(54) Title: POSITIVE FRICTION CONTROL COMPOSITION FOR RAILWAYS

Fig. 1

- **Step 1**: Charge water, water-insoluble hydrocarbon (e.g., SUTROL 220), wetting agent (e.g., VAN WET 9NB) to a container, begin mixing.

- **Step 2**: Charge the rheology additive (e.g., VAN GEL B) slowly into the container.

- **Step 3**: Dispense under high shear to form a thixotropic gel.

- **Step 4**: Charge water soluble polyalcohol freezing point depressant (e.g., GLYCERINE) slowly while mixing.

- **Step 5**: Charge liquid or solid lubricant (e.g., SUPERFINE WOLVESLEN), liquid or solid friction modifier (e.g., BARIUM SULFATE, TALC), corrosion inhibitor (e.g., AMMOMIUM BENZATE), and biocide/fungicide agent (e.g., BENZISOTHIAZOLINE) in the order above.

- **Step 6**: Dispense until grind is 5–7 on a HEGMAN GAUGE.

(57) Abstract: A friction control composition having high and positive frictional properties for sliding steel surfaces includes a water insoluble hydrocarbon that enables a reduced water content, a rheological additive, a freezing point depressant, a friction modifier, and a lubricant.

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REFERENCE TO RELATED APPLICATIONS

[0001] The present utility patent application claims priority to the following provisional patent applications: Friction Modifier Composition for Rail Tracks, Serial Number 61/848,596 filed on January 7, 2013; Friction Modifier Composition for Rail Tracks, Serial Number 61/850,690, filed on February 21, 2013; Friction Modifier Composition for Rail Tracks, Serial Number 61/850,923, filed on February 26, 2013; Friction Modifier with High and Positive Characteristics, Serial Number 61/958,789 filed on August 6, 2013; Friction Modifier with High and Positive Characteristics, Serial Number 61,962,265 filed on November 4, 2013; and Friction Modifier with High and Positive Characteristics, Serial Number 61/963,448 filed on December 4, 2013.

FIELD OF INVENTION

[0002] The present invention relates to friction control compositions with high and positive Motional properties for controlling friction in a positive manner between two sliding steel surfaces, namely steel wheels on steel rails in the railway industry.

BACKGROUND OF THE INVENTION

[0003] Steel transport wheels and steel rails have many issues that require the control of friction between the two surfaces. Failure to control the friction in a positive manner can result in noise problems, extensive wear, and sub-optimum performance (from an efficiency standpoint) caused by slip-stick oscillations due to negative friction between the two sliding steel surfaces. Accordingly, friction modifiers with high and positive friction to control sliding and rolling-sliding contact in steel-rail and steel-wheel transportation systems are well known in the art. Various patents from Kelsan Technologies describe friction modifier compositions (see, US Patents Nos, 6,759,372; 6,855,673; 7,045,489; 7,160,378; 7,244,695 and 7,939,476). These patents describe a water-based system commercialized by Kelsan.
However, there are various problems with some of these water based friction modifier products. Examples of the problems are set out below:

When ammonia is used in the water-based systems it can cause equipment to rust. Latex can cause problems with the gears and other movable mechanical parts as the compositions dry out. These compositions can form a skin over the liquid when it is pumped onto the rail head and the product is allowed to dry before taken up by a train wheel in a short period of time, and this can cause flowing out or splashing when contacted by the wheels of the train and accordingly not be carried down the track by the wheels of the train.

What is needed is a friction modifier composition for top of rail applicators that help prevent vast on equipment with which it comes into contact.

Another object of the invention is to eliminate the latex skin on the prior art compositions such that the product when applied does not have a skin but instead forms a soft, non-drying deposit on the rail head. This soft non-drying deposit is picked up by the train wheel and carried down the rail to form a continuous film which controls the friction between the wheel and the rail in a positive manner.

In accordance with a further object of the invention, the friction control composition is embodied as a thixotropic gel or liquid that is thinned by shear and returns to its thicker more viscous state under static conditions. The thixotropy of the composition is used to facilitate application to the rail and to promote retention on the rail in its thicker state without formation of a skin. Unlike the prior art compositions discussed above, the composition does not form a skin over a low viscosity fluid that is ejected or pushed away by the rolling contact of the train wheels. Thus, the composition may be mixed for liquid-like flow during application to the rail as by conventional rail-mounted pump systems. Following rail application, the composition thickens under static conditions without drying or forming a skin, and remains positioned on the rail until sheared by train wheel engagement for distribution along the track during "carry down".

Another object of the invention is to have freezing point depressant that does not degrade the rheology of the composition.
Yet another object of the invention is to have a composition that causes the thickener (e.g. the clay) to go into a matrix such that it has improved dispersability.

Also, solid stick prior art compositions, such as the one described in, U.S. Pat. No. 4,915,856, are well known in the art. But these solid stick friction modifiers have their own problems such as expense, and they need mechanical brackets and applicators to apply the product to the wheel. With solid stick compositions, practicality of use and duration of efficacy can also be a problem on freight trains.

Another object of the invention is to change the sliding friction (mm negative to positive.

Another object of the invention is to reduce noise by reducing or eliminating slip-stick oscillations between the wheel and rail.

Another object of the invention is to reduce lateral creep, which reduces lateral forces by changing the friction from negative to positive between the wheel and rail when a train, especially a freight train, goes through a curve. The benefits of reducing lateral forces include increasing the stability of the train as it travels down the track and in a related manner there is a reduction of the wear on the rail head, rail ties, and tie plates. Also, the bogey or truck goes around the curve in a much smoother fashion with reduction in jerking and jumping movement.

Another object of the invention is to reduce longitudinal creep wherein the wheel is sliding forward such as occurs in a transit system when the wheel is going around a mild curve. In longitudinal creep, if the wheels go slightly off-kilter, the locomotive pulls the wheel and the wheel slides in the longitudinal direction, if this creep happens all the time, you get short pitch corrugations. These are wear marks on the rail head in the nature of corrugations as encountered in a dirt road. By reducing this creep, the wheel will not slide as far and short pitch corrugations are inhibited.
Another object of the invention is to reduce spin creep wherein there is instability between the wheel and the rail and the wheel is almost making a small circle on the top of the rail head.

The friction control compositions of the present invention reduce, if not eliminate, these three different types of creep by changing negative friction to positive friction. Lateral forces are one of the main problems in the heavy hard railroad in North America, and it is preferably reduced in accordance with the present invention. Similarly, longitudinal creep is reduced, if not eliminated, in order to inhibit the formation of short pitch corrugations in the rail. The reduction or elimination of spin creep is also desirable in order to reduce wear on the wheel and rail.

All of these creeps are small and are, for example, in the micron size range. The friction control compositions herein are effective to change the Motion from negative to positive and thereby reduce or eliminate creep and the accompanying stick-slip.

SUMMARY OF THE INVENTION

The present invention relates to novel friction control compositions. More particularly, the present invention relates to friction control compositions that may be applied to steel-rails or steel-wheels that are potentially in sliding or rolling-sliding contact with each other.

The friction control compositions change the friction, or coefficient of Motion, between the steel surfaces from negative to positive and thereby reduce or eliminate the lateral, longitudinal and/or spin creeps with a corresponding reduction or elimination of lateral forces and wheel-rail wear while increasing stability of the train.

In accordance with preferred embodiments of the invention, the skin forming reactivity agents of the KelSan patents, supra, are avoided since skin formation is believed to inhibit uniform iehixotropic properties, effective shear of the applied composition upon train wheel passage, and the achievement of improved carry down. Accordingly, the present compositions preferably rely upon the iehixotropic properties to facilitate application of the composition, maintenance of position of the applied composition and subsequent train wheel
shear to provide viscosities desirable for distribution of increased amounts of composition over longer carry down distances.

[0022] The preferred liquid embodiments of the friction control compositions include a reduced amount of water compared with prior art liquid compositions. The water content is reduced by the use of a water insoluble hydrocarbon found to further enhance the stability of the friction control composition.

[0023] The friction control compositions herein are described in greater detail with reference to illustrative compositions. Compositional percentages are in weight percent (w/w%) unless otherwise specified.

[0024] The inventive friction control compositions for use on top of rail applications comprise:

(a) from about 4 to about 40 w/w% water;
(b) from about 2 to about 20 w/w% rheology additive;
(c) from about 10 to about 40 w/w% water insoluble hydrocarbon;
(d) from about 10 to about 40 w/w% water soluble polyalkohol freezing point depressant;
(e) from about 1 to about 7 w/w% liquid or solid friction modifier; and
(f) from about 1 to about 20 w/w% liquid or solid lubricant

[0025] Optionally, the composition may also contain one or more of:

(g) from 1 to 3 w/w% surfactant or wetting agent
(h) from 0.1 to 0.5 w/w% corrosion inhibitor, and/or
(i) from 0.05 to 0.2 w/w% biocide/fungicide agent

[0026] In preferred embodiments, the friction control compositions consist essentially of the foregoing components and, accordingly, the formation of a skin and the skin forming retentivity agents of the Kelsan patents, supra, are preferably avoided in favor of the thixotropic properties in the present compositions. Thus, the preferred compositions herein are substantially free of the film-forming retentivity agents described in the Kelsan patents as
being selected from the group consisting of acrylic, polyvinyl alcohol, polyvinyl chloride, oxazoline, epoxy, alkyd, urethane acrylic, modified alkyd, acrylic latex, acrylic epoxy hybrids, polyurethane, styrene acrylate, and styrene butadiene based compounds.

[0027] In preferred compositions, the water insoluble hydrocarbon is selected from the group consisting of isoparaffins, vegetable oils, bio-based triglycerides and fatty oils.

[0028] According to the present invention, another embodiment of the friction control composition comprises:

(a) from 15 to 29 w/w% water
(b) from 4 to 13 w/w% rheology additive
(c) from 11 to 28 w/w% water insoluble hydrocarbon (e.g. isoparaffins, vegetable oils, bio-based triglycerides or fatty oils).
(d) from 22 to 40 w/w% freezing point depressant
(e) from 9 to 24 w/w % liquid or solid friction modifier
(f) from 1 to 6 w/w% liquid or solid lubricant,

[0029] As noted above, the composition may also optionally contain one or more of:

(g) from 1 to 3 w/w% surfactant or wetting agent
(h) from 0.1 to 0.5 w/w% corrosion inhibitor, and/or
(i) from 0.05 to 0.2 w/w% bioeide/fongkide agent

[0030] Because of some of the problems with water-based systems, various other compounds were experimented with as a partial replacement of water. It has been found that adding a water insoluble hydrocarbon to the composition (e.g., an isoparaffin such as SOTROL 220) helps depress the freezing point and also helps stabilize or even improve the rheology of the formulation. This is especially true when the water insoluble hydrocarbon is compared with other freezing point depressants such as glycerin. Other water insoluble hydrocarbons that have environmental advantages over isoparaffins are vegetable oils, bio-based triglycerides and fatty oils such as canola oil. The oils do not have the same freezing point advantages as isoparaffins but they are environmentally friendly.
The addition of the wafer insoluble hydrocarbon (either isoparaffins or oils) in the partially water based system is counterintuitive because one would have guessed that it would not mix well with the water and would in all likelihood separate. However, we believe that the clay has receptor sites that allow the water insoluble hydrocarbon to bind onto the clay and keep the final product homogenous. The result is a composition that may contain lower amounts of water and in the case of isoparaffins lower amounts of soluble polyalcohol freezing point depressants such as glycerine. As pointed out above, water based systems have problem with maintenance of the system and typical freezing point depressants can cause negative theology effects on the composition.

BRIEF DESCRIPTION OF THE DRAWINGS

Fig. 1 shows a diagram of the process for making one of the embodiments of the friction control compositions disclosed herein;

Fig. 2 is a histogram showing L/V ratio on the high rail of a 5.7° curve for trail axles of a dry wheel-rail system and a wheel-rail system comprising a friction control composition in accordance with the present invention;

Fig. 3 is a histogram similar to Fig. 2 showing L/V ratio on the high rail of a 5.7° curve for lead axles of a dry wheel-rail system and a wheel-rail system comprising a friction control composition in accordance with the present invention;

Fig. 4 is a histogram showing lateral force distribution for the trail axles of the wheel-rail systems of Fig. 2

Fig. 5 is a histogram similar to Fig. 4 showing lateral force distribution for the lead axles of the wheel-rail systems of Fig. 2;

Fig. 6 is a histogram similar to Fig. 2 showing the high rail I/V ratio for lead axles of the wheel-rail systems of Fig. 2 at a distance of 6.9 miles from point of application of the friction control composition; and
[0038] Fig. 7 is a histogram similar to Fig. 6 showing the low rail L/V ratio for lead axles of the wheel-rail systems of Fig. 2 at a distance of 6.9 miles from point of application of the friction control composition.

DETAILED DESCRIPTION OF THE INVENTION

[0039] Turning now to Figure 1, the composition can be made in a batch process by adding the various components and then mixing them. Details of various methods of making the compositions are laid out in the examples below.

£0040] Figure 1 shows the process steps graphically. In step 1, charge water, water-insoluble hydrocarbon (e.g. SOTROL 220) and a wetting agent (e.g. VAN WET 9N9) into a batch container. In step 2 slowly charge the rheology additive (e.g. VAN GEL B, limc) into the container. In step 3 this mixture is dispersed under high shear to form a thixotropic gel. In step 4 slowly charge a water soluble poiyaalcohol freezing point depressant (e.g. glycerine) while mixing the composition. Step 5 requires adding the following components into the mixture while stirring: liquid or solid lubricant (e.g. superfine molybdenum, carbon black); liquid or solid friction modifier (e.g. barium sulfate and talc); corrosion inhibitor (e.g. ammonium benzoate); and bioeide/rangieide (e.g. benzisothiazolimone). In step 6 this mixture is dispersed until grind is 5-7 micrometers on the Hegman gauge.

£0041] In the mixing process, the composition is formed as a thixotropic gel or liquid. The thixotropic composition may be mixed or otherwise sheared to reduce its viscosity and increase its flow properties to values sufficient for application as a liquid to the rail head using conventional pump systems. For example, the viscosity achieved by mixing may be in the range of from about 5,000 to about 15,000 cP as measured using a Brookfield viscometer in accordance with ASTM D 29S3-02a. More preferably, the shear-mixed viscosity may be in the range of from about 8,000 to about 12,000 cP. The viscosity range may reflect the particular mode of application to the rail, and the foregoing range has been found satisfactory for pumping, spraying and other application techniques.

[0042] Upon termination of mixing and shear, the composition has a static thickness or cone penetration consistency in the range of from about 300 to about 400 tenths of a millimeter as measured using a standard cone test in an onworked condition in accordance with ASTM
217 -97. More preferably, the cone penetration may range from about 355 to about 375 tenths of a millimeter. The static thickness or cone penetration consistency may be varied to reflect weather conditions.

The foregoing thixotropic properties have been found to sufficient to allow ease of rail application and increased distribution or "carry down" along the rail road track as compared with prior art friction modifier formulations.

By the term "water-insoluble hydrocarbon" we mean hydrocarbons that are not typically miscible in water based solutions. The insoluble hydrocarbon has a solubility in water of less than or equal to 10 wt %, or even more specifically, less than or equal to 5 wt %, or even more specifically, less than or equal to 1 wt %, at ambient conditions of about 70°F and one atmosphere of pressure. Examples of such a water-insoluble hydrocarbon include isoparaffins such as SOTROL 220 (C13-C16 isoalkanes) and vegetable oils such as refined canola oil. Other potential water-insoluble hydrocarbons include bio-based triglycerides, fatty oils, polyalphaolefins such as DURASYN 162 and SYNFUID PA02, and synthetic esters such as di-octyl adipate and isopropyl oleate.

It has been found that the water-insoluble hydrocarbon gives the composition surprising advantages over water. Water based systems contain ammonia and this can cause equipment to rust. Also latex in water-based systems can cause problems with gears and other movable mechanical parts as the compositions dry out. Finally, the skin that can form over water based systems can cause flowing out or splashing when contacted by the wheels of the train and accordingly the friction modifier compound is then not carried down the track by the wheels of the train. Less water in the composition can help with all of these problems. We have found that replacing some of the water with a water-insoluble hydrocarbon is advantageous.

In the past as people have tried to lower the level of water in their friction control compositions they have at times attempted to replace some of the water with freeze depressants such as glycerine. However, this has the drawback that high amounts of water soluble polyalphaolefin freezing point depressants (e.g. glycerine) can have negative effects on the friction modifier composition since these type of freezing point depressants can cause rheology problems as the thickeners (e.g. clays) do not remain in the homogenous system.
We have found that replacing some of the water with a water-insoluble hydrocarbon (e.g. SOTROL 220) can improve the overall composition by eliminating or minimizing the negative effects set out above. The addition of a water-insoluble hydrocarbon into the system is counterintuitive since by definition a water-insoluble hydrocarbon should not mix well with a composition containing water. However, surprisingly we found that certain water-insoluble hydrocarbons (e.g. SOTROL 220 and refined canola oil) mix well with the compositions described herein because the water-insoluble hydrocarbon goes into a matrix with the clay in the composition.

By the term "friction modifier" we mean a solid powder which changes the coefficient of friction, in this case, from negative to positive. Examples of such liquid or solid friction modifiers include talc and barium sulfate. The friction modifiers can be chosen from the following list of friction modifiers, but are not limited to these friction modifiers, WHITING (calcium carbonate), BLANC FIXE (calcium sulphate), mineral fibre, whilasionite powder, powdered cashew nut shells, calcium carbonate, aluminum oxide, amorphous silica, silica oxide, magnesium oxide, magnesium carbonate, lead oxide and coal coke.

By the term "liquid or solid lubricant" we mean a liquid or solid material that reduces friction between two sliding metal surfaces. Examples of two preferred lubricants are superfine molybdenum disulfide and carbon black (in some embodiment used in combination with one another). A non-exclusive list of other potential liquid or solid lubricants includes graphite and zinc stearate. However this invention is not limited to these lubricants only.

By the term "wetting agent" we mean a surfactant which assists the liquid to wet out the solids in the formula. One preferred example of such a wetting agent includes Triton X-100. A non-exclusive list of other potential wetting agents include, UNIVAR propylene carbonate technical", CO630, TEXANOL, and TEXAPON P.

By the term rheology agent we mean a clay ore other substance that expands in water to produce a thixotropic mix. An example of one such rheology agents comprises clay such as VAN GEL B. Another example of a rheology agent usable with clay is hydrated lime. A non-exclusive list of other potential rheology agents includes methyl ethyl hydroxy cellulose and ethyl hydroxy cellulose,
By the term freezing point depressant we typically mean an alcohol which when mixed with water, lowers the freezing point of water. One preferred example of such freezing point depressant includes SUPER KPO glycerine. A non-exclusive list of other potential freezing point depressants includes ethyl alcohol, methyl alcohol isopropanol and butanol.

There are multiple ways of making the friction modifier composition disclosed herein. In one embodiment the composition typically comprises:

(a) from 24 to 25 w/w% water
(b) from 4 to 7 w/w% rheology additive
(c) from 14 to 18 w/w% liquid or solid friction modifier
(d) from 22 to 32 w/w% freezing point depressant
(e) from 14 to 18 w/w% liquid or solid friction modifier
(f) from 2 to 3 w/w% liquid or solid lubricant
(g) from 1 to 3 w/w% surfactant or wetting agent
(h) from 0.2 to 0.5 w/w% corrosion inhibitor
(i) from 0.1 to 0.2 w/w% bioeide/rungicide agent

The following Examples 1 to 6 illustrate other preferred friction control compositions in accordance with the invention comprising:
In one embodiment of the invention (see Example 5), it has been found that adding a water insoluble hydrocarbon (as a non-exclusive example paraffinic or isoparaffinic solvent SOTROL 220) helps depress the freezing point and also helps stabilize or even improve the rheology of the formulation. This is especially true when the water insoluble hydrocarbon is compared with other freezing point depressants such as glycerin. The addition of the water insoluble hydrocarbon is counterintuitive because one would have guessed that it would not mix well with the water based formulation of this invention and would in all likelihood separate. However, we believe that the clay has receptor sites that allow the water insoluble hydrocarbon to bind onto the clay and keep the final product homogenous.

In yet another embodiment of the invention (see Examples 1-4) it has been found that adding a vegetable oil (as a non-exclusive example, refined canola oil) has some of the advantages of paraffins or isoparaffins such as SOTROL 220 but also include environmental advantages such as having a higher flash point and being biodegradable.

In yet another embodiment of the invention (see Examples 2-3) it has been found that adding carbon black has some advantages. Carbon black was originally added as a solid lubricant to the formulation in order to lower the costs by using a less expensive lubricant than molybdenum disulfide. However, in addition to lowering the cost of the formulation, it was surprisingly found that carbon black also helps with the stability of the composition (i.e. less separation) and can give a surprising increase in viscosity which in some embodiments is also very helpful. The carbon black can be added in ranges from 0.5 to 5%.

In yet another embodiment of the invention (see Example 1) we use an aprotic solvent exhibiting limited water solubility (e.g. propylene carbonate, solubility in water is 17.5% at 25° C.) rather than the high amounts of glycerine used in other examples set forth infra. The propylene carbonate causes the thickener (e.g. the clay) to go into a matrix such that it has better solubility and can result in a higher friction product than can be achieved with glycerine. The propylene carbonate also helps as a free freeze point depressant and improves product efficacy at lower temperatures.

In yet another embodiment of the invention (see Example 7) it may be desirable in certain cold weather environments (e.g. at or below about negative 40 degrees centigrade) for the formulation to contain much higher amounts of freezing point depressants such as
glycerine or propylene glycol. In these extremely cold weather environments it may be desirable to replace some (or even all) of the water insoluble hydrocarbons with a freezing point depressant. In at least one embodiment for the cold weather composition the ratio of the glycerine to water shall be at least 63% glycerine to 37% water. For other freezing depressants the ratio of the depressant to water may differ as a function of the freezing point curve. Based upon the freezing point behaviors of these fluid blends they are commonly called eutectic mixtures. Propylene glycol: water mixtures maintain freezing points at or lower than -40° at any ratio of 55% or more propylene glycol. The ratio of glycerine: water shall be within the range of 63 – 70% glycerine for cold temperature flow down to -40° F.

[0060] Example 1 (the amounts below are weight percent).

Add UNTV All propylene carbonate technical to tap water is the vat and stir at slow speeds.

Add first portion of VAN GEL B (clay) slowly while Stirling for 30 min. under cowls mixer at high speeds.

Add SUPER KPO glycerine slowly and mix at high speed for 10 min.

Add a second portion of VAN GEL S (thy) slowly while stirring for 30 min. under cowls mixer at high speeds.

Add slowly refined canola oil AGRIPURE 60 and mix for 15 minutes at high speed.

Add in order while stirring, a tellurium disulfide THOMPSON CREEK SUPERFINE and mix at high speed for 15 minutes.

Add barium sulfate BARIMITE XF
and mix at high speeds for 15 minutes.

Add tide, magnesium silicate NICRON 604 and mix at high speeds for 15 minutes.

Add COUNTER RUST LT-26 and mix at medium speeds for 15 minutes.

Add anti-fungus, sitrobo®hölue BIOBAN P 1487 and mix at medium speeds for 15 minutes.

Disperse until grind is 5–7 on the Hegman gauge.

Example 2:

Add tap water to vessel
stir at low speeds.

Add first portion of VAN GEL B (day) slowly white stirring under cowls mixer at high speeds.

Add wetting agent, TRITON X-100 to mixture

Add SUPER KPO glycerine slowly
and mix at high speed for 10 min.

Add second portion of VAN GEL B (clay) slowly while stirring for 30 min. under cowls mixer at high speeds.

Add slowly refitted caaofa oil AGRIPURE 60 And mix for 15 minutes at high speed.

Add in order while stirring, molybdenum disulfide THOMPSON CREEK SUPERFINE
And mix at high speed for 15 minutes.

Add barytes, barium sulfate BARiMiTE XF and mix at high speeds for 15 minutes.

Add talc, magnesium silicate NICRON 604 and mix at high speeds for 15 minutes.

Add carbon black sad mix at high speeds for 15 minutes.

Add COUNTER RUST LT-267 and mix at medium speeds for 15 minutes.

Add anti-fungus, K 80078 and mix at medium speeds for 15 minutes.

Disperse until grind is 4-8 on the Hegman gauge.

[0062] Example 3:

Add tap water to vessel 25.0
stir at low speeds.

Add first portion of VAN GEL B (clay) 4.5
slowly while stilting under cowls mixer at high speeds.

Add wetting agent TRITON X-100 to mixture. 2.5

Add SUPER KPO glycerine slowly 27.9
and mix at high speed for 10 min.

Add second portion of VAN GEL B (day) 2.5
slowly while stirring for 30 min. under cowls mixer at high speeds.

Add slowly
refined canola oil AGRIPURE 60 14.6
and mix for 15 minutes at high speed.

Add in order while stirring,
molybdenum disulfide THOMSON CREEK SUPERFINE 2.0
And mix at high speed for 15 minutes.

Add barites, barium sulfate BARIMATE XF 3.0
and mix at high speeds for 15 minutes.

Add micro. magnesium silicate MICRON 604 15.0
and mix at high speeds for 15 minutes.

Add carbon black 2.5
and mix at high speeds for 15 minutes.

Add hydrated lime, Ca(OH)₂ 0.1
And mix at high speeds for 15 minutes.

Add COUNTER RUST LT-26? 0.2
and mix at medium speeds for 15 minutes.

Add anti-fungus, PRGXEL GXL Antimicrobial 0.2
and mix at medium speeds for 15 minutes.

Disperse till grind is 4-8 on the Hegman gauge.

[0063] Example 4

Add tap water to vessel 24.0 24.0
and stir at low speeds.
Add first portion of VAN GEL B (day) 4.5 slowly while stirring under cowls mixer at high speeds.

Add wetting agent, TRITON X-100 to mixture. 2.0

Add SUPER KPO glycerine slowly 28.5 and mix at high speed for .10 min.

Add second portion of VAN GEL B (clay) 2.5 slowly while stirring for 30 min. under cowls raker at high speeds.

Add slowly
refmed canola oil AGRIPURE 60 17.8 and mix for 15 minutes at high speed.

Add in order while stirring,
molybdenum disulfide THOMPSON CREEK SUPERFINE 3.0 and mix at high speed for 15 minutes.

Add barites, barium sulfate BARIMITE XF 3.0 and mix at high speeds for 15 minutes.

Add talc, magnesium silicate NiCRON 604 11.3 and mix at high speeds for 15 minutes

Add COUNTER RUST LT-267 0.2 and mix at medium speeds for 15 minutes.

Add anti-fungus, PRGXEL GXL antimicrobial 0.2 and mix at medium speeds for 15 minutes

Disperse until grind is 4-8 on the Hegman gauge.
Example 5

Add TRITON X-100 and tap water into the vat and stir.

Add slowly VAN GEL 8 (clay) while stirring under cowls mixer, it will become very thick.

Add slowly while stirring isoparaffinic solvent SOTR.OL 220
When well dispersed, add glycerol slowly while stirring, check that product is still thixotropic.

Add iotxfcr while stirs g,
molyhdcnm disulfide superfine grade, barjum sulfate, tale, hydrated lime, COUNTER RUST LT-267, K 78 biocide (1:10 dilution)

Disperse until grind is 5-7 on the Hegrao, gauge

Example 6

Add tap water into the vat and stir.

Add VAN GEL B (clay) and GARAMTIE clay
Add TRITON X-100 2.0
Add slowly while stirring under cowls mixer, it will become very thick.

Add glycerol slowly while stirring, 31.5

Add in order while stirring,
molybdenum disulfide superfine grade, 2.0
barum sulfate, 3.0
talc 15.0
carbon black, 2.5
COUNTER RUST LT-267 0.2
PROXEL antimicrobial 0.2
Add in canola oil 14.6

Disperse until homogeneous.

In the foregoing examples, the above mentioned thixotropic properties are achieved. That is, the shear-mixed composition has a thickness or viscosity in the range of from about 8,000 to about 12,000 cP to facilitate application to the rail using conventional techniques. The static composition has a thickness or penetration consistency of from about 355 to about 375 to maintain the composition on the rail for subsequent carry down by train wheel passage.

[0066] Example 7

Add TRITON X-100 2.0
and tap water 25.0
into the vat and stir.

Arid VAN GEL B (clay) 2.0
GARAM[T clay] 2.0
When well dispersed, add glycerol slowly while stirring, 46.0
check that product is still thixotropic,

Add in order while stirring,
molybdenum disulfide superfine grade 2.0
barium sulfate 3.0
talc 15.0
hydrated lime 0.1
carbon black 2.5
COUNTER RUST tX-26? 0.2
K 78 biocide {1:10 dilution} 0.2

[0067] Disperse until grind is 5-7 on the Hegman gauge.

[0068] Performance data using the friction control compositions disclosed herein have confirmed the compositions to be surprisingly more effective than prior art top of rail friction composition modifiers. In the following tests, "fresh" trains having 286,000 pound loaded coal cars passed over an applicator arranged to apply the composition of Example 2. The applicator was mounted to the field side of the track and pumped the shear-mixed liquid composition onto the rail head in a conventional manner and amount ahead of the die test curve in the track. The composition returns to its static condition on the rail prior to engagement with the train wheels. The trains are "fresh" in that the wheels were not previously treated, but rattier, the wheels were dry and had contaminants typically encountered in transportation. The product is picked-up by the wheels and carried in the wheel-rail "contact patch" down the track to the curve to provide the benefits of the invention.

[0069] Referring to Fig. 2, reduction of the lateral ibices and achievement of a positive coefficient of friction or friction is shown for use of the friction control composition in accordance with above Example 2. To that end, a dry wheel-rail system is compared with a wheel-rail system having the friction control of the present invention applied in liquid form to the top of the rail as described above.

[0070] Fig. 2 shows the distribution of the L/V ratio on the high rail of a 5.7° curve for trail axles of the dry wheel-rail system and the wheel-rail system comprising the friction control composition of Example 2. The friction control composition reduces the lateral forces as
indicated by the lower L/V ratio. That is, the composition of Example 2 changes the friction from negative to positive, limits the creep of the wheel on the rail head and reduces the lateral engagement force and/or contact by the wheel flange with the gauge side face of the rail. Accordingly, the L/V ratio is reduced.

(0071) It has also been found that the composition of Example 2 tends to limit the variation of the L/V ratio so as to result in a closer grouping of data points. This is also believed to be related to the smoothing of the train travel and increased train stability.

(0072) Referring to Fig. 3, the L/V ratio is shown for the lead axles of the cars of Fig. 2 for the dry wheel-rail system compared with the wheel-rail system having the friction control composition of Example 2. As stability increases, the L/V ratio for the treated wheel-rail system decreases to values less than those of the dry wheel-rail system.

(0073) Referring to Figs. 4 and 5, histograms show the lateral force distribution for the trail and lead axles of the train cars of Figs. 2 and 3. As shown, the lateral forces are reduced.

(0074) Referring to Figs. 6 and 7, the improved "carry down" of the compositions of the invention is shown. As noted above, carry down is how far the friction control composition is carried along the track in an effective amount from the application location. The former down the track the composition is carried the better for the rail-road customer since the friction control composition works over a longer distance (e.g. stick-slip and creep are reduced together with the achievement of the other benefits described above). This can save rail road customers significant money by requiring fewer applicators and also less friction control product.

(0075) To that end, the L/V ratios for the wheel-rail systems of Figs. 2-5 were measured at a distance of 6.9 miles from the point of application and shown in Figs. 6 and 7. It has been found in third party tests that the friction control compositions disclosed herein have carry down from two to three times further than prior art products. This dramatic increase is surprising to both the third party testers and potential customers.

(0076) The present invention has been described with regard to preferred embodiments. However, it will be obvious to persons skilled in the art that a number of variations and
modifications can be made without departing from the scope of the invention as described herein. In the specification the word "comprising" is used as an open-ended term, substantially equivalent to the phrase "including but not limited to", and the word "comprises" has a corresponding meaning. Citation of references is not an admission that such references are prior art to the present invention.
What is Claimed is:

1. A friction control composition for use on top of rail applications comprising:
   (a) from about 4 to about 40 w/w% water;
   (b) from about 2 to about 20 w/w% rheology additive;
   (c) from about 10 to about 40 w/w% water insoluble hydrocarbon;
   (d) from about 10 to about 40 w/w% water soluble polyalcohol freezing point depressant;
   (e) from about 1 to about 7 w/w% liquid or solid friction modifier; and
   (f) from about 1 to about 20 w/w% liquid or solid lubricant; and

   wherein said water insoluble hydrocarbon is selected from the group consisting of isoparaffins, vegetable oils, bio-based triglycerides, fatty oils, polyalpha olefins and synthetic esters.

2. The friction control composition of claim 1, comprising:
   (a) from about 15 to about 29 w/w% water;
   (b) from about 4 to about 13 w/w% rheology additive;
   (c) from about 11 to about 28 w/w% water insoluble hydrocarbon;
   (d) from about 22 to about 40 w/w% water soluble polyalcohol freezing point depressant;
   (e) from about 9 to about 24 w/w% liquid or solid friction modifier; and
   (f) from about 1 to about 6 w/w% liquid or solid lubricant.

3. The friction control composition of claim 1, comprising:
   (a) from about 24 to about 25 w/w% water;
   (b) from about 4 to about 7 w/w% rheology additive;
   (c) from about 14 to about 18 w/w% water insoluble hydrocarbon;
   (d) from about 22 to about 32 w/w% water soluble polyalcohol freezing point depressant;
   (e) from about 14 to about 18 w/w% liquid or solid friction modifier; and
   (f) from about 2 to about 3 w/w% liquid or solid lubricant.
4. The friction control composition of claim 1, wherein said water insoluble hydrocarbon is selected from the group consisting of isoparaffins and fatty oils.

5. The friction control composition of claim 4, wherein said water insoluble hydrocarbon is selected from the group consisting of C8 to C16 isomarkanes and canola oil.

6. The friction control composition of claim 1, wherein said composition has a viscosity achieved by shear mixing in the range of from about 5,000 to about 15,000, and upon termination of shear mixing and return to static conditions, the composition has a static viscosity or cone penetration thickness in the range of from about 300 to about 400 tenths of a millimeter.

7. The friction control composition of claim 1, further comprising one or more of the following:
   (g) from 1 to 7 w/w% aprotic solvent exhibiting limited water solubility;
   (h) from 0.1 to 4 w/w % corrosion inhibitor; and
   (i) from 0.05 to 2 w/w% biocide/fungicide agent.

8. The friction control composition of claim 1, further comprising: an aprotic solvent exhibiting limited water solubility, a corrosion inhibitor, a biocide/fungicide agent, carbon black, lime or a combination thereof.

9. A friction control composition for use on top of rail applications comprising:
   (a) from about 15 to about 29 w/w% water;
   (b) from about 4 to about 13 w/w% rheology additive;
   (c) from about 1 to about 28 w/w% water insoluble hydrocarbon;
   (d) from about 22 to about 40 w/w% water soluble polyalcohol freezing point depressant;
   (e) from about 9 to about 24 w/w% liquid or solid friction modifier; and
   (f) from about 1 to about 6 w/w% liquid or solid lubricant; and
wherein said water insoluble hydrocarbon is selected from the group consisting of isoparaffins, canola oil, polyalpha olefins and synthetic esters.
10. The friction control composition of claim 9, further comprising: an aprotic solvent exhibiting limited water solubility, a corrosion inhibitor, a btoctde/ftngticide agent, carbon black, lime or a combination thereof.

11. A method of modifying the friction of a rail road track from negative to positive comprising:

mixing the friction control composition of claim 1 by applying shear to thin or reduce the thickness or viscosity of the composition;

applying said thinned composition to the top of a rail; and

causing said thinned composition on the top of the rail to settle to a static condition of increased viscosity sufficient to maintain the composition in position on the top of the rail as an thinned composition for engagement by a train wheel with shear thinning and distribution of the composition along the rail to change the friction from negative to positive.

12. The friction control composition of claim 11, wherein said thinned composition has a viscosity or cone penetration achieved by shear mixing in the range of from about 5,000 to about 5,000, and upon termination of shear mixing and return to static conditions, the composition has a static viscosity or cone penetration thickness in the range of from about 300 to about 490 tenths of a millimeter.

13. The method of claim 11, wherein said step of applying said thinned composition as a liquid to the top of the rail includes pumping said thinned composition onto the top of the rail.

14. A friction control composition for use on top of rail applications consisting essentially of:

(a) from about 4 to about 40 w/w % water;
(b) from about 2 to about 20 w/w % rheology additive;
(c) from about 10 to about 40 w/w % water insoluble hydrocarbon;
(d) from about 10 to about 40 w/w % water soluble polyalcohol freezing point depressant;
(e) from about 1 to about 7 w/w % liquid or solid friction modifier; and
(f) from about 1 to about 20 w/w% liquid or solid lubricant; and
wherein said water insoluble hydrocarbon is selected from the group consisting of
isoparaffins, vegetable oils, bio-based triglycerides, fatty oils, poiyaipa olefins and
synthetic esters.

15. The friction control composition of claim 14, wherein said composition is substantially
free of a skin forming retentivity agent.

16. The friction control composition of claim 15, wherein said composition is substantially
free of a skin forming retentivity agent selected from the group consisting of acrylic,
polyvinyl alcohol, polyvinyl chloride, oxisolae, epoxy, alkyd, aretha acrylic, modified alkyd,
acrylic latex, acrylic epoxy hybrids, polyurethane, styrene acrylate, aad stymie butadiene based
compounds.
**Fig. 1**

### Step 1
Charge water, water-insoluble hydrocarbon (e.g. Sotrol 220), wetting agent (e.g. Van Wet 9N9) to a container. Begin mixing.

### Step 2
Charge the rheology additive (e.g. Van Gel B) slowly into the container.

### Step 3
Dispense under high sheer to form a thixotropic gel.

### Step 4
Charge water soluble polyalcohol freezing point depressant (e.g. Glycerine) slowly while mixing.

### Step 5
Charge liquid or solid lubricant (e.g. superfine molybdenum); liquid or solid friction modifier (e.g. barium sulfate, talc); corrosion inhibitor (e.g. ammonium benzoate) and biocide/fungicide agent (e.g. benzisothiazolinone) in the order above.

### Step 6
Dispense until grind is 5–7 on a Hegman gauge.
**Fig. 2**

Histogram of L/V ratio for trail axles of 286,000 pound cars.
High rail of 5.7° curve.

**Fig. 3**

Histogram of L/V ratio for lead axles of 286,000 pound cars.
High rail of 5.7° curve.
FIG. 6
HISTOGRAM OF HIGH RAIL L/V RATIO FOR LEAD AXLES OF 286,000 POUND CARS
DISTANCE FROM APPLICATOR TO INSTRUMENTED 5.7° CURVE – 6.9 MILES

FIG. 7
HISTOGRAM OF LOW RAIL L/V RATIO FOR LEAD AXLES OF 286,000 POUND CARS
DISTANCE FROM APPLICATOR TO INSTRUMENTED 5.7° CURVE – 6.9 MILES
INTERNATIONAL SEARCH REPORT

International application No.
PCT/US14/10188

A. CLASSIFICATION OF SUBJECT MATTER
IPC(8) - B00D 1/02; C10M 101/04, 113/10 (2014.01)
USPC - 508/1 10, 143, 449
According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED
Minimum documentation searched (classification system followed by classification symbols)
IPC(8): B00D 1 02; C10M 101/00, 101/04, 105/00, 105/02, 105/04, 113/00, 113/02, 113/10, 115/00, 115/02, 125/00, 125/14, 125/30, 127/02, 129/95, 159/08; C10N 20/02, 20/06, 30/00, 30/02, 30/08 (2014.01); USPC: 508/1 10, 113, 126, 136, 143, 167, 169, 449

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of database and, where practicable, search terms used)

C. DOCUMENTS CONSIDERED TO BE RELEVANT

<table>
<thead>
<tr>
<th>Category</th>
<th>Citation of document, with indication, where appropriate, of the relevant passages</th>
<th>Relevant to claim No.</th>
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Further documents are listed in the continuation of Box C.

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