COPOLYMER SEALANT COMPOSITIONS AND METHOD FOR MAKING

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ABSTRACT

Condensation curable poly(fluoroorgano)siloxane-poly(silarylene)siloxane block copolymer compositions having a glass transition temperature not exceeding about -54°C and excellent solvent resistance have been found useful as sealants. Polyalkoxysilylorgano compounds, such as 1,4-bis[trimethoxysilyl(ethyl)]benzene have been found to be effective as cross-linkers.

13 Claims, No Drawings
1 COPOLYMER SEALANT COMPOSITIONS AND METHOD FOR MAKING

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BACKGROUND OF THE INVENTION

The present invention is directed to copolymer compositions useful in the preparation of fuel tank sealants. More particularly, the present invention relates to condensation curable polyfluororganosiloxane-poly(dimethyldisiloxane) siloxane block copolymer compositions and the use of polyalkoxysilylorganic compounds as cross-linkers to facilitate the condensation cure of polyfluororganosiloxane-poly(dimethyldisiloxane) siloxane block copolymers.

As shown by Smith, U.S. Pat. No. 3,109,826, bis(alkoxysilyl) hydrocarbons, such as 1,2-bis(triethoxysilyl) ethane, can be used as cross-linkers in combination with a metal salt to effect the neutral condensation cure of hydroxy end-blocked polydiorganosiloxanes. However, the resulting cured silicone compositions have been found to be problematic as aircraft fuel tank sealants, as they do not meet the required solvent or fuel resistance. It is known that fluorosilicones, for example, made by polymerizing tris[(trifluoropropyl)methyl]cyclosiloxane, can provide excellent fuel resistance. However, fluorosilicones often do not meet the wide temperature stability requirements needed in aircraft sealants, such as temperatures in a range between -54°C and 177°C over an extended period of time. In addition, fluorosilicones are subject to depolymerization which can result in the formation of low molecular weight cyclics.

In an effort to enhance the thermal stability of fluorosilicones, non-siloxane groups, such as p-silphenylene, have been inserted into the polyfluorosiloxane backbone, as shown by Grassie and Beattie, "The Thermal Degradation of Polysiloxanes: Part 7", Polymer Degradation and Stabilization 8:177-193 (1984). It is also reported by Dvornic and Lenz, Macromolecules, 25, 3769 (1992), that copolymers having a glass transition temperature (Tg) of -51°C can be made by reacting methyl(3,3,3-trifluoropropyl)isocyanate, providing excellent fuel resistance. However, fluorosilicones often do not meet the wide temperature stability requirements needed in aircraft sealants, such as temperatures in a range between -54°C and 177°C over an extended period of time. In addition, fluorosilicones are subject to depolymerization which can result in the formation of low molecular weight cyclics.

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While fluorosilicones having improved thermal stability have been made by inserting non-siloxane groups, such as p-silphenylene into the polyfluorosiloxane backbone, such copolymers have been found to have a glass transition temperature which does not satisfy the minimum -54°C Tg flexibility requirements of aircraft fuel tank sealants.

Experience also has shown that in addition to being sensitive to depolymerization, condensation curable fluorosilicone compositions often suffer from an incomplete cure using a conventional curing catalyst, such as a tin salt, and a standard neutral condensation curable cross-linker, for example a polyalkoxysiloxane. One possible explanation, as discussed by Fujiaki, U.S. Pat. No. 5,236,997, is that the steric hindrance or electronic effects of bulky terminal trifluoropropyl groups inhibit crosslinking of the network.

Accordingly, depolymerization resistant silicone base copolymers which could be compounded to a fuel resistant condensation curable silicone composition convertible to the elastomeric state upon cure exhibiting stability over an operable temperature range in a range between about -54°C or below and at least about 177°C over an extended period of time are constantly being sought which also exhibit low temperature flexibility, in addition to high temperature stability.

2 BRIEF SUMMARY OF THE INVENTION

The present invention provides a condensation curable polyfluororganosiloxane-poly(dimethyldisiloxane) siloxane block copolymer exhibiting a glass transition temperature not exceeding about -54°C.

A further embodiment of the present invention provides a method for making a polyfluororganosiloxane-poly(dimethyldisiloxane) siloxane copolymer comprising effecting reaction between a bis(diorganohydroxysilyl)arylene and a polyfluoralkylarylene)cyclopolsiloxane.

In yet another embodiment of the present invention, there is provided a neutral condensation curable polyfluororganosiloxane-poly(dimethyldisiloxane) siloxane block copolymer sealant composition comprising:

(a) a polyfluororganosiloxane-poly(dimethyldisiloxane) siloxane block copolymer,
(b) a cross-linker, and
(c) a condensation catalyst.

In yet another embodiment of the present invention, there is provided a method for making a neutral condensation curable polyfluororganosiloxane-poly(dimethyldisiloxane) siloxane block copolymer sealant composition which comprises:

(a) effecting reaction between bis(diorganohydroxysilyl)arylene and polyfluoralkylarylene)cyclopolsiloxane to form a condensation curable polyfluororganosiloxane-poly(dimethyldisiloxane) siloxane block copolymer,
(b) shearing the copolymer, and
(c) blending a cross-linker and a condensation catalyst with the copolymer to form a sealant.

DETAILED DESCRIPTION OF THE INVENTION

The polyfluororganosiloxane-poly(dimethyldisiloxane) siloxane block copolymer, or "block copolymers" within the scope of the present invention, can be used to make aircraft fuel tank sealants having property profiles which include a glass transition temperature (Tg) of about -54°C or below and can be made by the ring opening polymerization of a polyfluoralkylarylene)cyclopolsiloxane in the presence of a bis(diorganohydroxysilyl)arylene.

As used hereinafter, the term "bis(diorganohydroxysilyl)arylene", or "bis(hydroxysilyl)arylene" is shown by the formula,

\[ HO-(R)SiO-(R)SiH \],

and the term "silarylenesiloxoy" can be represented by the formula,

\[ -(R)SiO(SiR)O- \],

where Q^2 is a C_{n-12} divalent aromatic organic radical, and R is a C_{1-4} alkyl radical. Preferably, the bis(diorganohydroxysilyl)arylene is 1,4-bis(dimethylhydroxysilyl)benzene.

"Polyfluoralkylarylene)cyclopolsiloxane," sometimes expressed as "polyfluoralkyl)cyclic silicone" is shown by the following formula:

\[ [(R^1R^2SiOP)]_{m} \],

where R^1 is a C_{1-12} polyfluoralkyl radical, R^2 is a C_{1-12} organic radical, and "m" is an integer in a range between about 3 and about 8 inclusive, and preferably 3 or 4. Polyfluoralkylarylene)cyclopolsiloxane can be used in a proportion in a range between about 0.5 and about 4 moles,
and preferably in a range between about 1 and about 2 moles, per mole of bis(diorganohydroxysilyl)arylene. Reaction to form the copolymer is typically effected with agitation under near conditions or in the presence of an inert organic solvent, for example, toluene, at temperatures in a range between about 60°C and about 150°C. Reaction is typically effected for a time period in a range between about 30 minutes and about 2 hours, and preferably, in a range between about 35 minutes and about 1.5 hours. It is preferred to operate within an inert atmosphere, for example, under a nitrogen blanket. Typically, there is also present an initiator, for example, an alkali hydroxide (e.g. sodium hydroxide) or an alkali fluorosilanolate (e.g. sodium fluorosilanolate) in a range between about 5 parts and about 15 parts by weight per 100 parts of block copolymer, based on the weight of the reaction mixture, and a quencher, for example, phosphoric acid or silyl phosphate in a range between about 10 ppm and about 60 ppm, based on the weight of reaction mixture.

C_{1-4} alkyl radicals which are included within R are, for example, methyl, ethyl, propyl and butyl; C_{3-6} polyfluoroalkyl radicals which are included within R^2 include but are not limited to, for example, trifluoropropyl units, tridecafluoro-1,1,2,2-tetrahydrocyclooctyl units, nonafluoro-1,1,2,2,5,5-hexahydrocyclohexyl units, and pentaffluoro-1,1,2,2-tetrahydrobutyl units. Radicals included within R^2 include, but are not limited to, for example, methyl, ethyl, propyl, butyl, and phenyl.

Among the poly(fluoroalkylorganosiloxanes) there is preferably included tris(trifluoropropyl)methylcyclosiloxane.

The sealant composition of the present invention includes a bis(polyalkoxysilyl)organoglycol cross-linker having the formula, [(RO)_{2-n}XSi]_{2n+1}{\text{O}},

where Q is a C_{2-12} divalent organic radical, R is as previously defined, and X is a member selected from the group consisting of R and RO. Some of the C_{2-12} divalent hydrocarbon radicals included within Q are dialkylene radicals such as dimethylene, trimethylene, tetramethylene, pentamethylene, and hexamethylene. Also included are divalent arylene radicals, for example, phenylene, tolylene, xylylene and naphthylene. Divalent aromatic organic radicals included within Q are for example phenylene, tolylene, xylylene and naphthylene.

The cross-linker can be used in combination with an effective amount of a condensation catalyst, such as a tin compound, to effect the neutral condensation cure of block copolymers comprising blocks of poly[furoorganosiloxy] organosiloxane units chemically combined with poly(silylarenyl)siloxane groups. The block copolymers are referred to as “poly(fluoro)organosiloxane-poly(silylarenyl)siloxane block copolymers”.

The sealant composition of the present invention is preferably made by initially compounding the poly(fluoroorganosiloxane-poly(silylarenyl)siloxane copolymer, or “block copolymer” into a curable sealant base, or masterbatch under shearing conditions. Shearing can be effected at a temperature in a range between about 25°C and about 200°C, preferably, in a range between about 100°C and about 150°C, at atmospheric pressure, and preferably under inert conditions, such as under an inert gas, for example nitrogen, in a mixer, for example, a double-planetary mixer. Shearing can be carried out over a period in a range between about 15 minutes and about 4 hours, preferably in a range between about 1 hour and about 2 hours.

The block copolymer can be blended with a filler which can include a reinforcing filler, such as fumed silica, or a combination thereof with an extending filler, such as diatomaceous earth, precipitated silica, ground quartz, or calcium carbonate. It is preferred to use fumed silica which has been pretreated with an effective amount of a cyclic siloxane, such as octamethyloctasiloxane, or a mixture thereof with an organosilazane, such as hexamethyldisilazane. A proportion of filler in a range between about 0 parts and about 30 parts by weight, per 100 parts of block copolymer can be used. Preferably, the filler is present in a range between about 5 parts and about 15 parts by weight per 100 parts of block copolymer. A heat stabilizer, such as iron oxide, in a range between about 0.1 and about 10 parts by weight, per 100 parts of block copolymer has been found to be effective.

After blending of the masterbatch ingredients, the mixture can be degassed under reduced pressure, and the resulting paste can be stored. The neutral condensation curable poly(fluoroorganosiloxane-poly(silylarenyl)siloxane copolymer sealant compositions can be made by blending the block copolymer masterbatch with an effective amount of a condensation catalyst and a suitable bis(polyhalosiloxyl)organoglycol cross-linker, referred to hereinafter sometimes as “cross-linker”. “Neutral” as used herein refers to a sealant composition which is substantially acid-free and substantially base-free.

Suitable condensation catalysts are present in a range between about 0.1 and about 2 parts, per 100 parts of block copolymer and include, for example, organo-metal compounds such as dibutyl tin diacetate, dimethyltin neodecanoate, dibutyl tin dilaurate, stannous octoate, dimethylhydroxysiloxane, or combinations thereof.

Some of the preferred bis(polyalkoxysilyl)organoglycol cross-linkers included within formula (1) are, for example, 1,2-bis(triethoxysilyl)ethane, 1,6 bis(trimethoxysilyl)hexane, 1,4-bis[trimethoxysilyl(ethyl)]benzene, 1,2-bis(methyl dichloxsilyl)ethane, and 1,6-bis(methyldiethoxysilyl)hexane. The cross-linkers are present in a range between about 1 and about 20 parts, per 100 parts of block copolymer.

In order that skilled in the art will be better able to practice the present invention, the following examples are given by way of illustration and not by way of limitation. All parts are by weight unless otherwise indicated.

**EXAMPLE 1**

A poly(fluoroorganosiloxane-siloxane-poly(silylarenyl)siloxane block copolymer useful in making a condensation curable sealant composition was prepared as follows:

A mixture of 2.26 grams (g) (0.010 mole) of 1,4-bis(dimethylhydroxysilyl)benzene and 9.42 g (0.020 mole) of tris[(trifluoropropyl)methyl]cyclosiloxane was heated with stirring to about 150°C under a nitrogen atmosphere. There was then added 0.04 g of sodium fluorosilanolate. After 60 minutes at 150°C, there was added to the mixture 0.065 g of a silyl phosphate. The reaction mixture was maintained with stirring for an additional at 150°C and then allowed to cool. Based on method of preparation, there was obtained a block copolymer, including chemically combined blocks of 1,4-dimethyl-silylphenylenesiloxane units and blocks of methyl(3,3,3-trifluoropropyl)siloxane units. A silicon-29 nuclear magnetic resonance spectrum indicated the 1,4-phenylenesiloxanesiloxane:methyl(3,3,3-trifluoropropyl)disiloxane ratio to be 1:5.7 as compared to the theoretical of 1:6. Based on gel permeation chromatography (GPC), its weight averaged molecular weight was 151,297; number average molecular weight=90,292; differential scanning calorimetry (DSC) indicated a Tg of −63°C, measured at a rate of 10°C per minute.
EXAMPLE 2

A viscous red sealant paste useful in making condensation curable sealant compositions was prepared as follows from a block copolymer made in accordance with the method of Example 1. The block copolymer was initially compounded in a Baker Perkins double-planetary mixer for 1 hour under a nitrogen blanket with a silazane/octamethylycyclotetrasiloxane treated fumed silica, and red iron oxide. A homogeneous filler dispersion was prepared based on the use of 1324 parts of block copolymer, 105 parts BTMSEB, 139 parts of a bis(alkoxyalkoxy)siloxane hydrocarbon as fuel resistant sealing materials, and 94 parts of a bis(trimethoxysilyl)ethane as fuel resistant materials.

Another test period, “B”, was heat aging at 177°C for 7 days after standard cure. The following results show that the compositions were resistant to degradation and were suitable for use in applications requiring chemically resistant materials.

Several neutral condensation curable sealant compositions were prepared by blending in a Semco 388 mixer, 4 parts of a bis(polyalkoxyalkoxy)silyl C_{2-10} hydrocarbon cross-linker and 100 parts of the above block copolymer red sealant paste contained in a Semco cartridge. The respective neutral condensation curable sealant compositions are identified in the tables below by the particular bis(polyalkoxyalkoxy)silyl Q cross-linker, where Q is as previously defined. More particularly, the bis(polyalkoxyalkoxy)silyl Q cross-linkers include 1,2-bis(triethoxysilyl)ethane, or “BTSE”, 1,4-bis(trimethoxysilyl)hexane, or “BTMSHEX”, and 1,4-bis(trimethoxysilyl)ethylene, or “BTMSEB”.

There was also mixed with the respective red sealant paste blends of block copolymer and cross-linker, 0.5 part of dibutyltin dilaurate, or “DBTDA”, as a condensation catalyst. Test specimens were prepared by compression molding the respective catalyzed formulations at ambient temperatures into 4 inchx5 inchx0.08 inch sheets. The sheets were allowed to cure to the elastomeric state for a minimum of 7 days at 25°C and 50% relative humidity (RH).

Tensile properties, such as tensile strength (psi), elongation (%), and 100% modulus (psi) of each of the respective compression molded elastomeric sheets were then measured after a series of seven day test periods. The initial test period, “A”, was standard cure at room temperature at 50% RH, while the second test period, “B”, was heat aging at 177°C for 7 days after standard cure. There was also included, “C”, immersion in Jet A fuel at 60°C for 7 days after standard cure, and, “D”, 2 days in Jet A at 60°C, followed by 5 days in air at 177°C after standard cure. The following results were obtained, where the respective cross-linked condensation cured compositions are denoted by their previously identified cross-linkers, namely BTMSEB, BTSE and BTMSHEX:

7 Day Test Periods

<table>
<thead>
<tr>
<th>Tensile Strength (psi)</th>
<th>Elongation (%)</th>
<th>100% Modulus</th>
</tr>
</thead>
<tbody>
<tr>
<td>BTMSEB</td>
<td>273</td>
<td>213</td>
</tr>
<tr>
<td>BTSE</td>
<td>239</td>
<td>185</td>
</tr>
<tr>
<td>BTMSHEX</td>
<td>268</td>
<td>199</td>
</tr>
<tr>
<td>BTMSEB</td>
<td>151</td>
<td>139</td>
</tr>
<tr>
<td>BTSE</td>
<td>146</td>
<td>145</td>
</tr>
<tr>
<td>BTMSHEX</td>
<td>192</td>
<td>92</td>
</tr>
<tr>
<td>BTMSEB</td>
<td>291</td>
<td>212</td>
</tr>
<tr>
<td>BTSE</td>
<td>253</td>
<td>186</td>
</tr>
<tr>
<td>BTMSHEX</td>
<td>275</td>
<td>191</td>
</tr>
</tbody>
</table>

The above results show that the copolymers of the present invention are depolymerization resistant and provide neutral condensation curable compositions which are useful as solvent resistant sealants.

The crosslinker in the present invention can be employed in non-corrosive cure systems to fabricate elastomers useful as fuel resistant scaling materials, electronic encapsulation, and in applications requiring chemically resistant materials. Particularly, the crosslinkers can be used for preparation of fuel tank sealants.

While typical embodiments have been set forth for the purpose of illustration, the foregoing description should not be deemed to be a limitation on the scope of the invention. Accordingly, various modifications, adaptations, and alternatives may occur to one skilled in the art without departing from the spirit and scope of the present invention.

What is claimed is:

1. A neutral condensation curable poly(fluoroorgano)siloxane-poly(silarylene)-siloxane block copolymer sealant composition comprising
   (a) a poly(fluoroorgano)siloxane-poly(silarylene)-siloxane block copolymer,
   (b) a cross-linker, and
   (c) a condensation catalyst, wherein the sealant composition has a glass transition temperature not exceeding about -54°C.

2. The sealant composition in accordance with claim 1, wherein the cross-linker comprises 1,4-bis(trimethoxysilyl)ethylene, or “BTSE”, and 1,6-bis(trimethoxysilyl)hexane, or combinations thereof.

3. The sealant composition in accordance with claim 2, wherein the cross-linker comprises 1,2-bis(tribromomethyl)ethylene.

4. A neutral condensation curable poly(fluoroorgano)siloxane-poly(silarylene)-siloxane block copolymer sealant composition comprising
   (a) a poly(fluoroorgano)siloxane-poly(silarylene)-siloxane block copolymer,
   (b) a cross-linker, and
   (c) a condensation catalyst, wherein the copolymer comprises the reaction product of a bis(diorganohydroxy-silyl)arylene chemically combined with a poly(fluoroalkylorgano)cyclopolsiloxane.

5. The sealant composition in accordance with claim 4, wherein the poly(fluoroalkylorgano)cyclopolsiloxane is present in a range between about 0.5 moles and about 4 moles, per mole of bis(diorganohydroxy-silyl)arylene.

6. The sealant composition in accordance with claim 4, wherein the poly(fluoroalkylorgano)cyclopolsiloxane is present in a range between about 0.5 moles and about 4 moles, per mole of bis(diorganohydroxy-silyl)arylene.
present in a range between about 1 mole and about 2 moles, per mole of bis(diorganohydroxysilyl)arylene.

7. The sealant composition in accordance with claim 4, wherein the bis(diorganohydroxysilyl)arylene comprises the formula

\[(\text{HO}-(\text{R})_{2}\text{Si})_{2}Q',\]

where \(Q'\) is a \(\text{C}_{(6.12)}\) divalent aromatic organic radical, and \(R\) is a \(\text{C}_{(1.4)}\) alkyl radical.

8. The sealant composition in accordance with claim 7, wherein the bis(diorganohydroxysilyl)arylene comprises 1,4-bis(dimethylhydroxysilyl)benzene.

9. The sealant composition in accordance with claim 4, wherein the poly(fluoroalkylorgano)cyclopolsiloxane comprises the formula

\[\text{[(RO)}_{2}\text{Si}]_{2}Q,\]

where \(Q\) is a \(\text{C}_{(2.12)}\) divalent organic radical, \(R\) is a \(\text{C}_{(1.4)}\) alkyl radical, and \(X\) is a member selected from the group consisting of \(R\) and \(RO\).

10. A neutral condensation curable poly(fluoroorgano)siloxy-poly(silarylene)siloxane block copolymer sealant composition comprising

(a) a poly(fluoroorgano)siloxane-poly(silarylene)siloxane block copolymer,
(b) a cross-linker,
(c) a condensation catalyst, and
(d) a heat stabilizer,

wherein the heat stabilizer is present in a range between about 0.1 and about 10 parts by weight per 100 parts of the copolymer.

11. A neutral condensation curable poly(fluoroorgano)siloxy-poly(silarylene)siloxane block copolymer sealant composition comprising

(a) a poly(fluoroorgano)siloxane-poly(silarylene)siloxane block copolymer,
(b) a cross-linker, and
(c) a condensation catalyst,

wherein the cross-linker comprises the formula

\[\text{[(RO)}_{2}X\text{Si}]_{2}Q,\]

where \(Q\) is a \(\text{C}_{(2.12)}\) divalent organic radical, \(R\) is a \(\text{C}_{(1.4)}\) alkyl radical, and \(X\) is a member selected from the group consisting of \(R\) and \(RO\).

12. A neutral condensation curable poly(fluoroorgano)siloxy-poly(silarylene)siloxane block copolymer sealant composition comprising

(a) a poly(fluoroorgano)siloxane-poly(silarylene)siloxane block copolymer,
(b) a cross-linker,
(c) a condensation catalyst, and
(d) a heat stabilizer,

wherein the heat stabilizer is present in a range between about 0.1 and about 10 parts by weight per 100 parts of the copolymer.

13. A neutral condensation curable poly(fluoroorgano)siloxy-poly(silarylene)siloxane block copolymer sealant composition comprising

(a) a poly(fluoroorgano)siloxane-poly(silarylene)siloxane block copolymer,
(b) a cross-linker,
(c) a condensation catalyst, and
(d) a heat stabilizer,

wherein the heat stabilizer comprises iron oxide.

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