact components to accelerate outgassing from the surface; heating the reactor 20 or other contact components, or any combination thereof. The drying step may comprise alternate sequences of purges,
during which a certain flow of inert gas is flown through the
vessel, and vacuuming steps. Alternatively, the drying step may be carried out by constantly flowing a purge gas while maintaining a low pressure in the reactor 20 or other contact components. The drying efficiency and end point may be assessed by measuring the trace H_2O level in the gas
emerging from the reactor 20 or other contact component.
With an inlet gas having less than 10 ppb H_2O , the outlet gas
should have a moisture content ranging from approximately 0 ppm to approximately 1 ppm, and more preferably ranging from approximately 0 ppb to approximately 200 ppb. During the purge steps and vacuum steps, heating the reactor 20 or other contact component is known to accelerate the drying time. Reactors 20 are typically maintained at a temperature ranging from approximately 40° C. to approximately 150 $^{\circ}$ C. during drying.

[0269] Once cleaned and dried, the reactor 20 must have a total leak rate below 1×10^{-6} std cm³/s, preferably $\leq 1 \times 10^{-8}$ std cm³/s.

 $[0270]$ Any gases used to prepare the system for catalysis or during the catalysis process must be of semiconductor grade (i.e. free of contaminants such as trace moisture and oxygen (<1 ppm, preferably <10 ppb), and particles (<5 particles per litre $(\partial_0 0.5 \mu m)$).

[0271] The reactor 20, the source vessels of trisilane 10 and optional disilane or tetrasilane 11, the polysilane product containers, and any other contact components may also be
passivated by exposure to a silylating agent such as silane,
disilane, or trisilane prior to the reaction. Passivation helps
minimize reaction between the lower or

Alternatively, the $Si₃H₈$ reactant 10 and optional $Si₂H₆$ or Si_4H_{10} reactant 11 may be directly introduced into reactor 20 via lines 12 and 13 (not shown). The Si_3H_8 reactant 10 and optional $\rm Si_2H_6$ or $\rm Si_4H_{10}$ reactant 11 may be added to the reactor 20 via a liquid metering pump (not shown), such as a diaphragm pump, peristaltic pump, or syringe pump.

[0273] Upon completion of the addition of the $Si₃H₈$ reactant 10 and optional $Si₂H₆$ or $Si₄H₁₀$ reactant 11, the reactor 20 may be heated to a temperature ranging from approximately 25 \degree C. to approximately 150 \degree C. or alternatively from approximately 15° C. to approximately 100° C.
The reactor 20 may be maintained at the desired temperature by a jacket (not shown). The jacket may have an inlet and outlet (not shown). Inlet and outlet may be connected to a heat exchanger/chiller (not shown) and/or pump (not shown) to provide recirculation of a heating or cooling fluid. Alternatively, the temperature of the reactor 20 may be maintained using heating tape (not shown) or a heating mantle (not shown), with the heating elements connected to a

temperature controlling unit (not shown). A temperature sensor (not shown) may be used to monitor the temperature of the contents of the reactor 20 .

[0274] The lower silane reactant and catalyst may be stirred for a time period ranging from approximately 0.1 hour to approximately 72 hours, alternatively from approximately 1 hour to approximately 30 hours. The mixing ma example, gas chromatography. The predominant reaction products are SiH_4 , $Si₅H₁₂$, etc.

[0275] Upon completion of the reaction, the reactor 20 is cooled to approximately room temperature. When the reactor 20 is jacketed, any heating fluid may be replaced with a cooling fluid to assist in cooling the reactor 20 and its contents. Liquid nitrogen, refrigerated ethanol, an acetone/ dry ice mixture, or heat transfer agents may be used to cool the reactor 20. Alternatively, any heating mechanism, such as heating tape or a heating mantel, may be turned off and natural cooling may occur. Any heavier liquid non-volatile silanes 23 are filtered from the catalyst and solid reaction products and removed from the reactor 20 via line 22. The volatile silanes 21 are stripped from reactor 20 by pressure difference .

[0276] The volatile silanes 21 may be collected in one or more traps 30 to produce a $\text{Si}_{n}\text{H}_{(2n+2)}$ mixture 31, wherein n=1-100. Exemplary traps 30 include a dry ice/isopropanol, dry ice/acetone, refrigerated ethanol, and/or liquid nitrogen trap.

[0277] The $\text{Si}_n\text{H}_{(2n+2)}$ mixture 31 may be collected in one or more containers and transported to a new location prior to performance of the next process steps. Alternatively, the mixture 31 may immediately be directed to a distillation unit 40 to further isolate the reaction product from any reactants and reaction by-products. The distillation unit 40 separates the desired polysilane product 45 from the $SiH₄$ reaction by - product 43, the volatile $\text{Si}_n\text{H}_{2n+2}$ with n≥5 reaction by products 44, and any unreacted Si_3H_8 reactant 41 and unreacted optional $Si₂H₆$ or $Si₄H₁₀$ reactant 42. The unreacted $Si₃H₈$ reactant 41 and unreacted optional $Si₂H₆$ or $Si₄H₁₀$ reactant 42 may be recycled for use in future processes .

[0278] Once again, the polysilane product 45 may be transported to a new location prior to performance of the next process steps. Alternatively, the polysilane product 45 may be directed to a fractional distillation unit 50 to separate the n-isomer 51 from other isomers 52. The fractional distillation may be performed using a static column or a spinning band column. The length of the spinning band distillation column is much smaller than that of the static column and may be preferred for use in crowded facilities because it takes up less space. A static column suitable to produce approximately 90% n-tetrasilane would require between approximately 90 to approximately 120 theoretical plates and would be approximately 6 to 7 meters tall.

[0279] FIG. 2 is a diagram of the flow process for catalytic conversion of the lower silane reactants to the $Si_nH_{(2n+2)}$ mixture. The same references numbers from FIG. 1 have been used for the same components in FIG. 2. all of the contact components of FIG. 2 must be clean and air- and moisture-free. As in FIG. 1, the catalysis of FIG. 2 may be performed under an inert atmosphere, such as N_2 , a noble gas (i.e., He, Ne, Ar, Kr, Xe), or combinations thereof.

[0280] Trisilane 10 and optionally disilane or tetrasilane 11 are added to flow reactor 25 via lines 12 and 13, respectively. As in FIG. 1, the Si₃H₈ reactant 10 and optional Si_2H_6 or Si_4H_{10} reactant 11 may be mixed in line 14 before introduction into flow reactor 25. Alternatively, the Si_3H_8 reactant 10 and optional Si_2H_6 or Si_4H_{10} reactant 11 may be directly introduced into flow reactor 25 via lines 12 and 13 (not shown). The Si_3H_8 reactant 10 and optional Si_2H_6 or $Si₄H₁₀$ reactant 11 may be added to the flow reactor 25 via a liquid metering pump (not shown), such as a diaphragm pump, peristaltic pump, or syringe pump. Preferably, the mixing is performed under an inert atmosphere at approximately atmospheric pressure.

[0281] As will be provided in further detail in discuss of FIG. 4 below, the catalyst (not shown) is located within the flow reactor 25. The flow reactor 25 is maintained at a temperature ranging from approximately 25° C. to approximately 250° C., alternatively from approximately 40° C. to approximately 250° C. or, in another alternative, from approximately 50° C. to approximately 100° C. The temperature selected will depend upon the catalyst selected, as well as the target reaction products. The flow reactor 25 is maintained at a pressure ranging from approximately 0.1 atm to approximately 10 atm. The flow of the trisilane 10 and optionally disilane or tetrasilane 11 reactants is selected to provide approximately 0.01 to approximately 100 minutes of residence time in flow reactor 25, alternatively between approximately 2 minutes to approximately 20 minutes residence time, alternatively between approximately 1 second to approximately 1,000 seconds or, in another alternative, from approximately 100 seconds to approximately 600 seconds.

[0282] The $Si_nH_{(2n+2)}$ mixture 26, wherein n=1-100, is collected in a receiver 35 after passing the flow reactor 25. The receiver 35 may be any sort of trap, including but not limited to dry ice/isopropanol, dry ice/acet

[0283] As in FIG. 1 above, the $\text{Si}_n H_{(2n+2)}$ mixture 31 may be collected in one or more containers and transported to a new location prior to performance of the next process steps. Alternatively, the mixture 31 may immediately be directed to a distillation unit 40 to further isolate the reaction product from any reactants and reaction by-products. The distillation unit 40 separates the desired polysilane product 45 from the SiH₄ reaction by-product 43, the volatile Si_nH_{2n+2} with n≥5 reaction by -products 44, and any unreacted $\overline{Si}_3\overline{H}_8$ reactant 41 and optional \overline{Si}_2H_6 or \overline{Si}_4H_{10} reactant 42. The unreacted Si_3H_8 reactant 41 and unreacted optional Si_2H_6 or Si_4H_{10} reactant 42 may be recycled. Real time analysis and purification of the unreacted Si_3H_8 reactant 41 and unreacted optional Si_2H_6 or Si_4H_{10} react

[0284] Once again, the polysilane product 45 may be transported to a new location prior to performance of the next process steps. Alternatively, the polysilane product 45 may be directed to a fractional distillation unit 50 to separate the n-isomer 51 from other isomers 52 . The fractional distillation may be formed with a static column or a spinning band column. The spinning band distillation column length is much smaller than that of the static column and may be preferred for use in crowded facilities beca less space. A static column suitable to produce approximately 90% n-tetrasilane would require between approximately 90 to approximately 120 theoretical plates and would

[0285] FIG. 3 is a diagram of the flow reactor 20 of FIG.
2. Please note that valves have not been included in this

figure to make the figure easier to read.
[0286] The $\text{Si}_a\text{H}_{(2a+2)}$ reactant 100 is pressurized with nitrogen in order to supply the $\mathrm{SL}_a\mathrm{H}_{(2a+2)}$ reactant to the flow reactor 120 via line 102. Line $10\overline{2}$ is also connected to vacuum 110. A flow regulator 101 controls the flow of the $Si_aH_{(2a+2)}$ reactant. The flow regulator 101 may be a graduated needle valve, electronic flow meter, etc. A gauge $103a$ measures the pressure and may communicate with the flow
regulator 101 to adjust accordingly.

[0287] Flow reactor 120 includes two thermocouples 121 and 122. More or fewer may be used without departing from the teachings herein. Exemplary thermocouples suitable for use in the teaching herein include Type K or Type J thermocouples.

[0288] The Si_nH_(2n+2) reaction mixture exits the flow reactor 120 via line 123. Pressure regulator 104 sets the pressure in the reactor 120 and provides the pressure differential that moves the Si_nH_(2n+2) reaction 120 to the dry ice/isopropanol trap 130. Gauge 103b indicates the pressure in the reactor 120. The dry ice/isopropanol cates the pressure in the reactor 120. The dry ice isopropanol trap 130 captures any $\text{Si}_{n}H_{(2n+2)}$ reaction products that condense above approximately -78° C.

[0289] Any volatile $\text{Si}_n H_{(2n+2)}$ reaction mixture that is not captured in the dry ice/isopropanol trap is condensed via line 131 to a liquid nitrogen trap 140. The liquid nitrogen trap 140 captures any Si_nH_(2n+2) reaction products that condense below approximately -78° C. and approximately -196° C. Line 131 is also connected to vacuum line 110. Pressure gauge $103c$ monitors pressure in line 131. SiH₄ by-product is sent to an exhaust scrubber (not shown) via line 150. N₂ 105 is used to dilute the SiH_4 by-product on its way to the exhaust scrubber. Check valve 106 prevent backflow of this

pyrophoric by-product.

[0290] FIG. 4 is a diagram of the flow reactor 120 of FIG.

3. In FIG. 4, valves 201 permit the stainless steel tube flow

reactor 220 to be accessed for troubleshooting or preventative maintenance. The stainless steel tube flow reactor 220 includes two thermocouples 221 and 222. As in FIG. 3, more or fewer thermocouples may be used without departing from the teachings herein. Glass wool 202 is located at the beginning and the end of the stainless steel tube flow reactor 220. The catalyst (not shown) may be packed between the glass wool 202 located at the beginning and end of the reactor or located on glass wool (not shown) packed between the glass wool 202 at the beginning and end of the flow reactor 220. As a result, the $\text{Si}_n\text{H}_{(2n+2)}$ reactant may be heated prior to catalysis when it passes through the glass
wool at the beginning of the flow reactor 220. One of
ordinary skill in the art will recognize that a layer of glass
beads and pellet catalysts may be used in plac

the stainless steel tube flow reactor 220. Thermal insulation 204 helps to maintain the temperature of the stainless steel tube flow reactor 220. One of ordinary skill in the art will recognize that alternative heating means may also be used
without departing from the teachings herein. For example,
the stainless steel tube flow reactor 220 may alternatively be placed in an oven (not shown). In that embodiment, thermal insulation 204 would not be needed.

[0292] One of ordinary skill in the art will recognize the sources for the equipment components of the systems used to practice the disclosed methods. Some level of customization of the components may be required based upon the desired temperature range, pressure range, local regulations, etc. Exemplary equipment suppliers include Parr Instrument Company equipment and components made from stainless steel.

[0293] Fractional distillation of the desired polysilane
product (50 in FIGS. 1 and 2) produces a Si-containing film
forming composition comprising between approximately
90% w/w to approximately 100% w/w n-Si_sH₁₂, n-S 95% w/w to approximately 100% w/w n- Si_5H_{12} , n- Si_6H_{14} , n- Si_7H_{16} , or n- Si_8H_{18} , and more preferably between approximately 97% w/w to approximately 100% w/w n- Si_5H_{12} , n- Si_6H_{14} , n- Si_7H_{16} , or n- Si_8H_{18} . The Si-containing film forming compositions further comprises between approximately 0% w/w of non n-isomers, preferably between approximately 0% w/w to approximately 5% w/w non n-isomers; and more preferably between approximately 0% w/w to approximately 3% w/w non n-isomers. For example, after fractional distillation of approximately 192 grams of a 3:1 n-Si₄H₁₀:i-Si₄H₁₀ mixture using a 1 cm diameter and 100 cm long spinning band distillation column, Applicants have been able to produce approximately 90% \overline{w}/w to approximately 95% w/w n-tetrasilane. One of ordinary skill in the art will recognize that higher purity n-tetrasilane would be obtained from mixtures having higher n-Si₄H₁₀:i-Si₄H₁₀ ratio and/or larger distillation columns.

[0294] The Si-containing film forming composition has a purity ranging from approximately 97% mol/mol to approximately 100% mol/mol, more preferably mol/mol to approximately 100% mol/mol, more preferably from approximately 99.5% mol/mol to approximately 100% mol/mol, and even more preferably from approximately

99.97% mol/mol to approximately 100% mol/mol.
[0295] The Si-containing film forming compositions preferably comprise between the detection limit and 100 ppbw of each potential metal contaminant (e.g., at least Ag, Al, Au,

[0296] The concentration of X (wherein X=Cl, Br, or I) in
the Si-containing film forming compositions may range
from approximately 0 ppmw to approximately 100 ppmw,
and more preferably from approximately 0 ppmw and to
app

confirmed by ¹H and/or ²⁹Si NMR.
[0298] As discussed in detail above and illustrated in the examples that follow, the Si-containing film forming composition must be stored in a clean dry storage vessel with which it does not react in order to maintain its purity.

[0299] The advantages of the disclosed synthesis methods are as follows:

- [0300] Lower process temperature and higher yield of the desired polysilane compared to pyrolysis process, which helps reduce cost and product isolation issues:
- [0301] Application of heterogeneous catalysts essentially eliminates the issues with the contamination of

reaction products with the catalyst , and eliminates steps of purification of the reaction products from the

- residual catalyst; [0302] Application of heterogeneous catalysts allows
better control of the reaction time, while the reported homogeneous catalysts continue polymerizing trisilane
- until quenched, mostly in the non-controlled fashion;
[0303] The process is solventless;
- $[0.903]$ The process is solventless,
- [0304] Purification only by distination, [0305] The waste generation is minimal and environ-
mentally benign; and
-

[0306] Many of the starting materials are inexpensive and readily available.
[0307] All of the above are advantageous from the stand-
point of developing a scalable industrial process. As a result, the reaction product maintains purity levels suitable for the semiconductor industry without the use of stabilizers, such as toxic organic amines which may be needed to quench

reactions using homogeneous catalysts.
[0308] Also disclosed are methods of using the disclosed
Si-containing film forming compositions for vapor deposi-
tion methods. The disclosed methods provide for the use of
the Si-co for fabrication of electronic or optoelectronic devices or circuits. The disclosed methods may be useful in the manu-
facture of semiconductor, photovoltaic, LCD-TFT, or flat panel type devices. The method includes: introducing the vapor of the disclosed Si-containing film forming compositions into a reactor having a substrate disposed therein and
depositing at least part of the disclosed Si-containing film
forming composition onto the substrate via a deposition

process to form a Si-containing layer.

[0309] The disclosed methods also provide for forming a bimetal-containing layer on a substrate using a vapor deposition process and, more particularly, for deposition of SiMO, or Si Nb, V, Hf, Zr, Ti, Al, B, C, P, As, Ge, lanthanides (such as Er), or combinations thereof.

[0310] The disclosed methods of forming silicon-containing layers on substrates may be useful in the manufacture of semiconductor, photovoltaic, LCD-TFT, or flat panel type devices. The disclosed Si-containing film forming composition may deposit Si-containing films using any vapor deposition methods known in the art. Examples of suitable vapor deposition methods include chemical vapor deposi tion (CVD) or atomic layer deposition (ALD). Exemplary CVD methods include thermal CVD , plasma enhanced CVD (PECVD), pulsed CVD (PCVD), low pressure CVD (LP-CVD), sub-atmospheric CVD (SACVD), atmospheric pressure CVD (APCVD), flowable CVD (f-CVD), metal organic chemical vapor deposition (MOCVD), hot-wire CVD (HW-CVD, also known as cat-CVD, in which a hot wire serves as an energy source for the deposition process), radicals incorporated

[0311] CVD, and combinations thereof. Exemplary ALD methods include thermal ALD, plasma enhanced ALD (PEALD), spatial isolation ALD, hot-wire ALD (HWALD), radicals incorporated ALD, and combinations thereof. Super
critical fluid deposition may also be used. Among these,
thermal CVD deposition is preferred for a process in which a high deposition rate, excellent film uniformity, and conformal film quality are required. Thermal ALD deposition is preferred for a process that forms a film having the high uniformity under severe conditions (e.g., trenches, holes, or vias). In one alternative, a PECVD deposition is preferred, particularly when fast growth, conformality, process-orientation and one direction films are require alternative, a PEALD deposition process is preferred, particularly when superior conformality of films deposited on challenging surfaces (e.g., trenches, holes, and vias) is required.

[0312] The vapor of the Si-containing film forming composition is introduced into a reaction chamber containing a substrate. The temperature and the pressure within the reaction chamber and the temperature of the substrate are held at conditions suitable for vapor deposition of at least part of the Si-containing film forming composition onto the substrate. In other words, after introduction of the vaporized composition into the chamber, conditions within the chamber are such that at least part of the vaporized precursor deposits onto the substrate to form the silicon-containing
film. A co-reactant may also be used to help in formation of
the Si-Containing Layer.
[0313] The reaction chamber may be any enclosure or
chamber of a device in wh

such as, without limitation, a parallel-plate type reactor, a cold-wall type reactor, a hot-wall type reactor, a single-wafer
reactor, a multi-wafer reactor, or other such types of deposition systems. All of these exemplary reaction chambers are capable of serving as an ALD reaction chamber. The reaction chamber may be maintained at a pressure ranging from about 0.5 m Torr to about 760 Torr. In addition, the temperature within the reaction chamber may range from about 20° C. to about 700° C. One of ordinary skill in the art will recognize that the temperature may be optimized through mere experimentation to achieve the desired result.

[0314] The temperature of the reactor may be controlled by either controlling the temperature of the substrate holder and/or controlling the temperature of the reactor wall. Devices used to heat the substrate are known in the art. The reactor wall may be heated to a sufficient temperature to obtain the desired film at a sufficient growth rate and with desired physical state and composition. A non-limiting
exemplary temperature range to which the reactor wall may
be heated includes from approximately 20° C. to approxi-
mately 700° C. When a plasma deposition process is a thermal process is performed, the deposition temperature may range from approximately 300° C. to approximately 700° C.

[0315] Alternatively, the substrate may be heated to a sufficient temperature to obtain the desired silicon-containing film at a sufficient growth rate and with desired physical state and composition. A non-limiting exemplary temperature range to which the substrate may be heated includes from 150° C. to 700° C. Preferably, the temperature of the

substrate remains less than or equal to 500° C.
[0316] The type of substrate upon which the siliconcontaining film will be deposited will vary depending on the final use intended . A substrate is generally defined as the material on which a process is conducted. The substrates include, but are not limited to, any suitable substrate used in
semiconductor, photovoltaic, flat panel, or LCD-TFT device manufacturing. Examples of suitable substrates include wafers, such as silicon, silica, glass, Ge, or GaAs wafers. The wafer may have one or more layers of differing mate-

rials deposited on it from a previous manufacturing step. For example, the wafers may include silicon layers (crystalline, amorphous, porous, etc.), silicon oxide layers, silicon nitride layers, silicon oxy nitride layers, carbon doped silicon oxide (SiCOH) layers, or combinations thereof. Additionally, the wafers may include copper layers, tungsten layers or metal layers (e.g. platinum, palladium, nickel, rhodium, or gold). The wafers may include barrier layers, such as manganese, manganese oxide, tantalum, tantalum nitride, etc. The layers may be planar or patterned. In some embodiments, the substrate may be coated with a patterned photoresist film. In some embodiments, the substrate may include layers of oxides which are used as dielectric materials in MIM, DRAM, or FeRam technologies (for example, ZrO₂ based materials, $HfO₂$ based materials, $TiO₂$ based materials, rare earth oxide based materials, ternary oxide based materials, etc.) or from nitride-based films (for example, TaN) that are used as electromigration barrier and adhesion layer between copper and the low-k layer. The disclosed processes may deposit the silicon-containing layer directly on the wafer or directly on one or more than one (when patterned layers form the substrate) of the layers on top of the wafer.
Furthermore, one of ordinary skill in the art will recognize
that the terms "film" or "layer" used herein refers to a thickness of some material laid on or spread over a surface the specification and claims, the wafer and any associated layers thereon are referred to as substrates. The actual substrate utilized may also depend upon the specific precur sor embodiment utilized. In many instances though, the preferred substrate utilized will be selected from hydrogenated carbon, TiN, SRO, Ru, and Si type substrates, such as polysilicon or crystalline silicon substrates.
[0317] The substrate may be patterned to include vias or

trenches having high aspect ratios. For example, a conformal Si-containing film, such as SiN or SiO₂, may be deposited

using any ALD technique on a through silicon via (TSV)
having an aspect ratio ranging from approximately $20:1$ to
approximately $100:1$.
 $[0318]$ When n=4-10, the Si-containing film forming com-
positions may be supplied be selected from, among others, C1-C16 saturated or unsaturated hydrocarbons.

[0319] For vapor deposition, the Si-containing film forming compositions are introduced into a reactor in vapor form
by conventional means, such as tubing and/or flow meters. The vapor form may be produced by vaporizing the Sicontaining film forming compositions through a conven tional vaporization step such as direct liquid injection, direct vapor draw in the absence of a carrier gas, by bubbling a carrier gas through the liquid, or by sweeping the vapors
with a carrier gas without bubbling through the liquid. The
Si-containing film forming compositions may be fed in liquid state to a vaporizer (Direct Liquid Injection) where it is vaporized and mixed with a carrier gas before it is introduced into the reactor. Alternatively, the Si-containing film forming compositions may be vaporized by passing a carrier gas into a container containing the composition or by bubbling the carrier gas into the composition. The carrier gas may include, but is not limited to, Ar, He, or N_2 , and mixtures thereof. The carrier gas and composition are then introduced into the reactor as a vapor.

[0320] If necessary, the Si-containing film forming composition may be heated to a temperature that permits the Si-containing film forming composition to have a sufficient vapor pressure. The delivery device may be maintained at temperatures in the range of, for example, $0-150^{\circ}$ C. Those skilled in the art recognize that the temperature of the delivery device may be adjusted in a known manner to control the amount of Si-containing film forming composition vaporized.

[0321] In addition to the disclosed composition, a reaction gas may also be introduced into the reactor . The reaction gas may be an oxidizing agent such as O_2 ; O_3 ; H_2O ; H_2O ; N_2O ; oxygen containing radicals such as O- or OH-; NO; $NO₂$; carboxylic acids such as formic acid, acetic acid, propionic acid; radical species of NO, $NO₂$, or the carboxylic acids; para-formaldehyde; and mixtures thereof. Preferably, the oxidizing agent is selected from the group consisting of O_2 , O_3 , H_2O , H_3O_2 , oxygen containing radicals thereof such as O- or OH-, and mixtures thereof. Preferably, when an ALD process is performed, the co-reactant is plasma treated oxygen, ozone, or combinations thereof. When an oxidizing gas is used, the resulting silicon containing film will also contain oxygen.

[0322] Alternatively, the reaction gas may H_2 , NH_3 ,

(SiH₃)₃N, hydridosilanes (such as SiH₄, Si₂H₆, Si₃H₈, Si₄H₁₀, Si₅H₁₀, Si₄H₁₂), chlorosilanes and chloropolysilanes (such as SiHCl₃, SiH₂Cl₂, SiH₃Cl₅, Si₂HCl₅, Si₃Cl₈), alkylsila NEt_2H , NMe₃, NEt₃, (SiMe₃)₂NH), diamines such as eth-ylene diamine, dimethylethylene diamine, pyrazoline, pyridine, B-containing molecules (such as B_2H_6 , trimethylboron, triethylboron, borazine, substituted borazine, dialkylaminoboranes), alkyl metals (such as trimethylaluminum, triethylaluminum, dimethylzinc,

diethylzinc), radical species thereof, or mixtures thereof.
When H_2 or an inorganic Si containing gas is used, the
resulting silicon containing film may be pure Si.
[0323] Alternatively, the reaction gas may be a hydro tene, pentyne, cyclopentane, butadiene, cyclobutane, terpinene, octane, octene, or combinations thereof.

[0324] The reaction gas may be treated by a plasma, in order to decompose the reaction gas into its radical form. N_2 may also be utilized as a reducing agent when treated with plasma. For instance, the plasma may be generated with a power ranging from about 50 W to about 500 W, preferably from about 100 W to about 200 W. The plasma may be generated or present within the reactor itself. Alternatively, the plasma may generally be at a location removed from the reactor, for instance, in a remotely located plasma system. One of skill in the art will recognize methods and apparatus
suitable for such plasma treatment.
[0325] The desired silicon-containing film also contains

another element, such as, for example and without limitation, B, P, As, Zr, Hf, Ti, Nb, V, Ta, Al, Si, or Ge.

[0326] The Si-containing film forming composition and one or more co-reactants may be introduced into the reaction chamber simultaneously (chemical vapor deposition), sequentially (atomic layer deposition), or in other combinations. For example, the vapor of the Si-containing film

forming composition may be introduced in one pulse and two additional metal sources may be introduced together in
a separate pulse (modified atomic layer deposition). Alternatively, the reaction chamber may already contain the co-reactant prior to introduction of the Si-containing film
forming composition. The co-reactant may be passed through a plasma system localized within or remote from the reaction chamber, and decomposed to radicals. Alternatively, the Si-containing film forming composition may be introduced to the reaction chamber continuously while other
precursors or reactants are introduced by pulse (pulsedchemical vapor deposition). In another alternative, the Sicontaining film forming composition and one or more coreactants may be simultaneously sprayed from a shower head under which a susceptor holding several wafers is spun (spatial ALD).

[0327] In one non-limiting exemplary atomic layer deposition process, the vapor phase of the Si-containing film forming composition is introduced into the reaction chamber, where it is contacted with a suitable substrate. Excess composition may then be removed from the reaction cham ber by purging and/or evacuating the reaction chamber. An oxygen source is introduced into the reaction chamber where it reacts with the absorbed Si - containing film forming com position in a self-limiting manner. Any excess oxygen source is removed from the reaction chamber by purging and/or evacuating the reaction chamber. If the desired film is a silicon oxide film, this two-step process may provide the desired film thickness or may be repeated until a film having the necessary thickness has been obtained.

[0328] Alternatively, if the desired film is a silicon metal/ metalloid oxide film (i.e., SiMO_x, wherein x may be 0-4 and M is B, Zr, Hf, Ti, Nb, V, Ta, Al, Si, Ga, Ge, or combinations thereof), the two-step process above may be followed by
introduction of a vapor of a metal- or metalloid-containing
precursor into the reaction chamber. The metal- or metalloid-containing precursor will be selected based on the nature of the silicon metal/metalloid oxide film being deposited. After introduction into the reaction chamber, the metalor metalloid-containing precursor is contacted with the substrate. Any excess metal- or metalloid-containing pre-
cursor is removed from the reaction chamber by purging and/or evacuating the reaction chamber. Once again, an oxygen source may be introduced into the reaction chamber
to react with the metal- or metalloid-containing precursor. Excess oxygen source is removed from the reaction chamber
by purging and/or evacuating the reaction chamber. If a desired film thickness has been achieved, the process may be terminated. However, if a thicker film is desired, the entire four-step process may be repeated. By alternating the provision of the Si-containing film forming composition, metal-
or metalloid-containing precursor, and oxygen source, a film

of desired composition and thickness can be deposited.
[0329] Additionally, by varying the number of pulses, films having a desired stoichiometric M:Si ratio may be obtained. For example, a SiMO_2 film may be obtained having one pulse of the Si-containing film forming composition and one pulse of the metal- or metalloid-containing precursor, with each pulse being followed by a pulse of the oxygen source. However, one of ordinary skill in the art will recognize that the number of pulses required to obtain the desired film may not be identical to the stoichiometric ratio of the resulting film .

[0330] The silicon-containing films resulting from the processes discussed above may include $SiO₂$; SiC ; SiN ; SiON; SiOC; SiONC; SiBN; SiBCN; SiCN; SiMO, SiMN in which M is selected from Zr, Hf, Ti, Nb, V, Ta, Al, Ge, depending of course on the oxidation state of M. One of ordinary skill in the art will recognize that by judicial selection of the appropriate Si-containing film forming composition and co-reactants, the desired film compo

[0331] Upon obtaining a desired film thickness, the film may be subject to further processing, such as thermal annealing, furnace-annealing, rapid thermal annealing, UV or e-beam curing, and/or plasma gas exposure. Those skilled in the art recognize the systems and methods utilized to perform these additional processing steps. For example, the silicon-containing film may be exposed to a temperature ranging from approximately 200° C. and approximately 1000° C. for a time ranging from approximately 0.1 second to approximately 7200 seconds under an inert atmosphere, a H-containing atmosphere, a N-containing atmosphere, or combinations thereof. Most preferably, the temperature is 600° C. for less than 3600 seconds. Even more preferably, the temperature is less than 400° C. The annealing step may
be performed in the same reaction chamber in which the
deposition process is performed. Alternatively, the substrate
may be removed from the reaction chamber, wit ing/flash annealing process being performed in a separate apparatus. Any of the above post-treatment methods, but especially UV-curing, has been found effective to enhance the connectivity and cross linking of the film, and to reduce the H content of the film when the film is a SiN containing film. Typically, a combination of thermal annealing to $\leq 400^\circ$ C. (preferably about 100° C.-300° C.) and UV curing is used to obtain the film with the highest density.

[0332] The disclosed Si-containing film forming compositions may also be used in coating deposition processes to form silicon, silicon nitride, silicon oxide, silicon carbide, or silicon oxynitride films used in the electronics and optics industry. The silicon oxide films are obtained from thermal treatment of the deposited film under an oxidative atmo-
sphere, containing at least one of O_2 , O_3 , H_2O , H_2O_2 , NO, N_2O , and combinations thereof. The disclosed Si-containing film forming compositions may also be used to form protective coatings or pre-ceramic materials (i.e., nitrides and oxynitrides) for use in the aerospace, auto

[0333] For coating processes, the Si-containing film forming composition preferably comprises an isolated Si_nH_(2n+2) compound or a Si_nH_(2n+2) mixture, wherein n=10-100, preferably from 10-30 or 30-50. The Si_nH_(2n+2) mixture may have a Mn ranging from approximately 400 Da to approximately 1000 Da, a Mw ranging from approximately approximately 1 to approximately 10.
[0334] The Si-containing film forming compositions used

in coating processes may further comprise a solvent or solvent system having different boiling points in order to adjust the coating composition's properties, such as viscosity or layer thickness. Exemplary solvents include hydrocarbons such as benzene, toluene, xylene, mesitylene, or n-hexane; ketones, such as methylethylketone, cyclohexanone, or 2-heptanone; ethers, such as ethyl ether, dibutyl ether, or tetrahydrofuran; silanes such as m-tolyl

silane, o-tolyl silane, p-tolyl silane, p-ethylphenyl silane, m-ethylphenyl silane, o-ethylphenyl silane, m-xylene, o-xylene, o-xylene, or combinations thereof; and amines, such as pyridine, xylene, or methyl pyridine; est thereof. An exemplary solvent system may contain one solvent that boils at lower temperatures, with a boiling point (BP) between 30 $^{\circ}$ C. and 100 $^{\circ}$ C., such as pentane, hexane, benzene, diethylether, methylethylether, cyclohexane, acetone etc. The solvent system may also comprise a second solvent may have a higher boiling point, isobutyl ketone, cyclohexanones, cyclopentanone, glycols,
etc. To be suitable for coating methods, the Si-containing
film forming composition should have a molecular weight
ranging from approximately 500 to approximately 1 approximately 80% w/w to approximately 99% w/w, and more preferably from approximately 85% w/w to approxi-

more preferably from approximately 85% w/w to approximately 95% w/w.

[0335] The Si-containing film forming composition may

further comprise a perhydropolysilazane. One particularly

preferred perhydropolysilazane is disc

factants, pigments, UV absorbers, pH adjusters, surface modifiers, plasticizers, dispersing agents, catalysts, and combinations thereof. The catalyst may be the same or different from the catalyst used to synthesize the Si-containing film forming composition. Exemplary catalysts may be selected to facilitate further densification of the Si-containing film forming composition in subsequent processing steps
by catalyzing desilylative coupling (DSC), cross-linking, or
H₂ elimination. Such catalysts should be selected for their low activity at room temperature to keep the composition stable upon storage, and only induce reactions when heated to a temperature higher than room temperature, and ideally between 50 $^{\circ}$ C. and 200 $^{\circ}$ C. For instance, P(Ph)₃, P(Tolyl)₃ or metal carbonyl may be suitable catalysts for high tem toactive materials that will induce further cross-linking upon exposure to photons, such as photo-acid generators and photoinitiators such as radical initiators, cationic initiators, anionic photoinitiators, like mono-aryl ketones, trimethyl-
benzoyldiphenyl phosphinates, and/or phospine oxides.
[0337] The catalyst may also promote the conversion of
the Si-containing film forming composition to silica.

[0338] The Si-containing films may be deposited using any coating methods known in the art. Examples of suitable coating methods include spin coating, dip coating, spray
coating, fiber spinning, extrusion, molding, casting, impregnation, roll coating, transfer coating, slit coating, etc. For
usage in non-semiconductor applications, t containing film forming compositions may also contain a ceramic filler, such as BN, SiN, SiCN, SiC, Al_2O_3 , ZrO_2 ,

 Y_2O_3 , and/or Li₂O powders. The coating method is preferably spin coating in order to provide suitable film thickness control and gapfill performance.

[0339] The disclosed Si-containing film forming compositions may be applied directly to the center of the substrate and then spread to the entire substrate by spinning or may be applied to the entire substrate by spraying. When applied directly to the center of the substrate, the substrate may be spun to utilize centrifugal forces to evenly distribute the composition over the substrate. One of ordinary skill in the art will recognize that the viscosity of the Si-containing film forming compositions will contribute as to whether rotation
of the substrate is necessary. Alternatively, the substrate may be dipped in the disclosed Si-containing film forming compositions. The resulting films may be dried at room temperature for a period of time to vaporize the solvent or volatile components of the film or dried by force-drying or baking or by the use of one or a combination of any following suitable process including thermal curing and irradiations, such as, ion irritation, electron irradiation, UV and/or visible light irradiation, etc.

[0340] The spin-on Si-containing film forming compositions may also be used for the formation of transparent silicon oxynitride films suitable for optics applications. [0341] When used for spin coating, dip coating or spra

silicon nitride barrier layers that are useful as moisture or oxygen barriers, or as passivation layers in displays, light

extern parameters and photovoltaic devices.
 [0342] In semiconductor applications the Si-containing film forming compositions may be used for forming sacrificial layers such as etching hard masks, ion implantation masks, anti-reflective coatings, tone inversion layers. Alternatively, the Si-containing film forming compositions may be used for forming non-sacrificial layers ("leave behind" films), such as gapfill oxide layer, pre-metal dielectric layers, transistor stressing layers, etch stop layers, inter-layer

dielectric layers,

[0343] For Gap-fill applications, the trench or hole may

have an aspect ratio ranging from approximately 0.5:1 to

approximately 100:1. The Si-containing film forming compositions is typically spun on the substrate, pre-baked at 50° C -300 $^{\circ}$ C. to evaporate the solvent(s), and eventually converted to silicon oxide by annealing the substrate in an oxidizing atmosphere, typically containing O_2 , O_3 , H_2O , H_2O_2 , N_2O , NO, at a temperature ranging from 300 to 1000° C. The oxide quality may be improved by a multi-step annealing process in various atmospheres (oxidative or inert).

[0344] FIG. 5 is a flow chart diagraming exemplary processes for the preparation of the Si-containing film forming compositions, preparation of the silicon substrate, and the steps of a spin-coating process. One of ordinary skill in the art will recognize that fewer or additional steps than those provided in FIG. 5 may be performed without departing from the teachings herein. For example, the characterization step utilized in a R&D setting may not be required in commercial operations . One of ordinary skill in the art will under an inert atmosphere to prevent undesired oxidation of the film and/or in a clean room to help prevent particle contamination of the film.

[0345] In Step A, the desired $\text{Si}_{n}H_{(2n+2)}$ product, wherein $n=10-30$ or $n=30-50$, is mixed with the solvent to form a 1-50 wt $%$ mixture. Mixing mechanisms known in the art may be used to mix these two components (e.g., mechanical stirring, mechanical shaking, etc.). Depending on the ingredients, the mixture may be heated to a temperature ranging from 27° C. to approximately 100° C. The heating temperature should always remain lower than the pre-baking temperature. Depending on the specific ingredients, mixing may occur for 1 minute to 1 hour.

[0346] In Step B, the optional catalyst, optional perhy-
dropolysilane, such as that disclosed in US Pat App No
2018/072571, or both may be added to the mixture and mechanically stirred in the same manner. Depending on the ingredients, the mixture may be heated to a temperature ranging from 27° C. to approximately 100° C. Depending on the specific ingredients, mixing may occur for 1 minute to 1 hour.

[0347] In Optional Step C, the mixture may be aged to allow any reaction between the additives to reach equilibrium. After mixing, the mixture may age for 1 hour to 2 weeks prior to use. Depending on the ingredients, the mixture may be aged at a temperature ranging from 27° C.
to approximately 100° C. For catalyst-containing compositions, the catalyst and polysilane may partially react for a short period of time. Therefore, aging is recommended prior to use to stabilize the composition. Initial aging test results indicate that the composition reaches an equilibrium at which further shrinking of the resulting oxide film does not occur. One or ordinary skill in the art would be able to perform the necessary aging tests to determine the proper
aging duration.
[0348] After Step B or Optional Step C, the mixture may

be filtered to remove any particles or other solid content.
One of ordinary skill in the art would recognize that the filter
must be compatible with the components of the Si-containing film forming composition. PolyTetraFluoroEthylene (PTFE) is typically a suitable filtration material. The filter size ranges from approximately 0.02 micron to approximately 1 micron.
[0349] One of ordinary skill in the art will also recognize that other addition sequences are possible, such as the

pre-blending of the catalyst in the solvent or one of the solvents to facilitate the mixing and enable a more homogeneous mixture with the desired $Si_nH_{(2n+2)}$ product.

[0350] An optional process to prepare a substrate for the spin-coating process is also provided in FIG. 5.
[0351] The planar or patterned substrate on which the Si-containing film is to be deposited may be prepared for the

deposition process in Steps 1 and 2 and alternative Steps 3a preparation process. Gases are typically of semiconductor grade and free of particle contamination . For semiconductor 100 particles/mL $(0.5 \mu m)$ particle, more preferably less than 10 particles/mL) and free of non-volatile residues that would lead to surface contamination. Semiconductor grade solvents having less than 50 ppb metal contamination (for each element, and preferably less than 5 ppb) are advised. In Optional Step 1, the substrate may be sonicated in a cleaning
solvent, such as acetone, at room temperature (between
approximately 20° C. and approximately 25° C.) for
approximately 60 seconds to approximately 1 patterned substrate may then be sonicated at room temperature in another cleaning solvent, such as isopropyl alcohol (IPA) , for approximately 60 seconds to approximately 120 seconds, and preferably for approximately 90 seconds. One of ordinary skill in the art will recognize that these steps may be performed in the same or different sonicators. Different sonicators require more equipment, but provide an easier process . The sonicator must be thoroughly cleaned between Step 1 and 2 if used for both to prevent any contamination of the substrate . Exemplary sonicators suitable for the dis closed methods include Leela Electronics Leela Sonic Models 50, 60, 100, 150, 200, 250, or 500 or Branson's B Series. [0352] In Optional Step 2, the substrate may be removed from the IPA sonicator and rinsed with fresh cleaning solvent. The rinsed substrate is dried using an inert gas, such as N, or Ar.

[0353] In Optional Step 3a, the substrates of Step 2 may be treated by UV-ozone for 1 hour at 25° C. and atmospheric pressure to generated OH-terminated hydrophilic surfaces when a hydrophilic surface is desired. Step 3a also further removes organic contaminations.

[0354] In Optional Step 3b, the substrates of Step 2 are dipped into a 1% HF water solution at 25° C. for 1-2 minute to etch away the top native oxide layer, and generate H-terminated hydrophobic surfaces when a hydrophobic surface is desired.

[0355] One of ordinary skill in the art will recognize that Optional Steps 1, 2 and alternative Steps 3a and 3B provide exemplary wafer preparation processes. Multiple wafer preparation processes exist and may be utilized without departing from the teachings herein. See, e.g., Handbook of Silicon Wafer Cleaning Technology, 3rd Edition, 2017 (William Andrew). One of ordinary skill in the art may determine the appropriate wafer preparation process based at least
upon the substrate material and degree of cleanliness

required.
[0356] The substrates may proceed to the spin coating process after any of steps 2, 3a, or 3b.

[0357] The flow chart of FIG. 5 also diagrams an exemplary spin-coating process.
[0358] The substrate as optionally prepared above is trans-

ferred to the spin coater. Exemplary suitable spin coaters include Screen's Coat/Develop Track DT-3000, S-cubed's Scene12, EVG's 150XT, Brewer Science's Cee® Precision spin coaters, Laurell's 650 series spin coaters, Specialty Coating System's G3 spin coaters, or Tokyo Electron's CLEAN TRACK ACT equipment family. In Step 4, the Si-containing film forming compositions of Step B or C are dispensed onto the substrate of Step 2, 3a, or 3b. The wafer substrate is spun in Step 5. One of ordinary skill in the art will recognize that Step 4 and Step 5 may be performed
sequentially (static mode) or concurrently (dynamic mode).
Step 4 is performed using a manual or auto-dispensing
device (such as a pipette, syringe, or liquid flow met approximately 999 rpm, preferably between approximately 5 rpm to approximately 300 rpm). After all of the Sicontaining film forming composition is dispensed (i.e., when Step 4 is complete in either static or dynamic mode), rate ranges between approximately 1000 rpm to approximately 4000 rpm. The wafer is spun until a uniform coating is achieved across the substrate, which typically takes between approximately 10 seconds and approximately 3

minutes. Steps 4 and 5 produce a Si-containing film on the wafer. One of ordinary skill in the art will recognize that the required duration of the spin coating process, the acceleration rate, the solvent evaporation rate, etc., are adjustable parameters that require optimization for each new formulation in order to obtain the target film thickness and uniformity (see, e.g., University of Louisville, Micro/Nano Technology Center—Spin Coating Theory, October 2013).

[0359] After the Si-containing film is formed, the wafer is pre-baked or soft baked in Step 6 to remove any remaining volatile organic components of the PHPS composition and/ or by-products from the spin-coating process. Depending on the activation temperature of the catalyst, catalyzation may also commence in Step 6. Step 6 may take place in a thermal chamber or on a hot plate at a temperature ranging from approximately 30° C. to approximately 300° C., preferably 80° C. to 200° C. for a time period ranging from approximately 1 minute to approximately 120 minutes. Exemplary hot plates include EVG's 105 Bake Module Brewer Scie

plates.

[0360] In step 7, the substrate is cured to produce the desired material. 3 non-limiting options are shown in FIG. 5. Any of the 3 options may be performed using an inert or reactive gas. Exemplary inert gases include N2, Ar, He, Kr,

Xe, etc. The reactive gas may be used to introduce oxygen,
nitrogen, or carbon into the film.
[0361] Exemplary reactive gases that introduce oxygen
into the film include oxygen-containing gases, such as O_2 ,
 O_3 , air, Alternatively, curing may occur under a H_2O_2 at temperatures ranging from approximately 300° C. to approximately 500° C. H_2O_2 is a strong oxidizer and may permit consistent Si oxide film consistency further into t

[0362] Exemplary reactive gases that introduce carbon into the film include carbon-containing gases, and specifically unsaturated carbon-containing gases, such as alkenes and alkynes (ethylene, acetylene, propylene, etc.).

[0363] Exemplary reactive gases that introduce nitrogen into the film must have at least one N—H bond to enable the DHC reaction to proceed. For a completely C-free film, this means that the curing gas may comprise $NH₃$ or $N₂H₄$. Alternatively, C-containing N-sources may be used, but may
yield some C in the film. Exemplary C-containing N sources
include substituted hydrazines (i.e., N_2R_4 , wherein each R is
independently H or a C1-C4 hydrocarbo least one R is H)(e.g., MeHNNH₂, Me₂NNH₂, MeHN-NHMe, phenyl hydrazine, t-butyl hydrazine, 2-cyclohexyl-NHMe, phenyl hydrazine, t-butyl hydrazine, 2-cyclohexyl-
1, 1-dimethyhydrazine, 1-tert-butyl-1,2,2-trimethylhydrazine,
2ine, 1,2-diethylhydrazine, 1-tert-butyl-1,2,2-trimethylhydrazine,
1-(2-methylphenyl)hydrazine, 1-(1-ph Ethylmethylamine, Hexylamine, Isobutylam in e, Isopro-
pylamine, Methylhexanamine, Pentylamine, Propylamine, cyclic amines like pyrrolidine or pyrimidine), ethylene
diamines (i.e., $R_2N-C_2H_4-NR_2$ wherein each R is indep diamine, tetramethylethylene diamine), pyrazoline, pyri-
dine, radicals thereof, or mixtures thereof. If the desired
Si-containing film also contains oxygen, C-containing N
source may include $H_2N-C_xH_{2x}$ —OH, with x=1-4 carbon, such as ethanolamine. Preferably the reactant is $NH₃$, radicals thereof, or mixtures thereof.

[0364] In Step 7a, the substrate is subject to thermal curing
at a temperature ranging from approximately 101° C. to
approximately $1,000^{\circ}$ C., preferably from approximately
 200° C. to approximately 800° gas . A furnace or rapid thermal processor may be used to perform the thermal curing process. Exemplary furnaces include the ThermoFisher Lindberg/Blue M^{TM} tube furnace, the Thermo Scientific Thermolyne™ benchtop tube furnace or muffle furnace , the Inseto tabletop quartz tube furnace , the NeyTech Vulcan benchtop furnace, the Tokyo Electron TELINDYTM thermal processing equipment, or the ASM
International ADVANCE^o vertical furnace. Exemplary rapid thermal processors include Solaris 100, ULVAC RTP-6, or Annealsys As-one 100.

[0365] Alternatively, in Step 7b, the substrate is subject to UV-curing at a wavelength ranging from approximately 190 nm to approximately 400 nm using a monochromatic or polychromatic source. Exemplary VUV- or UV-curing systems suitable to perform Step 8b include , but are not limited to, the Nordson Coolwaves® 2 UV curing system, the Heraeus Noblelight Light Hammer® 10 product platform, or the Radium Xeradex® lamp.

[0366] In another alternative of Step 7c, both the thermal and UV process may be performed at the same temperature and wavelength criteria specified for Steps 7a and 7b. The thermal and UV curing may be performed simultaneously or sequentially. One of ordinary skill in the art will recognize that the choice of curing methods and conditions will be determined by the target silicon-containing film desired.
[0367] In another alternative, the thermal curing process

may proceed in a stepwise fashion. More particularly, the thermal curing may start at a temperature ranging from approximately 50° C. to approximately 500° C. under an inert or reactive gas for a time period ranging from approximately 10 to approximately 30 minutes. The temperature may be increased by approximately 50° C. to approximately 150 $^{\circ}$ C. and maintained for an additional 10 to 30 minutes. Additional incremental temperature increases may be used, if necessary. Alternatively, the temperature may be increased using a specified ramp and then maintained at specific temperatures for a short period of time. For example, the wafer may be placed in a room temperature chamber being heated at a ramping rate of approximately 1° C./minute to approximately 100° C./minute, preferably from approximately 5° C./minute to approximately 40° C./minute, and more preferably from approximately 10° C./minute to approximately 20° C./minute. Once the temperature reaches the desired heating temperature, for example approximately 100° C. to approximately 400° C., the ramping may be stopped for a specified period of time, for example ranging from approximately 5 minutes to approxi utes. The same or a different ramping temperature rate may

then be used to increase the chamber temperature to the next
desired heating temperature, for example approximately
300° C. to approximately 600° C. and be maintained for
another specified period of time, for example rangi This may be repeated for again if a third heating temperature
is desired, for example approximately 500° C. to approximately $1,000^{\circ}$ C. and maintained for another specified
period of time, for example ranging fro native, the curing may use a slow, steady heating ramp
without any specified time spent at any specific temperature
(e.g., approximately 0.5/minute to approximately 3° C./minute). Once curing is complete, the furnace is allowed to cool to room temperature at a cooling rate ranging from approximately 1° C./minute to approximately 100° C./minute. Applicants believe that any of these thermal curing steps may help to reduce formation of cracks and voids in the

resulting film.

[0368] Additionally, shrinkage may be further reduced by

controlling the $O_2:H_2O$ ratio when an oxygen-containing

atmosphere is required. Preferably, the $O_2:H_2O$ ratio ranges

from approximately 6:1 disclosed Si-containing film forming compositions may pro-
vide oxide shrinkage ranging from approximately -5% to
approximately 15%, preferably from approximately 0% to
approximately 10%, and more preferably from appro film ranges from approximately 0 atomic % to approximately 7 atomic %, preferably from approximately 0 atomic % to approximately 5 atomic %. The Si, O, and C concentrations may be determined by X-ray photoelectron spectroscopy (XPS). The wet etch rate ratio of the cured SiO2 film using a 1% HF-water solution ranges from approximately 1:1 to approximately 5:1 as compared to thermal

oxide grown at 1100° C.
[0369] In Step 8, the cured film is characterized using standard analytic tools. Exemplary tools include, but are not
limited to, ellipsometers, x-ray photoelectron spectroscopy,
atomic force microscopy, x-ray fluorescence, fourier-trans-
form infrared spectroscopy, scanning el

processes discussed above may include $SiO₂$; $SiNi$; $SiON$; SIOC; SIONC;; SICN; SIMCO, in which M is selected from Zr, Hf, Ti, Nb, V, Ta, Al, Ge, B, Nb. One of ordinary skill in The art will recognize that by judicial selection of the appropriate Si-containing film forming composition and co-reactants, the desired film composition may be obtained.

[0371] The disclosed Si-containing film forming compositions provide less shrinkage of Si-containing films than
prior art NH-containing PHPS compositions for applications
in shallow trench isolation dielectrics, pre-metal copy (XPS) or Energy Dispersive X-ray (EDXO spectros-
copy. Applicants further believe that the resulting oxide films will have a thin film stress measurement ranging from
approximately -160 MPa to approximately $+160$ MPa as
determined by profilometer.
[0372] The recipe for the curing of the film and conversion
to SiO₂ is also

to SIO₂ is also widely investigated to decrease the shrinkage,
as it is believed that the shrinkage is related to the loss
(volatilization) of short oligomers before they are oxidized
during the curing step. As such, th

EXAMPLES

[0375] The following non-limiting examples are provided
to further illustrate embodiments of the invention. However,
the examples are not intended to be all inclusive and are not
intended to limit the scope of the inventio

suitable means, such as by gas chromatography (GC) using
part of the product stream or an aliquot of the product. In the
following examples, GC analysis was performed on Agilent
7890A and Agilent 6890 Gas Chromatographs e

T=250 \Box C.; Reference flow: 20 mL/min; Makeup flow: 5 mL/min; Carrier gas: 5 mL/min (Helium); Oven: 35 \Box C., $\frac{m}{200}$ $\frac{m}{20}$ $\frac{m}{20}$ $\frac{m}{200}$ $\frac{m}{200}$ C., 13 min; Injector: 200 \Box C.; Splitless mode; Sample Size: 1.0 μ L.

Example 1

General Overview of Catalysts

[0378] Applicants have surprisingly discovered the catalytic activity of heterogeneous catalysts comprising elements selected from Fe, Group I, Group II, and Group III of the Periodic Table, as summarized in the Tables 1 and 2:

TABLE 1

Activity of claimed catalysts for converting liquid $Si3H8$										
Catalyst	$\mathbf T$ (h)	$i-Si4H10$	$n-Si4H10$	Si ₅ H ₁₂ A. B. C	Si ₆ H ₁₄	Si ₇ H ₁₆	Si_8H_{18}	Silanes \ge Si _o	Selectivity $n-Si_AH_{10}/i$ Si ₄ H ₁₀	Selectivity $Si5H12 C/Si5H12$ B/Si ₅ H ₁₂ A
Vitride TM /Silica		2.0	19.3	0.2: 2.9: 7.6	6.4	4.0	2.3	1.4	9.7	2.6:1:0.07
KSiPh ₃		1.6	18.1	0.2; 2.5; 6.7	49	2.7	1.5	0.9	11.0	2.7:1:0.08
KN(SiMe ₃) ₂ , 41 \Box C *		1.3	15.3	$0.1:1.8:3.5*$	$1.4 *$	$0.1 *$	$0.0*$	$0.0*$	11.8	1.9:1:0.06
$KN(SiMe2)2$.	22	1.3	14.3	0.1: 2.2: 5.3	5.9	2.6	1.6	1.5	10.7	2.4.1.0.05

	IABLE 1-continued									
Activity of claimed catalysts for converting liquid Si ₂ H _o										
Catalyst	T (h)	$i-Si_4H_{10}$	$n-Si_4H_{10}$	Si ₅ H ₁₂ A, B, C	Si ₆ H ₁₄	Si ₇ H ₁₆	Si_8H_{18}	Silanes \geq Si _o	Selectivity $n-Si4H10/i-$ Si ₄ H ₁₀	Selectivity $Si5H12 C/Si5H12$ B/Si ₅ H ₁₂ A
KH	6	0.5	5.8	0.0; 0.6; 1.2	0.6	0.2	0.1	0.0	12.1	2.0:1:0
MgH ₂	6	2.0	7.3	0.1; 0.7; 0.7	0.2	0.1	0.0	$_{0.0}$	3.6	1.0:1:0.14
LiAlH ₄		0.3	3.3	0.0; 0.1; 0.2	0.0	0.0	0.0	$_{0.0}$	10.2	2.9:1.0
n-BuLi		0.6	1.6	0.1; 1.1; 2.8	1.0	0.8	0.5	0.3	2.5	2.6:1:0.09
ActiveGel ™	$\overline{2}$	0.8	5.9	0.0; 1.1; 2.1	1.2	0.5	0.2	0.1	7.9	19:1:0
ActiveGel ™	21	1.5	11.2	0.0, 2.5, 4.1	3.6	2.3	1.4	1.5	7.3	1.6:1:0
Activelsom ™	$\overline{2}$	1.2	9.6	0.0, 2.2, 4.6	3.7	2.3	1.3	0.4	8.2	2.2:1:0
Activelsom ™	21	1.6	14.0	0.0; 2.8; 5.7	5.1	3.3	2.3	1.9	8.6	2.0:1:0
FeCl ₃ on silica-	$\overline{2}$	2.3	12.0	3.5	1.0	0.3	0.1	0.0	5.1	
49 equiv. LiNMe ₂ .										
FeCl ₂ on silica- 5.3 equiv. BuLi	3	1.1	2.4	2.9	3.4	0.7	0.4	0.4	2.2	

 TA DIE 1

All numbers in $%$ w/w.
1-5% w/w catalyst taken.

All experiments are at room temperature, except test with $KN(SiMe_3)_2$ at 41 \Box C *.

* For test with $KN(SiMe_3)_2$ at 41 \Box C, % given for volatile fraction in dry ice trap stripped from reaction products, the total amoun

Vitride ™ = Na|AlH₂(OCH₂CH₂OMe)] sold under the trademark Vitride ™ by Vertellus Holdings LLC or Red-Al ® by Sigma-Aldrich Biotechnology LP, ActiveGel ™ = 30-40%
Na/Silica sold under the trademark ActiveGel ™ by SiG

[0379] At the applied conditions, selectivity toward the n-Si₄H₁₀: KH>KN(SiMe₃)₂=KSiPh₃>LiAlH₄>VitrideTM/
Silica>ActivelsomTM>ActiveGelTM>FeCl₃/silica/
LiNMe₂>MgH₂>BuLi=FeCl₃/silica/BuLi [0380] Activity in transformation of liquid $Si₃H₈$: VitrideTM/Silica>KSiPh₃>KN(SiMe₃) r_{2} ActivelsomTM>ActiveGelTM>n-BuLi>FeCl₃/silica/
LiNMe₂>FeCl₃/silica/BuLi>MgH₂>KH>LiAlH₄.

Structures:

[0381]

 $H_3S_1 \longrightarrow H_2S_1 \longrightarrow SH_2 \longrightarrow H_3S_1 \longrightarrow SH_3 \longrightarrow SH_3$ H_12C n - tetrasilane iso - tetrasilane

$$
H_3Si \longrightarrow Si
$$

 $H_3S_1 \longrightarrow S_1 \longrightarrow S_1H_3$

 $\frac{\text{SiH}_3}{\text{H}_3\text{Si}} \times \frac{\text{SiH}_2}{\text{SiH}_2} \text{SiH}_2 \times \frac{\text{SiH}_2}{\text{SiH}_3}$

-continued

 $\widetilde{\text{SiH}}^{\text{SIII}_3}$

 \sum SiH₂

TABLE 2

Temperature range tested 50-300 \Box C., the highest activity reported in the table. No solid formed at the reported conditions. [0382] The n-Si₄H₁₀/i-Si₄H₁₀ isomeric ratio and quantity

of higher silanes is much higher for the liquid phase reactions.

[0383] Additional synthesis details are provided in the examples that follow.

Comparative Example 1

Summary of Results for Prior Art Catalysts with Liquid $Si₃H₈$

[0384] Catalysis of liquid $Si₃H₈$ using the prior art homogeneous catalysts $Cp₂ZrCl₂/BuLi$, $Cp₂ZrCl₂/LiNMe₂$, $RuCl₄(p-Cymene)₂$ and $Ni(COD)₂ (COD = cyclooctadieny)$ was performed [catalysts from Joyce Y. Corey, "Dehydrocoupling of Hydrosilanes to Polysilanes and Silicon Oli-
gomers: A 30 Year Overview", Advances in Organometallic Chemistry, Volume 51, 2004 Elsevier Inc.]. Catalysis of liquid $Si₃H₈$ using the prior art heterogeneous catalysts Ru (5%) C and Rh (5%) C was also performed [catalysts from " Method for Producing a Semiconductor Material", Keizo Ikai; Masaki Minami; Mitsuo Matsuno, Nippon Oil Co., Ltd., U.S. Pat. No. 5,700,400 A, Aug. 14, 1995]. FeCl₃ on silica and in combination with MMAO (MMAO=modified methylaluminoxane, formula $[(CH_3)_0, _{95} (n-C_8H_{17})$ _{0.05}AlO] $_n$) were also tested.

TABLE 3

α . α is an α , i.e., α , α , α , α , α , α Bull. Korean Chem. Soc. 2000, 21,	Comparative test of catalysts toward the liquid trisilane.						
Example 3	Si_4H_{10} Si ₅ H ₁₂ %	Result	time	Additive	Catalyst (g) per 3- 4 g Si_3H_8		
	n.d.	Polymer	1 min		$Cp_2ZrCl_2 (0.32 g)$ BuLi 2.5M		
$Si3H8$ and $FeCl3(5%)$ on s	n.d.	5 min Polymer		in Hexane BuLi 2.5M			
				in Hexane	$Cp_2TiCl_2 (0.27 g)$		
A similar reaction to Exa [0390] with comparable quantities of $Si3H$	n.d.	Polymer	1.5 _h	LiNMe ₂	Cp_2ZrCl_2 (<0.01		
and with 45-50 equivalents of Li	n.d.	Polymer	0.5 _h		$Cp_2TiCl_2 (0.27 g)$ LiNMe ₂		
n-BuLi solution. The reaction pro- separated by distillation affording a	similar	No reaction	5 h	Tiba $(A^{\dagger}Bu_3)$ (0.15 g)	$Cp_2ZrCl_2 (0.02 g)$		
ticipated Mn 600-900 DA) and y	similar	No reaction	3 _h	No	FeCl ₃ $(5%)$ on Silica (0.11 g)		
mixture of silanes (according to G	similar	No reaction	3 _h	MMAO	$FeCl3$ (5%) on		
Si_2 — Si_6 silanes containing 3.5% Si				$(Fe:AI = 1:100)$	Silica (0.11 g)		
iso- Si_4H_{10} and 12.0% n- Si_4H_{10} ,	similar	No reaction	24 _h	N0	$RuCl4(p-$		
$n-Si_5H_{12}$).					$Cymene$ ₂		
	similar	No reaction	24 h	MMAO12	$RuCl4(p-$		
Example 4				$(35%)$ on silica	$Cymene$ ₂		
	similar	No reaction	24 _h	Al ⁱ Bu ₃ /	$RuCl4(p-$		
Liquid Si_3H_8 and $Na[A1H_2(C)]$				MMAO12 $(35%)$ on	$Cymene$ ₂		
$(35%)$ on silic				silica			
	similar	No reaction	24 h	No.	$Ni(COD)_{2}$		
A 35% w/w $Na[A1H2(OC)$ [0391]	similar	No reaction	24 h	No.	Ru (5%)/C		
catalyst was prepared by mixing \sim 0.000 \sim \sim \sim \sim \sim \sim \sim	similar	No reaction	24 h	No	Rh (5%)/C		

COD = cyclooctadiene

 $n.d. = not detected.$

 \sin similar = relative amount of higher silanes after test is similar to that in trisilane before test within $(\pm 0.2\%)$

[0385] The CpTiCl₂ and CpZrCl₂ homogeneous catalysts polymerized trisilane to a non-volatile solid in a noncontrollable fashion. As a result, these catalysts are not useful for controllable synthesis of isomerically enriched tetrasilane or liquid higher silanes .

[0386] The RuCl₄(p-Cymene)₂, Ni(COD)₂, and FeCl₃ homogeneous catalysts and Ru (5%)/C and Rh (5%)/C heterogeneous catalysts are not active for the transformation of non-substituted liquid trisilane to a higher silanes.

Example 2

$Si₃H₈$ and $FeCl₃(5%)$ on silica—BuLi

[0387] Liquid $Si₃H₈$ (24.7 g, 0.27 mol) and solid commercially available FeCl₃ (5% w/w)/Silica (2.7 g, 0.83 mmol $FeCl₃$) were charged in to reactor under the nitrogen atmosphere. Then n-BuLi solution (4 mL, 2.5 Molar in hexane, 10 mmol) was added to the mixture. The reaction mixture under 1 atm. of nitrogen was stirred for 24 hours at room temperature and then filtered. The flask with the liquid after filtration was connected to the vacuum manifold and all volatiles were distilled off in a trap cooled with the liquid nitrogen. GC of distillate in the liquid N_2 trap revealed a mixture of Si_2-Si_8 silanes and hexane isomers, containing 31.6% Si_3H_8 ; 3.9% iso- Si_4H_{10} ; 16.4% n- Si_4H_{10} ; 11.7% Si_5H_{12} ; and 3.4% of silanes with **6-8** or more silicon atoms.

4.4 g of undistilled liquid remained.

[0388] GPC of the undistilled liquid was obtained using

the Refractive index detector. The GPC results are provided

in FIG. 6. Polysilane has Mn=810±63 DA, Mw/Mn=2.2±0.

4, Mn corr

Li metal in THF or DME afforded only dimer trimer and tetramer [Park, M. J.; Lee, S. L.; Park, M. K.; Han, B. H. Bull. Korean Chem. Soc. 2000, 21, 336].

$Si₃H₈$ and $FeCl₃(5%)$ on silica—LiNMe₂

[0390] A similar reaction to Example 2 was performed with comparable quantities of $Si₃H₈$ and FeCl₃(5%)/Silica and with 45-50 equivalents of $LiNMe₂$ in place of the n-BuLi solution. The reaction products were filtered and separated by distillation affording a nonvolatile liquid (anticipated Mn 600-900 DA) and volatile liquid, being a mixture of silanes (according to GC analysis a mixture of \rm{Si}_2 — \rm{Si}_6 silanes containing 3.5% \rm{Si}_2H_6 , 78.0% \rm{Si}_3H_8 ; 2.3% iso- Si_4H_{10} and 12.0% n- Si_4H_{10} , 1.5 iso- Si_6H_{12} , 2.0% $n-Si_5H_{12}$).

Example 4

Liquid $Si₃H₈$ and $Na[AlH₂(OCH₂CH₂OH₂)$ $(35%)$ on silica

[0391] A 35% w/w Na $[AlH_2(OCH_2CH_2OMe)_2]$ on silica catalyst was prepared by mixing silica, pre-dried in a vacuum at 300° C., with a 65% w/w toluene solution of Na[AlH₂(OCH₂CH₂OMe₎₂] (sold under the trademark VitrideTM by Vertellus Holdings LLC or Red-Al® by Sigma-
Aldrich Biotechnology LP). The mixture was stirred 12 hours at room temperature. All volatiles were removed under dynamic vacuum . The remaining solid was utilized as

[0392] Liquid Si_3H_8 (3.1 g, 33.6 mmol) and the catalyst (0.18 g, Na [AlH₂ (OCH₂CH₂OMe)₂] 0.063 g, 0.32 mmol)

were stirred for 3 hours at room temperature in a vial in a glove box. The stirring was stopped after 3 hours. The solids were allowed to settle and an aliquot of the clear supernatant collected for GC analysis. GC analysis of the clear solution revealed a mixture of $Si₂ - Si₈$ silanes containing 44.8% Si_3H_8 ; 2.0% iso- Si_4H_{10} and 19.3% n- Si_4H_{10} , 10.8% Si_5H_{12} total and 14.1 % of silanes with 6 or more silicon atoms in total.

[0393] VitrideTM on Silica transformed the liquid trisilane in the pot reaction to a mixture of silanes with excellent selectivity toward n-Si₄H₁₀ [n-Si₄H₁₀:i-Si₄H₁₀=9.7:1]. In order to illustrate the applicability of the process for indus trial applications, a flow reaction was performed with gaseous and liquid trisilane .

w/w VitrideTM on silica (3.7 g of active component) on 3.0 g of glass wool. The effluent was collected in the dry ice trap, followed by liquid nitrogen trap. The content of the traps was analyzed by GC . The contents of the dry ice trap were distilled, affording 127.1 g of distillate and 4.4 g of nondistilled liquid. The non-distilled liquid was a mixture of silanes with 6 and more silicon atoms [GC]. GPC of non-distilled liquid dissolved in p-tolylsilane: Mn=360-760 DA, average Mn corresponds to 19 SiH₂ units. The GC results are provided in FIGS. 7 and 8. The distillate is a mixture of volatile silanes with 1-8 silicon atoms . The results are summarized in the Tables 6 and 7 below. It is anticipated that the nondistilled liquid contains a mixture of higher silanes with 14-27 silicon atoms, similarly to Examples 2 and 3.

TABLE 6

Results of flow reaction applying the liquid trisilane and Vitride TM on Silica.									
Precursor content (g)	Si_3H_8 consumed (g)	Selectivity: $n-Si_4H_{10}$. Si ₄ H ₁₀	Yield: $n-Si4H10$ $\frac{0}{0}$	Yield: Si ₅ H ₁₂ $\frac{0}{0}$	Av. T (° C.)	Residence time (Seconds)			
(149.5 g) [99.7% \rm{Si}_3H_8]	77.0	6.3:1	13.6	6.3	52.0 ± 0.7	275-421 $Av. = 358$			
(163.8 g) [99.7% Si_3H_8]	$44.1*$	7.2:1	10.5	2.3	50.7 ± 1.1	271-418 Av. $= 369$			
(143.9 g) [99.6% Si_3H_8]	38.2	7.2:1	10.2	2.5	53.7 ± 2.3	210-432 $Av = 359$			

TABLE 7

1) Similar catalyst used in all experiments, 1 and 14 days after packing.
2) Vitride TM on Silica is significantly active

3) 129 g of styring consumed by //3 g of 40.8% virtue ²¹⁰ on since (3./1 g of active component) in 3 now experiments (1 pass each, results of 2 experiments are shown here) producing n-Si₄H₁₀ with the reasonable se

Liquid $Si₃H₈$ and $Na[AlH₂(OCH₂CH₂OMe)₂],$ Flow Process

[0394] Three separate samples of liquid $Si₃H₈$ passed through a stainless steel tube reactor (20.9 cm Lx1 cm Inner Diameter) at $52.0 \pm 0.7 \square$ C. and pressure 31.1 ± 0.6 psig with the flow 1.1 ± 0.1 g/min. The reactor contained 7.9 g of 46.8%

Example 5 Example 6

Liquid Si_2H_6/Si_3H_8 Mixture and Na[AlH₂ (OCH₂CH₂OMe)₂], flow process

[0395] Liquid Si₂H₆ (42.5% w/w)—Si₃H₈ (57.5% w/w) (174.0 g) passed through the reactor (20.9 cm L×1 cm ID) at 51.9 ± 3.5 OC. and pressure 30.6 ± 0.4 psig with the flow 1.4 \pm 0.3 g/min, corresponding to the residence time 442 \pm 77

g of liquid, GC: 74.2% Si_3H_8 ; 1.3% iso- Si_4H_{10} and 15.3% n-Si₄H₁₀, 1.8% i-Si₅H₁₂, 3.5% n-Si₅H₁₂ and 1.6% of silanes with 6 or more silicon atoms in total] and liquid nitrogen trap

[4.7 g, GC: 45.9% SiH₄, 49.1% Si₂H₆, 5.0% Si₃H₈]. Undistilled reaction product [3.2 g] remaining in the pot was filtered and also analyzed by GC [mixture of Si₅—Si₁₂ silanes]. It is anticipated that the nond

ing siliconations, similarly to Examples 2 and 3. [0397] KN(SiMe_3)₂ transformed the liquid trisilane in the pot reaction to a mixture of silanes with excellent selectivity

toward the n Si_4H_{10} [n Si_4H_{10} : $Si_4H_{10} = 11.8$: 1] and selec-

sec. The reactor contained 7.0 g of 46.8% w/w VitrideTM on silica (3.3 g of active component) on 2.6 g of glass wool. The effluent was collected in the dry ice trap, followed by liquid nitrogen trap. The content of the traps was analyzed by GC.
The content of dry ice trap was distilled, affording 2.0 g of
nondistilled liquid. The non-distilled liquid was a mixture of silanes with 6 and more silicon atoms [GC]. GPC of non-distilled liquid dissolved in p-tolylsilane and 10% w/w Si_4H_{10} : Mn=561 DA, Mw/Mn=1.03, Mn corresponds to 19 $SiH₂$ units. The distillate is a mixture of volatile silanes with 1-8 silicon atoms . The results are summarized in the Tables 8 and 9 below.

TABLE 8

Results of flow reaction using disilane-trisilane mixture and Vitride TM on Silica.									
Precursor content (g)	Si ₂ H ₆ formed (g)	Si ₂ H ₈ consumed (g)	Selectivity: $n-Si4H10$:i- Si ₄ H ₁₀	Yield: $n-Si_AH_{10}$ $\frac{0}{n}$	Yield: Si ₅ H ₁₂ $\frac{0}{0}$	Volatile Heavies \ge Si6	Non distilled		
100.05 g Si_3H_8 $(57.5\%) \&$ 73.95 g Si_2H_6 (42.5%)	$+29.0$	-52.67	5.5.1	9.9	2.6	0.31 g	2.0 g		

TABLE 9

As seen from the tables, the relative amount of heavier silanes (Siz 6) is lower for the mixture of Si₂H₆—Si₃H₈.

$Si₃H₈$ and $KN(SiMe₃)₂$ Example 8

[0396] Liquid $Si₃H₈$ (28.8 g, purity 99.8% w/w) and solid KN($SiMe₃$), (0.3 g, 1.5 mmol) were charged into a reactor equipped with a thermocouple in a glove box. Two traps were installed after the reactor . The first trap after the condenser remains empty at room temperature during the reaction. The second trap is cooled with liquid nitrogen to trap SiH_4 and Si_2H_6 during the reaction. The reactor was connected to a manifold and the N_2 atmosphere replaced
with a He atmosphere. The condenser was filled with dry ice.
The reaction mixture under 1 atm. of helium was heated to $41-50$ ^o C. and stirred for 3 hours. After 3 hours, the heating was stopped and the reaction mixture cooled to room temperature. Dry ice was removed from the condenser. The first trap was cooled with dry ice and the second trap remained cooled with liquid N₂. The reaction products were stripped under the dynamic vacuum in the dry ice trap [20.2]

tivity toward the n-Si_sH₁₂. In addition, heavier silanes are produced. In order to illustrate the applicability of the process for industrial applications, flow reaction was per-Example 7 formed with gaseous and liquid trisilane.

Flow Process with the Liquid $Si₃H₈$ and Solid $KN(SiMe₃)$, Catalyst

[0398] Si₃H₈ liquid (178.2 g, 99.8% w/w) passed through the stainless steel tube reactor $(20.6 \text{ cm L} \times 1 \text{ cm } \text{Inner})$ Diameter) at 73.2 ± 1.8 ° C. and pressure 27.2 ± 0.5 psig with the flow 1.2 ± 0.3 g/min, corresponding to the residence time 467 ± 106 sec. The reactor is the 1 cm inner diameter, 20.8 cm long stainless steel tube containing 3.6 g of KN(SiMe₃)₂ on 3.6 g of glass wool. After passing of liquid trisilane through
the flow reactor, the products are collected in the dry ice trap
(175.5 g), followed by liquid nitrogen trap (2.7 g). The gas
and liquid phases of the traps to separate the higher silanes from the products collected in dry ice trap, the contents of the dry ice trap were stripped out under vacuum leaving 1.7 g of nondistilled liquid, which is

considered as mixture of "non volatile" silanes under the applied conditions 50 mtorr Vacuum and 45 \Box C. The nondistilled liquid was a mixture of silanes with 6 and more silicon atoms [GC]. 173.8 g of volatile silanes were stripped from the dry ice trap. The volatile silanes fraction was a mixture of silanes Si_nH_{2n+2} (n=1-8). It is anticipated that the remaining nondistilled liquid contains mixture of higher silanes with 14-27 silicon atoms, similarly to two above given examples 3 and 4. Results are in the Tables 10 and 11 below.

passed through the stainless steel tube reactor $(20.6 \text{ cm L} \times 1)$ cm Inner Diameter) at $81.4\pm2.3^{\circ}$ C. and pressure 29.3 ± 0.5 psig with the flow 1.4 ± 0.3 g/min corresponding to the residence time 414±85 sec. The effluent was collected in a dry ice trap (137.1 g) , followed by a liquid nitrogen trap (4.2 g) g). The gas and liquid phases of the traps were analyzed by GC. In order to separate the higher silanes from the products collected in dry ice trap , the contents of the dry ice trap were stripped out under vacuum leaving 4.2 g of nondistilled

TABLE 10

	Results of flow reaction applying the liquid trisilane and $KN(SiMe3)$.					
Precursor content (g)	Si ₃ H ₈ consumed (g)	Selectivity: $n-Si_4H_{10}$: $i-Si_AH_{10}$	Yield: $n-Si_4H_{10}$ $\frac{0}{0}$	Yield: Si ₅ H ₁₂ $\frac{0}{0}$	P (psig)	Residence time (Seconds)
(178.2 g) [99.8% Si ₃ H ₈	34.9	7.7:1	6.8	1.7	27.2 ± 0.5	310-587 Av. $= 467$

TABLE 11

[0399] KN(SiMe_3)₂ is significantly active in flow reaction with the liquid trisilane and produces a low amount on nonvolatile silanes.

[0400] The present example illustrates a potential for scaling.

Example 9

Exemplary Experiment for the Flow Process with the Mixture of Liquid $Si₃H₈$ and $Si₄H₁₀$ and Solid $KN(SiMe₃)₂$ Catalyst.

[0401] $Si₃H₈$ liquid (141.3 g), containing 93.7% w/w Si_3H_8 , 4.4% w/w i- Si_4H_{10} and 1.8% w/w n- Si_4H_{10} , was liquid, which is considered as mixture of "non volatile" silanes under the applied conditions 50 mtorr Vacuum and $45 \square$ C. The nondistilled liquid was a mixture of silanes with 6 and more silicon atoms $[GC]$. 130.0 g of volatile silanes were obtained from the dry ice trap. The volatile silanes fraction was a mixture of silanes Si_nH_{2n+2} (n=1-8). It is anticipated that the remaining nondistilled liquid contains a mixture of higher silanes with 14-27 silicon atoms, similarly to Examples 2 and 3. The results are provided in the Tables 12 and 13 below.

TABLE 12

	Results of flow reaction applying the liquid trisilane and $KN(SiMe3)$.					
Precursor content (g)	Si ₂ H ₈ consumed (g)	Selectivity: $n-Si4H10$: $i-Si_4H_{10}$	Yield: $n-Si4H10$ $\%$	Yield: Si ₅ H ₁₂ $\frac{0}{n}$	Av. T $(^\circ$ C.)	Residence time (Seconds)
(141.3 g) [93.7% Si_3H_8 , 4.4% i- $Si4H10$ 1.8% n- Si_4H_{10}	32.4	-2.38 g $i-Si_AH_{10}$; $+8.59$ g $n-Si4H10$	6.5	3.0	81.4 ± 2.3	291-497 Av. $= 414$

TABLE 13

24

[0402] Addition of Si_4H_{10} to Si_3H_8 increases pentasilane yield (4.22 g vs 3.06 g for pure Si_3H_8) and yield of heavier silanes (4.2 g vs. 1.8 g for pure $Si₃H₈$). The process is more beneficial for synthesis of heavier silanes.

[0403] While embodiments of this invention have been shown and described, modifications thereof can be made by one skilled in the art without departing from the spirit or teaching of this invention. The embodiments described herein are exemplary only and not limiting. Many variations and modifications of the composition and method are possible and within the scope of the invention. Accordingly the scope of protection is not limited to the embodiments described herein, but is only limited by the claims which follow, the scope of which shall include all equivalents of the subject matter of the claims.
We claim:

1. A method of producing $\text{Si}_{n}\text{H}_{(2n+2)}$, wherein n=4-100, the method comprising: reacting a liquid $\text{Si}_{a}\text{H}_{(2a+2)}$ reactant, wherein a=1-4, with a catalyst to produce $\text{Si}_{n}\text{H}_{(2n+2)}$, wherein n>a and the cata

- a) a heterogeneous catalyst selected from a Group I, II or III element from the Periodic Table or oxides, alkyls,
- III hydrides, silanides, or silyl amides thereof; or b) a catalyst comprising a Fe halide combined with an alkyl lithium LiR or lithium amide LiNR₂, with each R

alkyl lithium Lik or lithium amide LINK₂, with each K
independently a C1-C4 alkyl group.
2. The method of claim 1, wherein the $\text{Si}_aH_{(2a+2)}$ reactant
is Si_3H_8 .
3. The method of claim 1, wherein the $\text{Si}_aH_{(2a$

a Fe halide combined with an alkyl lithium LiR or lithium amide $LiNR_2$, with each R independently a C1-C4 alkyl group.

10. The method of claim 1, further comprising fractionally distilling Si_n $H_{(2n+2)}$ to produce a Si-containing film forming composition comprising approximately 95% w/w to approximately 100% w/w n-Si_s H_{12} .

11. The method of claim 1, further comprising fractionally distilling $\text{Si}_{n}H_{(2n+2)}$ to produce a Si-containing film forming composition comprising approximately 95% w/w to approximately 100% w/w n- \overline{Si}_6H_{14} .
12. The method of claim 1, further comprising fraction-

ally distilling $Si_nH_{(2n+2)}$ to produce a Si-containing film forming composition comprising approximately 95% w/w to approximately 100% w/w n-Si₇H₁₆.

13. The method of claim 1, further comprising fractionally distilling Si_n $H_{(2n+2)}$ to produce a Si-containing film forming composition comprising approximately 95% w/w to

approximately 100% w/w n-Si₈H₁₈.
14. A method of selectively synthesizing an isomerically enriched polysilane having a formula $Si_nH_(2n+2)$, wherein

n=5-8, the method comprising:
catalyzing a liquid $\text{Si}_a\text{H}_{(2a+2)}$ reactant, wherein a=1-4, to produce the isomerically enriched polysilane having a ratio of one isomer to another isomer ranging from

and of one isomer to another isomer ranging nom
approximately 2:1 to approximately 15:1.
15. The method of claim 14, wherein the heterogeneous
16. The method of claim 14, wherein the heterogeneous
catalyst is KN(SiMe₃)

approximately 100% w/w n-Si₃H₁₂.
19. The method of claim 14, further comprising fractionally distilling Si_nH_(2n+2) to produce a Si-containing film forming composition comprising approximately 95% w/w to

approximately 100% w/w n-Si₆H₁₄.
20. The method of claim 14, further comprising fractionally distilling Si_nH_(2n+2) to produce a Si-containing film forming composition comprising approximately 95% w/w to approximately 100% w/w n-Si₇H₁₆.

* * * *