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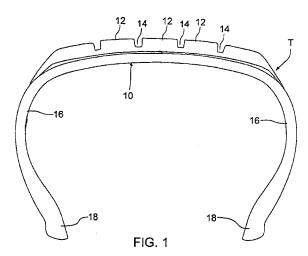
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(57) Abstract: A rubber composition including guayule rubber and 15 to 90 phr silica. The rubber can be cis-1,4 polyisoprene and include 15 to 90 phr silica and contain less than 20% of the proteins associated with Hevea rubber and further include at least a trace of resins present in other natural rubbers. The rubber can be cis-1,4 polyisoprene and further include between at least a trace and 3% by weight of resins present in guayule rubber. The rubber composition can further be comprised of a synthetic rubber and include between about 40-100 phr of guayule rubber, and silica. The silica can be hydrophobated.



## NATURAL RUBBER COMPOUNDS WITH SILICA AND USE WITH TIRES

### **BACKGROUND**

The present exemplary embodiment relates to a method for producing a rubber including silica and natural rubber, and the rubber produced thereby. It finds particular application in conjunction with guayule rubber, and will be described with reference thereto. However, it is to be appreciated that the present exemplary embodiment is also amenable to other similar natural rubber materials. It finds particular suitability for use in association with tires and will be described in association therewith. However, it is to be appreciated that the present exemplary embodiments are also amenable to other applications, such as engineered product applications such as hoses and belts.

[0002] Natural rubber, derived from the plant Hevea brasiliensis is a core component of many consumer goods, including medical devices and tires. The United States has a strong reliance on natural rubber, primarily because synthetic alternatives cannot match the high performance properties of natural rubber required for many applications and tend to be expensive. Hevea brasiliensis rubber is a critical material for all types of tires but especially for heavy truck and aircraft tires. These types of tires have a very large percentage of natural rubber because of the low heat buildup of natural rubber based compounds. Similarly, passenger tires may advantageously include a significant amount of natural rubber in specific components such as sidewall and bead filler to provide low heat build-up and flex fatigue resistance.

Over 90% of the Hevea-derived natural rubber imported by the United States originates in Indonesia, Malaysia and Thailand. Natural rubber sources in these countries are under intense threat from potential diseases and blights due to the genetic similarity of the rubber plants. Furthermore, the crop is limited to a restricted geographic area and labor-intensive harvesting methods. In addition, the Southeast Asian natural rubber crop contains many protein contaminants which are responsible for Type-I latex allergies, which are estimated to affect as many as 20 million Americans. The high cost of importation to the United States, as well as the potential for the entire crop to be wiped out by disease and the ubiquity of latex allergies, make non-allergenic domestic natural rubber alternatives particularly attractive.

[0004] Accordingly, attention is being directed to the production of natural rubber from plants such as guayule (Parthenium argentatum) and Russian dandelion (Taraxacum kok-saghyz), which yield polymeric cis 1,4-isoprene essentially identical to that produced by Hevea rubber trees in Southeast Asia. A major difference between Hevea rubber (HR) and guayule rubber (GR) is in the amount and types of proteins contained in each species. Hevea rubber possesses many different types of proteins with a 14-kDa "rubber elongation factor" and a 24-kDa "small rubber particle protein" (SRPP) dominating. Both of these are known allergens. In guayule, there are few proteins. A 53-kDa monoxygenase P450 (an allene oxide synthase) comprises about 50% of the rubber particle protein. (See M. Whalen, C. McMahan and D. Shintani, "Development of Crops to Produce Industrially Useful Natural Rubber, Isoprenoid Synthesis in Plants and Microorganisms: New Concepts and Experimental Approaches. pages 329-345, T. Bach and M. Rohmer eds, 2013; the disclosure of which is herein incorporated by reference). Thus, although the chemical structure of the rubber in both species are similar (cis-1,4 polyisoprene), the overall composition of the rubbers is not the same. A comparison of three different types of rubber - Hevea, Guayule, and TSK "Dandelion rubber" is shown in the table below.

Table

Protein sequences	Presence in	rubber particles	
Function	Hevea	Guayule	Russian dandelion
Rubber-associated	THE REAL PROPERTY OF THE PARTY		
Small rubber particle protein	+		+
Rubber elongation factor	+	***	+
Allene oxide synthase		+	4
Defense- or stress-related	terdag ong Elicioth (first a commission described a men a s <del>edi</del> ation at a tip described a m	a meneral de anno anno en emitente d'an de de de en 1900 de de de de desarrous alemandes de entre de la compan	, et de français au de français de la company de la co
Lipoxygenase, chitinase, PGase inhibitor	and the second s	+	+
Proteases, protease inhibitors		+	+
Phospholipase C, lipases, peroxidase, acid phosphatase	+		+
Dehydration-, wound-, stress-inducible proteins	+		
Annexin	+		+
Endomembrane-associated	+	4	+
Mitochondria-associated		+	+

[0005] The use of silica as a filler in tire rubbers has grown tremendously since its introduction into main line tires in the early 1990's. It has particularly been used in association with synthetic styrene-butadiene and polybutadiene rubbers. This is due to up to a 30% reduction in rolling resistance and up to a 15% increase in wet adhesion in poor weather conditions when a silica filler is used instead of carbon black. To achieve

these advantages, it is beneficial to react the silica with a silane coupling agent either before it is mixed into the rubber (an ex-situ process) or during the mixing of the rubber with the silica (in-situ process). Almost all commercial tire compounds using silica use the in-situ process for preparation of silica compounds.

The in-situ silica/silane technology has not been widely adopted in natural rubber compounds due to problems with rolling resistance, wear, handling and tear strength. This may be due to the presence of non-rubber materials in the natural rubber not present in the styrene-butadiene or polybutadiene rubber used in passenger tires. It may also be that these materials, especially the proteins and protein metabolites, may interfere with the silanization reaction to reduce overall physical properties and adversely affect processing. Investigators have tried to overcome these shortcomings by using pretreated (ex-situ) silica such as the Agilon family of products (see Justin Martin and Timothy Okel – "Bringing innovation to the surface: Functionalized slicas for improved natural rubber truck tire vulcanizates" – Technical paper at 184<sup>th</sup> Rubber Division meeting October 8, 2013) in natural rubber compounds, but this is an extremely costly solution. There remains a need to be able to improve the incorporation of silica/silane into natural rubber compounds using conventional in-situ silica mixing technology.

Ideally, very high molecular weight polymers incorporating various functional groups could be used in compounds to achieve maximum performance, but in many cases it is difficult to mix these types of materials. There is always a tradeoff between tire performance and the ability to mix the desired ingredients. There are several ways to measure the processability of compounds. One of the more common techniques is to measure the compounds Mooney viscosity. For the most part, the lower the compound Mooney viscosity, the better the processability. A less common approach, but one that can be applied to more tire building operations is capillary rheometery. In this test, a rubber sample is subjected to various shear rates, and a plot of polymer viscosity versus shear rates is obtained. Since different plant operations operate at different shear rates, the plots generated from the capillary rheometer can be used to predict performance for all of the plant operations, including extrusion, mixing and calendaring.

### **BRIEF DESCRIPTION**

[0008] The exemplary embodiment has been described with reference to the preferred embodiments. Obviously, modifications and alterations will occur to others upon reading and understanding the preceding detailed description. It is intended that the exemplary embodiment be construed as including all such modifications and alterations insofar as they come within the scope of the appended claims or the equivalents thereof.

[0009] According to a first embodiment, a rubber composition including guayule rubber and 15 to 90 phr silica is disclosed. The rubber can be cis-1,4 polyisoprene and include 15 to 90 phr silica and contain less than 20% of the proteins associated with Hevea rubber and further include at least a trace of resins present in other natural rubbers. In certain embodiments the Hevea protein content can be less than 5% or even less than 2%. The rubber can be cis-1,4 polyisoprene and further including between at least a trace and 3% by weight of resins present in guayule rubber. The rubber composition can further be comprised of a synthetic rubber and include between about 40-100 phr of guayule rubber, and silica. The silica can be hydophobated.

### BRIEF DESCRIPTION OF THE DRAWINGS

[0010] FIGURE 1 illustrates a tire in cross-section suitable for construction using the rubber compounds of this disclosure;

[0011] FIGURE 2 graphically illustrates the effect of oil type on viscosity for silica tread compounds (100 phr R) demonstrating lower viscosity for guayule compounds and a higher effect of oil on Hevea compounds;

[0012] FIGURE 3 graphically illustrates the effect of filler type on viscosity in sidewall compounds (40 phr R) showing guayule and Hevea black compounds had similar viscosities, black compounds had lower viscosity than silica compounds, and guayule silica compounds had lower viscosity than Hevea silica compounds;

[0013] FIGURE 4 graphically illustrates the effect of oil type on viscosity in silica containing sidewall compounds (40 phr R) demonstrating that a lower level of natural rubber in a recipe leads to smaller viscosity differences, guayule compounds had lower viscosity than Hevea compounds, and no significant effect of oil type on viscosity;

[0014] FIGURE 5 graphically illustrates the effect of filler type on viscosity of apex compounds (60 phr R) wherein silica compounds had higher viscosities than black compounds, Hevea silica compounds had significantly higher viscosities than guayule compounds, and silica containing Hevea and guayule black compounds had similar viscosities;

[0015] FIGURE 6 graphically illustrates the effect of oil type on viscosity in silica apex compounds (60 phr R) demonstrating that Hevea silica had a higher viscosity than guayule silica;

[0016] FIGURE 7 plots interactions between polymer, filler, polymer-filler, and oil level on Mooney viscosity in tread compounds;

[0017] FIGURE 8 plots interactions between polymer, filler, polymer-filler, and oil level on Mooney viscosity in apex compounds;

[0018] FIGURE 9 plots interactions between polymer, filler, polymer-filler, and oil level on Mooney viscosity in sidewall compounds; and

[0019] FIGURE 10 plots interactions for all stocks combined on Mooney viscosity.

### **DETAILED DESCRIPTION**

[0020] In the description of this invention, the terms "compounded" rubber compositions and "compounds" where used refer to the respective rubber compositions which have been compounded with appropriate compounding ingredients such as, for example, carbon black, oil, stearic acid, zinc oxide, silica, wax, anti-degradants, resin(s), sulfur accelerator(s), and silica coupler where appropriate. The terms "rubber" and "elastomer" may be used interchangeably. The amounts of materials are usually expressed in parts of material per 100 parts of rubber polymer by weight (phr).

The processing of plants for the extraction of biopolymers such as natural rubber latex can be achieved using chemical and/or mechanical processing. The methods utilized typically follow the general steps of: pre-grinding, wet milling, filtration, clarification, separation of liquid phases, purification, creaming, and concentration. Exemplary harvesting/processing techniques are described in US Patent 7,923,039 and US Patent Publication 2008/0015336, the disclosures of which are herein incorporated by reference.

[0022] Non-limiting examples of plant materials that can be used include, but are not limited to, guayule plant (Parthenium argentatum), gopher plant (Euphorbia lathyris), mariola (Parthenium incanum), rabbitbrush (Chrysothamnus nauseosus), milkweeds (Asclepias L.), goldenrods (Solidago), pale Indian plantain (Cacalia atripilcifolia), rubber vine (Crypstogeia grandiflora), Russian dandelions (Taraxacum kok-saghyz), mountain mint (Pycnanthemum incanum), American germander (Teucreum canadense) and tall bellflower (Campanula americana).

[0023] In practice, pneumatic rubber tires conventionally have relatively thin rubber sidewalls, a relatively thicker rubber tread, and a rubber inclusive bead, which are normally expected to be able to be subject to significant punishment under typically operating conditions by undergoing considerable dynamic distortion and flexing, abrasion due to scuffing, fatigue cracking and weathering such as, for example, atmospheric ozone aging. The rubber compounds of this disclosure are suitable for each of these applications.

[0024] With reference to FIG. 1 there is shown a tire T having a crown 10 with external treads 12 and grooves 14. In cross-section the tire T has the crown 10 extending radially outwardly along an arcuate path to a pair of oppositely disposed sidewalls 16 which define the maximum radial extent of the tire T. The sidewalls 16 curve inwardly from such maximum radial extent to a narrower area terminating at a pair of oppositely disposed beads 18. The rubber compounds of this disclosure are suitable for use in construction of either or all of the tread, sidewalls, and beads of a tire.

[0025] In accordance with this invention, a tire including a rubber is provided wherein the rubber composition is comprised of: (A) natural non-Hevea cis-1,4-polyisoprene rubber; (B) from about 15 to about 90, alternately from about 20 to about 80, and alternately from about 25 to about 70 phr of particulate reinforcing fillers comprised of: (1) amorphous synthetic silica, such as precipitated silica, and, optionally (2) rubber reinforcing carbon black. The composition can further optically include (C) an emulsion polymerized styrene/butadiene or a solution polymerized styrene/butadiene rubber or polybutadiene rubber.

[0026] The styrene/butadiene copolymer and/or polybutadiene may be prepared, for example, using conventional aqueous emulsion copolymerization of styrene and/or 1,3-butadiene monomers in an aqueous medium in the presence of a catalyst system.

[0027] The silica can be hydrophobated. More specifically, the silica can have a material attached to its surface to enhance incorporation into the rubber. Hydrophobating materials, processes of hydrophobating, and processes of making a rubber masterbatch and rubber products are disclosed in U.S. Published Application 2006/0137575 A1 (in-situ) and U.S. Patent 8,865,799 (ex-situ), the disclosures of whichare herein incorporated by reference. Also incorporated herein by reference is "Optimization of Rubber Formulations for Silica-Reinforced Rubber Compounds", Rubber Chemistry and Technology, Vol. 86, No. 2, pp. 313-329 (2013) by W. Kaewsakul, K. Sahakaro, W.K. Dierkes, and J.W.M. Noordermeer.

[0028] The silica may be used in conjunction with a silica coupler to couple the silica to the elastomer(s), to thus enhance its effect as reinforcement for the elastomer composition. Use of silica couplers for such purpose are well known and typically have a moiety reactive with hydroxyl groups (e.g. silanol groups) on the silica and another moiety interactive with the elastomer(s) to create the silica-to-rubber coupling effect.

The coupling agent for said silica reinforcement may be, for example, (A) [0029] a bis-(3-triakloxysilylalkyl) polysulfide such as, for example, a bis-(3-triethoxysilylpropyl) polysulfide, or (B) a bis-(3-triethoxysilylpropyl) polysulfide having an average of from about 2 to about 2.6 connecting sulfur atoms in its polysulfidic bridge or a bis-(3triethoxysilylpropyl) polysulfide having an average of from about 3.4 to about 4 connecting sulfur atoms in its polysulfidic bridge, wherein said polysulfide having an average of from 2 to about 2.6 connecting sulfur atoms in its polysulfidic bridge (to the exclusion of such polysulfide having an average of from 3 to 4 connecting sulfur atoms in its polysulfidic bridge) is blended with said rubber composition in the absence of sulfur and sulfur vulcanization accelerator and wherein said polysulfide having an average of from about 3.4 to about 4 connecting sulfur atoms in its polysulfidic bridge is thereafter blended with said rubber composition in the presence of sulfur and at least sulfur vulcanization accelerator, or (C) an organoalkoxymercaptosilane. Representative examples of various organoalkoxymercaptosilanes are, for example, triethoxy mercaptopropyl silane, trimethoxy mercaptopropyl silane, methyl dimethoxy mercaptopropyl silane, methyl diethoxy mercaptopropyl silane, dimethyl methoxy mercaptopropyl silane, triethoxy mercaptoethyl silane, tripropoxy mercaptopropyl silane, ethoxy dimethoxy mercaptopropylsilane, ethoxy diisopropoxy mercaptopropylsilane,

ethoxy didodecyloxy mercaptopropylsilane and ethoxy dihexadecyloxy mercaptopropylsilane.

[0030] The coupling agent may, for example, be added directly to the elastomer mixture or may be added as a composite of precipitated silica and such coupling agent formed by treating a precipitated silica therewith or by treating a colloidal silica therewith and precipitating the resulting composite.

[0031] It is readily understood by those having skill in the art that the rubber compositions can be compounded by methods generally known in the rubber compounding art, such as mixing the various sulfur-vulcanizable constituent rubbers with various commonly used additive materials such as, for example, curing aids, such as sulfur, activators, retarders and accelerators, processing additives, such as oils, resins including C5-C9 resins, C5 resins, coumarone-indene resins, terpene resins and terpene-phenolic resins, the aforesaid silica, and other plasticizers, fillers, pigments, fatty acid, zinc oxide, microcrystalline waxes, antioxidants and antiozonants, peptizing agents and carbon black reinforcing filler. As known to those skilled in the art, depending on the intended use of the sulfur vulcanizable and sulfur-vulcanized material (rubbers), the additives mentioned above are selected and commonly used in conventional amounts.

[0032] The vulcanization can be conducted in the presence of a sulfur-vulcanizing agent. Examples of suitable sulfur vulcanizing agents include elemental sulfur (free sulfur) or sulfur donating vulcanizing agents, for example, an amine disulfide, polymeric polysulfide or sulfur olefin adducts. Preferably, the sulfur-vulcanizing agent is elemental sulfur.

[0033] Accelerators are used to control the time and/or temperature required for vulcanization and to improve the properties of the vulcanizate. In one embodiment, a single accelerator system may be used, i.e., primary accelerator. In another embodiment, combinations of two or more accelerators in which the primary accelerator is generally used in the larger amount, and a secondary accelerator which is generally used in smaller amounts in order to activate and to improve the properties of the vulcanizate. Combinations of these accelerators have been known to produce a synergistic effect on the final properties and are somewhat better than those produced by use of either accelerator alone. In addition, delayed action accelerators may be used

which are not affected by normal processing temperatures but produce satisfactory cures at ordinary vulcanization temperatures. Suitable types of accelerators that may be used in the present invention are amines, disulfides, guanidines, thioureas, thiazoles, thiurams, sulfenamides, dithiocarbamates and xanthates. Preferably, the primary accelerator is a sulfenamide. If a second accelerator is used, the secondary accelerator is preferably a guanidine, dithiocarbamate or thiuram compound. One description of a suitable rubber processing methodology is provided in GB 1,400,930, incorporated herein by reference.

In general, compounds with guayule rubber (GR) in place of Hevea rubber (HR) had similar processing results as found in the corresponding black compounds. Surprisingly however, in silica compounds, GR compounds had lower Mooney viscosity, complex torque, elastic torque and high tan  $\delta$ . In capillary rheometry, GR compounds had similar viscosities to HR compounds with black and again surprisingly lower viscosities with silica. These results indicate that compounds with GR have better processability than their HR equivalents in silica compounds and similar processability in black compounds. GR compounds were generally less affected by changing the filler or oil type suggesting that the processability of GR compounds is less likely to be affected by compounding changes that HR compounds.

[0035] With silica as the filler, GR compounds had lower Mooney viscosity with naphthenic oil than those with NR and naphthenic oil in recipes A and B (see below). GR compounds with silica had lower S\* and higher tanð than those with HR and silica in recipe A with both oils and in recipe C with TDAE. With silica as the filler, GR compounds with both oil types had lower S' than equivalent HR compounds in recipe A. In recipe C, this was only observed with TDAE. These results indicate that in recipe A, changing the polymer from HR to GR decreases the viscosity and improves the processability of the rubber. In recipes B and C this is dependent on the oil type. In recipe B GR compounds with naphthenic oil and recipe C GR compounds with TDAE have better processability than their HR equivalents. In silica compounds, increasing naphthenic oil had a slight effect of decreasing Mooney viscosity, S\*, and S' with HR and no effect with GR. Increasing the level of TDAE in silica compounds decreased Mooney viscosity S\*, and S' and increased them with GR. This indicates that GR

interacts with oils differently than HR and its processability is not necessarily improved by an increase in oil.

[0036] For purposes of this disclosure a compound with better processability is considered as having a lower Mooney viscosity. A reference to Mooney (ML 1+4) viscosity of an elastomer or sulfur vulcanizable polymer represents the viscosity of the respective elastomer or sulfur vulcanizable polymer in its uncured state. The Mooney (ML 1+4) viscosity at  $100^{\circ}$ C. uses a one minute warm up time and a four minute period of viscosity measurement, a procedural method well known to those having skill in such art. Processability of compounds will also be improved if the tan  $\delta$  of the green compound is higher, or if the viscosity as a function of shear (as observed in a capillary rheometer plot) is lower.

[0037] A tire can be built, shaped, molded and cured from the presently disclosed composition by various methods which will be readily apparent to those having skill in such art.

[0038] The invention may be better understood by reference to the following example in which the parts and percentages are by weight unless otherwise indicated.

### Examples:

[0039] The elastomers used were SIR-20 for the Hevea rubber compounds and Guayule rubber isolated from latex and purchased from the Yulex Corporation. Hevea and Guayule compounds were made according to the formulas in recipes A (Tread), B (Sidewall) and C (Apex). Mooney viscosity was determined using ASTM D1649, and capillary rheometery data was obtained using ASTM D5099. All testing was performed at 100°C.

Reci	pe A	- T	read
1 1001	$\circ \circ \circ \circ$		JOUG

Component (phr)	A1	A2	АЗ	A4	A5	A6	Α7	A8
SIR-20	100	100	100	100				
GR			—		100	100	100	100
N234	50	50			50	50	_	
Ultrasil 7000	_	_	50	50		<del></del>	50	50
TESPD			4.3	4.3	—	_	4.3	4.3
Naphthenic oil	5		5	_	5	_	5	
TDAE		5		5		5		5
Stearic acid	1	1	1	1	1	1	1	1
Zinc oxide	5	5	5	5	5	5	5	5
TMQ	2	2	2	2	2	2	2	2
Sulfur	1.4	1.4	1.4	1.4	1.4	1	1.4	1.4
TBBS	1	1	1	1	1	1	1	1

## Recipe B - Sidewall

Component (phr)	B1	B2	B3	B4	B5	B6	B7	B8
SIR-20	40	40	40	40		_		
GR					40	40	40	40
BR 1280	60	60	60	60	60	60	60	60
N550	50	50	_	_	50	50		
Zeosil 1085	_		62	62	_		62	62
TESPD			5.3	5.3	_	_	5.3	5.3
Naphthenic oil	7		7	_	7	_	7	
TDAE		7	_	7		7	<del></del>	7
Stearic acid	1.5	1.5	1.5	1.5	1.5	1.5	1.5	1.5
Zinc oxide	3	3	3	3	3	3	3	3
TMQ	1	1	1	1	1	1	1	1
DAPD	1	1	1	1	1	1	1	1
6-PPD	3	3	3	3	3	3	3	3
Sulfur	1.5	1.5	1.5	1.5	1.5	1.5	1.5	1.5
CBS	1	1	2.3	2.3	1	1	2.3	2.3

Recipe C - Apex

Component (phr)	CI	C2	C3	C4	C5	C6	C7	C8	C9	C10	C11	C12	C13	C14	C15	C16
SIR-20	60	60	60	60	60	б0	60	60								
GR			-	******					60	60	60	60	60	60	60	60
SBR 1502	40	40	40	40	40	40	40	40	40	40	40	40	40	40	40	40
N326 carbon black	80	80	80	80					80	80	80	80			******	
Ultrasil 7000					80	80	80	80		******			80	80	80	80
TESPD					6.8	6.8	б.8	6.8					6.8	6.8	б.8	6.8
Naphthenic oil	8	12			8	12			8	12			8	12		
TDAE			8	12			8	12			8	12			8	12
Stearic acid	2	2	2	2	2	2	2	2	2	2	2	2	2	2	2	2
Zinc oxide	3	3	3	3	3	3	3	3	3	3	3	3	3	3	3	3
TMQ	2	2	2	2	2	2	2	2	2	2	2	2	2	2	2	2
P-tert-octylphenol formaldehyde resin	3	3	3	3	3	3	3	3	3	3	3	3	3	3	3	3
Phenolic novalac resin	8	8	8	8	8	8	8	8	8	8	8	8	8	8	8	8
Hexamethy lenetetramine	0.8	0.8	0.8	8.0	8.0	0.8	8.0	0.8	1	0.8	8.0	0.8	8.0	0.8	8.0	8.0
Sulfur	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1
Insoluble polymeric sulfur	6.3	6.3	6.3	6.3	6.3	6.3	6.3	6.3	6	6.3	6.3	6.3	6.3	6.3	6.3	6.3
OBTS*	2	2	2	2	2	2	2	2	2	2	2	2	2	2	2	2

N-oxydiethylene-2-benzothiazole sulfenamide

[0040] Rubber compounds were mixed in a 300 cc Brabender mixer using the following mixing protocols.

### Carbon black compounds:

Mixing Step	Mixing Time (min)
Master Pass	
Add polymer, ½ of black, and additives	0:00
Increase rotor speed to 40 rpm	0:10
Increase rotor speed to 60 rpm	0:20
Add ½ of remaining black	1:00
Increase rotor speed to 80 rpm	1:15
Add oil	2:30
Ram sweep	4:00
Drop batch	5:00
Final Pass	
Add master pass and curatives	0:00
Increase rotor speed to 40 rpm	0:30
Ram sweep	1:30

Drop batch	2:30
Silica com	npounds
Mixing Step	Mixing Time (min)
Master Pass	
Add polymer, ½ of silica, and additives	0:00
Increase rotor speed to 40 rpm	0:10
Increase rotor speed to 60 rpm	0:20
Add ½ of remaining silica	1:00
Increase rotor speed to 80 rpm	1:15
Add oil	2:30
Ram sweep	3:30
Maintain temperature at 160-170°C	4:30
Drop batch	8:30
Final Pass	
Add master pass and curatives	0:00
Increase rotor speed to 40 rpm	0:30

### Compound Nomenclature for Capillary Rheometry

Ram sweep

Drop batch

Component	Hevea (H)	Guayule (G)
Polymer	Hevea (H)	Guayule (G)
Filler	Carbon black (B)	Silica (S)
Oil	Naphthenic (N)	TDAE (T)

1:30

2:30

[0041] In general, compounds with GR in place of HR had similar results in black compounds. In silica compounds, GR had lower Mooney viscosity, complex torque,

elastic torque and higher tanð. In capillary rhemoetry, GR compounds had similar viscosities to HR compounds with black and lower viscosities with silica. These results indicate that compounds with GR have better processability than HR equivalents in silica compounds and similar processability in black compounds. GR compounds were generally less affected by changing the filler or oil type suggesting that the processability of GR compounds is less likely to be affected by compounding changes than HR compounds. In short, silica compounds with GR have better processing characteristics than equivalent compounds with HR.

[0042] This written description uses examples to enable any person skilled in the art to make and use the disclosure. The patentable scope of the disclosure is defined by the claims, and may include other examples that occur to those skilled in the art. Such other examples are intended to be within the scope of the claims if they have structural elements that do not differ from the literal language of the claims, or if they include equivalent structural elements with insubstantial differences from the literal language of the claims. Moreover, this disclosure is intended to seek protection for a combination of components and/or steps and a combination of claims as originally presented for examination, as well as seek potential protection for other combinations of components and/or steps and combinations of claims during prosecution.

### [0043] CLAIMS:

1. A rubber composition comprising guayule rubber and 15 to 90 phr silica.

- 2. A rubber composition comprising cis-1,4 polyisoprene and 15 to 90 phr silica, said composition containing less than 20% of the proteins associated with Hevea rubber and further including at least a trace of resins present in other natural rubbers.
- 3. A rubber composition comprised of cis-1,4 polyisoprene and 15 to 90 phr silica and further including between at least a trace and 3% by weight of resins present in guayule rubber.
- 4. The rubber composition of claims 1-3 wherein the silica includes a silane coupling agent.
- 5. The rubber composition of claim 4 wherein the silane coupling agent is selected from the group including bis(triethoxysilypropyl)tetrasulfide (TESPT), bis(triethoxysilypropyl-y)disulfide (TESPD), 3-mercaptopropyl trimethoxy silane (A-189), and 3-mercaptoproppyl triethoxy silane (A-1891).
- 6. The rubber composition of claims 1-3 wherein the silica is pretreated with a hydrophobating agent prior to mixing with the rubber.
- 7. The rubber composition of claims 1-3 including at least a trace of allene oxide sythase.
- 8. A rubber composition of claims 1-7 where the surface area of the silica is between 175 and 90 m<sup>2</sup>/gm.
- 9. A rubber composition comprising synthetic rubber and between about 40-100 phr of guayule rubber and 15 to 90 phr silica.

10. The rubber composition of claim 9 wherein the silica is hydrophobated.

- 11. The rubber composition of claim 10 including carbon black.
- 12. The rubber composition of claim 10 including about 20 to 80 phr silica
- 13. The rubber composition of claim 10 including about 25 to 70 phr silica
- 14. A rubber composition comprising a synthetic rubber and between 40-100 phr of guayule rubber and between 50-80 phr silica.
- 15. Finished rubber articles containing the rubber composition of claims 1-14.
- 16. A tire containing the rubber composition of claims 1-14.
- 17. A tread component of a tire containing the rubber composition of claims 1-14.
- 18. A tread component of a truck tire containing the rubber composition of claims 1-
- 14.
- 19. A tread component of an aircraft tire containing the rubber composition of claims 1-14.
- 20. A rubber composition comprised of:
- (A) natural non-Hevea 1,4-polyisoprene rubber;
- (B) from about 15 to 90 phr silica; and
- (C) optionally a synthetic rubber;
- said composition having a Mooney (ML1+4) viscosity that is less than a comparative rubber composition that contains an equivalent amount of Hevea natural rubber instead of a non-Hevea 1,4-polyisoprene rubber natural rubber.
- 21. The composition of claim 20 wherein said natural non-Hevea rubber is guayule or Russian dandelion.

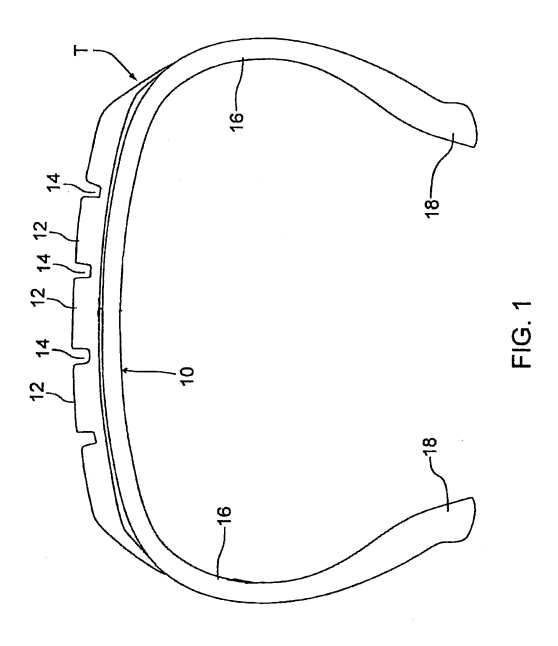
22. A method of forming a rubber compound comprising the steps of: hydrophobating silica, obtaining a non-Hevea natural rubber, mixing the hydrophobated silica with the rubber, and making a nonproductive compound, and wherein additional rubber may or may not be used in making the nonproductive compound; mixing curatives with the nonproductive compound to make a final compound; making a rubber item comprising the final compound; and vulcanizing the rubber item.

### 23. A method of forming a rubber compound comprising the steps of:

combining a non-Hevea natural rubber, a precipitated silica, a silica coupler, optionally a processing oil, and optionally a synthetic rubber in a mixer to form a nonproductive compound;

combining said nonproductive compound with curatives to obtain a final compound;

making a rubber item comprising the final coupound; and vulcanizing the rubber item.



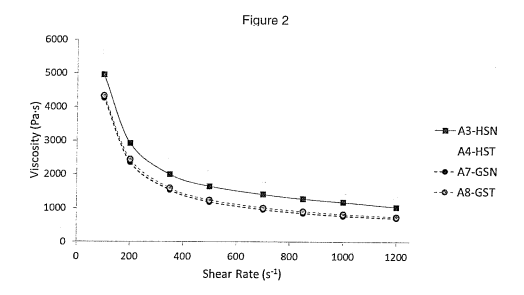
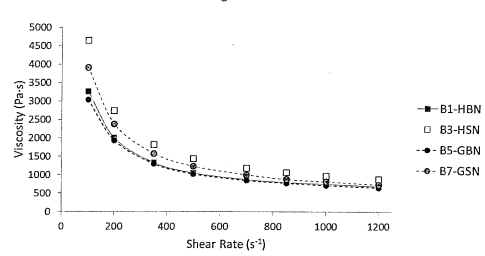
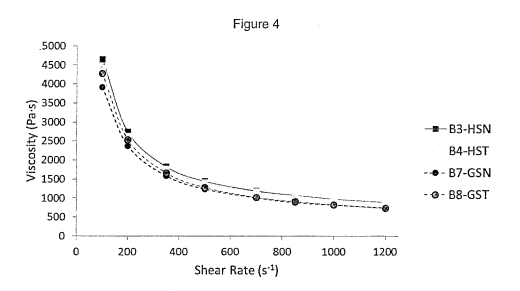


Figure 3





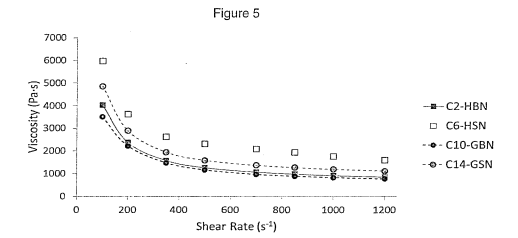


Figure 6

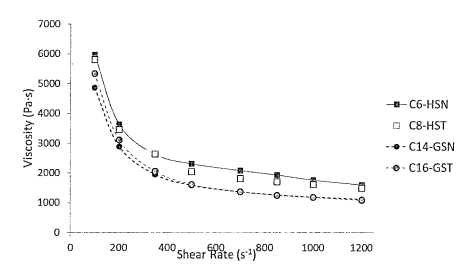


Figure 7

Interactions – Tread Compounds – Mooney Viscosity

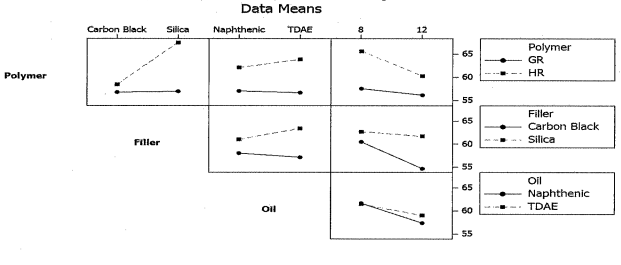
## Interaction Plot for Mooney 1+4 Data Means

Carbon Black Silica Naphthenic TDAE Polymer 64 GR --- HR 56 Polymer 48 Filler 64 Carbon Black Silica 56 Filler 48 Oil

Figure 8

Interactions – Apex Compounds – Mooney Viscosity

### Interaction Plot for Mooney 1+4



4/5

Oil Level

Figure 9

Interactions – Sidewall Compounds – Mooney Viscosity

## Interaction Plot for Mooney 1+4 Data Means

Silica Carbon Black Naphthenic TDAE 50 Polymer GR HR 45 Polymer - 40 - 50 Filler Carbon Black Silica 45 Filler Oil

Figure 10

Interaction – All Stocks Combined – Mooney Viscosity

# Interaction Plot for Mooney 1+4 Data Means

Carbon Black Naphthenic Silica TDAE Polymer 60 GR **----** HR 56 52 Filler 60 Carbon Black Silica Filler 56 52

Polymer

Oil

International application No. PCT/US2016/013043

#### CLASSIFICATION OF SUBJECT MATTER

C08K 3/36(2006.01)i, C08L 7/00(2006.01)i, B60C 1/00(2006.01)i

According to International Patent Classification (IPC) or to both national classification and IPC

#### FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols) C08K 3/36; C07F 7/08; C07F 7/10; C08L 9/08; C08J 3/215; C08K 5/541; C08K 5/5419; C08L 7/00; C08L 9/00; B60C 1/00

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Korean utility models and applications for utility models Japanese utility models and applications for utility models

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used) eKOMPASS(KIPO internal) & Keywords: rubber, silica, guayule, hydrophobated, cis-1,4 polyisoprene, synthetic rubber, curative

### DOCUMENTS CONSIDERED TO BE RELEVANT

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	US 2013-0253088 A1 (AGARWAL, S.) 26 September 2013 See paragraphs [0024], [0029], [0034], [0036], [0038]; and claim 1.	1-3,20,21,23
Y	See paragraphs [0024], [0024], [0004], [0000], and craim 1.	9-14,22
Y	CN 103772760 A (EVE RUBBER INSTITUTE CO., LTD.) 07 May 2014 See abstract; and claims 1, 4, 5.	9-14
Y	US 2010-0022684 A1 (WALLEN, P. J. et al.) 28 January 2010 See abstract; and claim 1.	10-13,22
A	US 2013-0203915 A1 (ARIGO, M. et al.) 08 August 2013 See claims 1, 11-14.	1-3,9-14,20-23
A	US 2013-0203940 A1 (SOTO, J.) 08 August 2013 See claims 1, 12.	1-3,9-14,20-23

l		Further documents are listed in the continuation of Box C.		See patent family annex.
	* "A"	Special categories of cited documents: document defining the general state of the art which is not considered to be of particular relevance	"T"	later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention

- "E" earlier application or patent but published on or after the international filing date
- document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)
- document referring to an oral disclosure, use, exhibition or other means
- document published prior to the international filing date but later than the priority date claimed
- nd
- "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone
- document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art
- "&" document member of the same patent family

Date of mailing of the international search report Date of the actual completion of the international search 13 May 2016 (13.05.2016) 12 May 2016 (12.05.2016)

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### INTERNATIONAL SEARCH REPORT

International application No.

### PCT/US2016/013043

Box No. II	Observations where certain claims were found unsearchable (Continuation of item 2 of first sheet)
This internat	ional search report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
	ims Nos.: ause they relate to subject matter not required to be searched by this Authority, namely:
bec ext	ims Nos.: 5 ause they relate to parts of the international application that do not comply with the prescribed requirements to such an ent that no meaningful international search can be carried out, specifically: aim 5 is unclear since it is referring to the multiple dependent claims which do not comply with PCT Rule 6.4(a).
	ims Nos.: 4,6-8,15-19 ause they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Box No. III	Observations where unity of invention is lacking (Continuation of item 3 of first sheet)
This Internat	ional Searching Authority found multiple inventions in this international application, as follows:
	all required additional search fees were timely paid by the applicant, this international search report covers all searchable ms.
	all searchable claims could be searched without effort justifying an additional fees, this Authority did not invite payment any additional fees.
	only some of the required additional search fees were timely paid by the applicant, this international search report covers y those claims for which fees were paid, specifically claims Nos.:
	required additional search fees were timely paid by the applicant. Consequently, this international search report is ricted to the invention first mentioned in the claims; it is covered by claims Nos.:
Remark on	Protest  The additional search fees were accompanied by the applicant's protest and, where applicable, the payment of a protest fee.  The additional search fees were accompanied by the applicant's protest but the applicable protest fee was not paid within the time limit specified in the invitation.  No protest accompanied the payment of additional search fees.

### INTERNATIONAL SEARCH REPORT

Information on patent family members

International application No.

Information on 1	PCT/U	PCT/US2016/013043				
Patent document cited in search report	Publication date	Patent family member(s)	Publication date			
US 2013-0253088 A1	26/09/2013	None				
CN 103772760 A	07/05/2014	WO 2015-109791 A1	30/07/2015			
US 2010-0022684 A1	28/01/2010	CA 2724798 A1 CA 2724798 C CN 102083892 A EP 2303951 A1 EP 2303951 A4 JP 2011-522925 A JP 2013-079400 A JP 5220189 B2 JP 5643350 B2 KR 10-1248051 B1 KR 10-2012-0068659 A US 2013-0192746 A1 US 2014-0083598 A9 US 8357733 B2 US 8865799 B2 WO 2010-011345 A1	28/01/2010 26/08/2014 01/06/2011 06/04/2011 03/12/2014 04/08/2011 02/05/2013 26/06/2013 17/12/2014 27/03/2013 27/06/2012 01/08/2013 27/03/2014 22/01/2013 21/10/2014 28/01/2010			
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