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(54) **OIL SOLUBLE POLYOXYBUTYLENE POLYMERS AS FRICTION MODIFIERS FOR LUBRICANTS**

**ÖLLÖSLICHE POLYOXYBUTYLENPOLYMERE ALS REIBUNGSMODIFIKATOREN FÜR SCHMIERMITTEL**

**POLYMÈRES DE POLYOXYBUTYLÈNE SOLUBLES DANS L'HUILE EN TANT QUE MODIFICATEURS DE FROTTEMENT POUR DES LUBRIFIANTS**

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(73) Proprietor: **Dow Global Technologies LLC**  
**Midland, MI 48674 (US)**

(72) Inventors:  
• **GREAVES, Martin R.**  
**CH-ZG 6340 Baar (CH)**  
• **ZAUGG-HOOZEMANS, Evelyn A.**  
**CH-8810 Horgen (CH)**

(74) Representative: **Houghton, Mark Phillip**  
**Patent Outsourcing Limited**  
**1 King Street**  
**Bakewell, Derbyshire DE45 1DZ (GB)**

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**EP-A1- 0 438 709**      **WO-A1-2013/062682**  
**WO-A2-2011/011656**

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**Description**Field

5 **[0001]** This invention relates generally to a lubricant composition. More specifically, the invention relates to a lubricant composition containing a base oil and a polyoxybutylene polymer as a friction-modifier additive, wherein the polyoxybutylene polymer and the base oil are soluble in one another.

Background

10 **[0002]** Increasing interest in developing lubricants that provide low friction and which are energy efficient in equipment for both the automotive and industrial lubricants industries is a macro trend across the lubricant industry today. One important way to reduce friction is to use friction-modifier additives in lubricant compositions. There has been considerable research carried out to explore the performance and mechanisms of action of friction-modifying additives across the lubricant industry. During this time, tribological tools for studying surfaces have greatly enhanced our understanding of how additives perform. Researchers have shown that friction can be reduced in boundary lubricating conditions by adsorbing or reacting additives on metal surfaces to form thin low-shear-strength films. Furthermore some of these additives can have a significant effect on friction in the mixed lubrication regime.

15 **[0003]** Two examples of friction-modifying additive chemistries are organic friction modifiers and organo-molybdenum compounds. The former are usually long-chain polar compounds based on carboxylic acid, amine, ester and alcohol groups. Examples include glycerol mono-oleate, oleylamide, stearic acid and trimethylolpropane esters. These tend to function through their polar heads absorbing on to surfaces with the lipophilic tail aligned perpendicular to the surface. There are some practical challenge in using these materials in lubricant compositions. For example esters are prone to hydrolysis if there is ingress of water into the lubricant. Amine containing materials are known to cause elastomer incompatibility issues. Amides, such as oleylamide, are known to have a high degree of surfactancy character and can lead to emulsion formation. Carboxylic acids can react with metals to form carboxylate salts that are sometimes not desired.

20 **[0004]** When formulating lubricants, it is highly desirable that all additives, including friction-modifiers, be soluble in the composition. Such solubility is preferably maintained across a wide range of temperature and other conditions in order to enable shipping, storage, and/or prolonged use of these compositions.

25 **[0005]** International patent application WO 2011/011656 A2 discloses lubricant compositions that contains certain hydrocarbon oils and certain polyalkylene glycols. International patent application WO 2013/062682 A1 discloses lubricant compositions that comprise a Group I, II, III or IV hydrocarbon oils and a polyalkylene glycol prepared by reacting a C<sub>8</sub>-C<sub>20</sub> alcohol and a mixed butylene/propylene oxide feed using a double metal cyanide catalyst catalysed oxyalkylation process.

30 **[0006]** Lubricant additives that provide significant friction modification benefits without the disadvantages of current additives, such as hydrolytic instability, and that are also readily soluble in the lubricant base oil would be highly beneficial to the lubricant industry.

Statement of Invention

35 **[0007]** We have now found that polyoxybutylene polymers as described herein function as excellent friction modifier additives for lubricants. In particular, it has been found that selection of polyoxybutylene polymers having a number average molecular weight ranging from 800 to 1200 g/mol and prepared from a monol initiator, as herein described, significantly outperform similar materials that are otherwise of lower or higher molecular weight or that are prepared from a non-monomer initiator. Advantageously the polyoxybutylene polymers are also soluble in hydrocarbon base oils.

40 **[0008]** Moreover, the inventive materials outperform conventional ester based friction modifiers and provide the added benefit of having greater hydrolytic stability over the esters, thus making them more tolerant of water that may be present in lubricant compositions.

45 **[0009]** In one aspect, therefore, there is provided a lubricant composition comprising: a hydrocarbon base oil; and a friction modifier comprising a polyoxybutylene polymer, the polyoxybutylene polymer having been prepared by polymerizing butylene oxide with a monol initiator, and having a number average molecular weight ranging from 800 g/mol to 1200 g/mol; the polyoxybutylene polymer further characterized as being a propylene glycol n-butyl ether initiated butylene oxide homo-polymer having a kinematic viscosity at 40 °C of 60 mm<sup>2</sup>/s (60 cSt).

50 **[0010]** In another aspect, there is provided a method for reducing friction between lubricated surfaces, the method comprising: lubricating a surface with the lubricant composition as described herein, wherein friction is reduced relative to a composition free of the polyoxybutylene polymer.

55 **[0011]** In a further aspect, there is provided a method of lubricating a mechanical device, the method comprising using

the lubricant composition as described herein to lubricate the mechanical device.

#### Brief Description Of the Figures

5 **[0012]**

FIG. 1 shows friction profiles for various comparative and inventive polymers in a representative mineral hydrocarbon base oil (NEXBASE™ 3080).

10 FIG. 2 shows friction profiles for various comparative and inventive polymers in a representative polyalphaolefin hydrocarbon base oil (SPECTRASYN™ 8).

#### Detailed Description

15 **[0013]** Unless otherwise indicated, numeric ranges, for instance as in "from 2 to 10," are inclusive of the numbers defining the range (e.g., 2 and 10).

**[0014]** Unless otherwise indicated, ratios, percentages, parts, and the like are by weight. Unless otherwise indicated, the phrase "molecular weight" refers to the number average molecular weight as measured in a conventional manner.

20 **[0015]** "Propyleneoxy" or "PO" as used herein refers to  $-\text{CH}_2-\text{CH}(\text{CH}_3)-\text{O}-$  or  $-\text{CH}(\text{CH}_3)-\text{CH}_2-\text{O}-$ , and "butyleneoxy" or "BO" refers to  $-\text{CH}_2-\text{CH}(\text{CH}_2\text{CH}_3)-\text{O}-$  or  $-\text{CH}(\text{CH}_2\text{CH}_3)-\text{CH}_2-\text{O}-$ . "Alkyl" encompasses straight and branched chain aliphatic groups having the indicated number of carbon atoms.

**[0016]** The invention provides lubricant compositions comprising a hydrocarbon base oil and a polyoxybutylene polymer as a friction modifier and methods for its use. Advantageously, the hydrocarbon base oil and the polyoxybutylene polymer are soluble in each other.

25 **[0017]** Hydrocarbon base oils useful in the composition of the invention include the hydrocarbon base oils designated by the American Petroleum Institute as falling into Group I, II, III or IV. Of these, the Group I, II, and III oils are natural mineral oils. Group I oils are composed of fractionally distilled petroleum which is further refined with solvent extraction processes to improve properties such as oxidation resistance and to remove wax. Group II oils are composed of fractionally distilled petroleum that has been hydrocracked to further refine and purify it. Group III oils have similar characteristics to Group II oils, with Groups II and III both being highly hydro-processed oils which have undergone various steps to improve their physical properties. Group III oils have higher viscosity indexes than Group II oils, and are prepared by either further hydrocracking of Group II oils, or by hydrocracking of hydroisomerized slack wax, which is a byproduct of the dewaxing process used for many of the oils in general. Group IV oils are synthetic hydrocarbon oils, which are also referred to as polyalphaolefins (PAOs). Mixtures of the foregoing oils may be used. Lubricant compositions of the invention preferably contains at least 90 percent of the hydrocarbon base oil, alternatively at least 95 percent, by weight based on the total weight of the hydrocarbon base oil and the polyoxybutylene polymer. In some embodiments, the lubricant compositions contains up to 99 weight percent, alternatively up to 98 weight percent of the hydrocarbon base oil based on the total weight of the hydrocarbon base oil and the polyoxybutylene polymer.

35 **[0018]** The polyoxybutylene polymer useful herein (also referred to as a BO homopolymer) may be prepared by polymerizing butylene oxide with a monol initiator. Such polymerization processes are known to those skilled in the art (see for instance United States Patent publication number 2011/0098492) and suitable polymers are commercially available. In a typical polymerization procedure, the initiator is alkoxyated with butylene oxide in the presence of acidic or alkaline catalysts, or by using metal cyanide catalysts. Alkaline polymerization catalysts may include, for instance, hydroxides or alcoholates of sodium or potassium, including NaOH, KOH, sodium methoxide, potassium methoxide, sodium ethoxide and potassium ethoxide. Base catalysts are typically used in a concentration of from 0.05 percent to about 5 percent by weight, preferably about 0.1 percent to about 1 percent by weight based on starting material.

45 **[0019]** The addition of butylene oxide may, for instance, be carried out in an autoclave under pressures from about 10 psig to about 200 psig, preferably from about 60 to about 100 psig. The temperature of alkoxylation may range from about 30 °C to about 200 °C, preferably from about 100 °C to about 160 °C. After completion of oxide feeds, the product is typically allowed to react until the residual oxide is reduced to a desired level, for instance less than about 10 ppm. After cooling the reactor to an appropriate temperature ranging for instance from about 20 °C to 130 °C, the residual catalyst may be left unneutralized, or neutralized with organic acids, such as acetic, propionic, or citric acid. Alternatively, the product may be neutralized with inorganic acids, such as phosphoric acid or carbon dioxide. Residual catalyst may also be removed using, for example, ion exchange or an adsorption media, such as diatomaceous earth.

50 **[0020]** The monol initiator for use in the invention is propylene glycol n-butyl ether (available from The Dow Chemical Company as DOWANOL™ PnB).

55 **[0021]** Sufficient butylene oxide is used in the polymerization reaction with the initiator to provide a polyoxybutylene polymer having a number average molecular weight ranging from 800 g/mol to 1200 g/mol, alternatively 900 g/mol to 1100 g/mol, alternatively 950 g/mol to 1050 g/mol, or alternatively about 1000 g/mol.

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**[0022]** In some embodiments, the polyoxybutylene polymer is included in the lubricant compositions of the invention at a concentration of up to 10 percent by weight, alternatively up to 5 percent by weight, based on the total weight of the hydrocarbon base oil and the polyoxybutylene polymer. In some embodiments, the polyoxybutylene polymer is included in the lubricant compositions at a concentration of at least 1 percent by weight, alternatively at least 2 percent by weight, based on the total weight of the hydrocarbon base oil and the polyoxybutylene polymer. In some embodiments, the lubricant composition comprises from 5 to 10 weight percent of the polyoxybutylene polymer based on the total weight of the hydrocarbon base oil and the polyoxybutylene polymer.

**[0023]** Polyoxybutylene polymers as described herein function as highly effective friction modifier additives for lubricant compositions. Thus, the polyoxybutylene polymers reduce friction between lubricated surfaces relative to a composition free of the polyoxybutylene polymer. In some embodiments, the polyoxybutylene polymers reduce friction between lubricated surfaces by at least 10 percent, alternatively by at least 20 percent, at speeds of 10 and 20 mm/sec relative to a composition free of the polyoxybutylene polymer as measured by a Mini-Traction Machine in which a steel ball (diameter of 19mm) rotates on a steel disc (diameter of 45mm) at a slide-roll-ratio of 50% and a contact load of 50N and temperature of 80 °C.

**[0024]** Surprisingly, the polyoxybutylene polymers of the invention are significantly more effective friction modifiers than other materials with similar chemical structures and/or molecular weights. For instance, as demonstrated by the examples below, polyoxybutylene polymers prepared from a monol initiator and having a number average molecular weight ranging from 800 to 1200 g/mol, as herein described, significantly outperform polyoxybutylene polymers that are also prepared from a monol initiator but are otherwise of lower or higher molecular weight. In addition, the polyoxybutylene polymers of the invention outperform polyoxybutylene polymers that have very similar molecular weight but that were not prepared from a monol initiator.

**[0025]** Further advantageously, the polyoxybutylene polymers are soluble in hydrocarbon base oils. Moreover, they outperform conventional ester based friction modifiers and also provide the added benefit of having greater hydrolytic stability over the esters, thus making them more stable in the presence of water.

**[0026]** Lubricant compositions of the invention may contain other additives including, for instance, antioxidants, corrosion inhibitors, antiwear additives, foam control agents, yellow metal passivators, dispersants, detergents, extreme pressure additives, additional friction reducing agents, and/or dyes.

**[0027]** The compositions of the invention are useful as lubricants for a variety of mechanical devices including, for example, internal combustion engines such as automotive engines, gear boxes, hydraulic pumps, compressors and transmissions.

**[0028]** Some embodiments of the invention will now be described in detail in the following Examples.

### EXAMPLES

**[0029]** The materials in the following Table 1 are used in the example compositions.

Table 1

Material	Chemistry
PO/BO-550	Dodecanol initiated random copolymer (PO/BO, 50/50 by wt) with a typical kinematic viscosity at 40°C = 18 mm <sup>2</sup> /s (cSt). Its average molecular weight is 550 g/mole.
PO/BO-760	Dodecanol initiated random copolymer (PO/BO, 50/50 by wt) with a typical kinematic viscosity at 40°C = 32 mm <sup>2</sup> /s (cSt). Its average molecular weight is 760 g/mole.
PO/BO-1000	Dodecanol initiated random copolymer (PO/BO, 50/50 by wt) with a typical kinematic viscosity at 40 °C = 46 mm <sup>2</sup> /s (cSt). Its average molecular weight is 1000 g/mole.
PO/BO-1300	Dodecanol initiated random copolymer (PO/BO, 50/50 by wt) with a typical kinematic viscosity at 40 °C = 68 mm <sup>2</sup> /s (cSt). Its average molecular weight is 1300 g/mole.
Monol-BO-600	Propylene glycol n-butyl ether initiated BO homo-polymer with a typical kinematic viscosity at 40 °C of 25mm <sup>2</sup> /s (cSt). Its average molecular weight is 600 g/mole.

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(continued)

<b>Material</b>	<b>Chemistry</b>
Monol-BO-1000	Propylene glycol n-butyl ether initiated BO homo-polymer with a typical kinematic viscosity at 40 °C of 60mm <sup>2</sup> /s (cSt). Its average molecular weight is 1000 g/mole.
Monol-BO-2000	Propylene glycol n-butyl ether initiated BO homo-polymer with a typical kinematic viscosity at 40°C of 185mm <sup>2</sup> /s (cSt). Its average molecular weight is 2000 g/mole.
Diol-BO-1000	Monopropylene glycol (diol) initiated BO homopolymer with a typical kinematic viscosity at 40°C = 80 mm <sup>2</sup> /s (cSt). Its average molecular weight is 1000 g/mole.
SPECTRASYN™ 8 (available from Exxon Mobil Chemical)	A polyalphaolefin with a typical kinematic viscosity at 100°C = 8 mm <sup>2</sup> /s (cSt).
NEXBASE® 3080 (from Neste)	An API Group III mineral oil with a typical kinematic viscosity at 100°C = 8 mm <sup>2</sup> /s (cSt).
SYNATIVE™ TMTC (from BASF)	Trimethylol propane (TMP) ester of C8/C10 fatty acids with a typical kinematic viscosity at 40oC (KV40) = 19cst, KV100 = 4.3cst and Viscosity index (VI) = 142 (from BASF literature)
SYNATIVE™ ES 2960 (from BASF)	Dibasic acid ester with KV40 = 17.5cst, KV100 = 4.3cst and VI = 162 (from BASF literature)
SYNATIVE™ ES DITA (from BASF)	Dibasic acid ester with KV40 = 27.5cst, KV100 = 5.2cst and VI = 136 (from BASF literature). The ester is chemically known as Di-(triiso-decyl) adipate

Method of measuring traction (friction coefficient)

**[0030]** Friction coefficients are measured using a Mini-Traction Machine (from PCS Instruments) in which a steel ball is rotated on a steel disc. The disc used is steel (AISI 52100), diameter of 45 mm and hardness 750HV with a Ra <0.02 micrometers. The ball is steel (AISI 52100), diameter of 19 mm and hardness 750HV with a Ra <0.02 micrometers. Traction coefficients are measured at 80 °C at a slide-roll ratio of 50 % and speed 0-2500 mm/s and at a contact load of 37N. Traction values are reported at 5, 10 and 20 mm/sec.

**[0031]** The slide roll ratio, SRR, is the ratio of sliding speed to entrainment speed, i.e.

$$SRR = [U2-U1]/U$$

Where entrainment speed (U) is defined as the mean speed of the two surfaces as follows

$$U = \frac{1}{2}(U1 + U2)$$

Where U1 and U2 are the ball and disc speeds.

**[0032]** The compositions described in Tables 2 and 3 below are prepared by simply adding the ester or oil soluble polyalkylene glycol to the hydrocarbon base oil (either SPECTRASYN™ 8 or NEXBASE® 3080). The mixtures are stirred at ambient temperature until clear and homogeneous.

**[0033]** In the Tables 2 and 3, blends that represent the invention are labeled as "Inv. Ex." Comparative Examples are labeled as "C. Ex."

Table 2. Friction data in a representative mineral base oil

Example (Ex) or comparative Example (C.Ex)	Composition	5mm/sec	10mm/sec	20mm/sec
C. Ex	NEXBASE® 3080	0.111	0.094	0.074

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(continued)

Example (Ex) or comparative Example (C.Ex)	Composition	5mm/sec	10mm/sec	20mm/sec
C. Ex	NEXBASE® 3080 + PO/BO-550 (10%)	0.101	0.087	0.070
C. Ex	NEXBASE® 3080 + PO/BO-760 (10%)	0.113	0.091	0.074
C. Ex	NEXBASE® 3080 + PO/BO-1000 (10%)	0.119	0.096	0.072
C. Ex	NEXBASE® 3080 + PO/BO-1300 (10%)	0.097	0.080	0.065
C. Ex	NEXBASE® 3080 + Monol-BO-600 (10%)	0.104	0.086	0.074
Ex	NEXBASE® 3080 + Monol-BO-1000 (10%)	0.084	0.068	0.058
C. Ex	NEXBASE® 3080 + Monol-BO-2000 (10%)	0.098	0.087	0.069
C. Ex	NEXBASE® 3080 + Diol-BO-1000 (10%)	0.093	0.093	0.070
C. Ex	NEXBASE® 3080 + PO/BO-550 (5%)	0.099	0.088	0.074
C. Ex	NEXBASE® 3080 + PO/BO-1300 (5%)	0.108	0.091	0.080
Ex	NEXBASE® 3080 + Monol-BO-1000 (5%)	0.075	0.055	0.046
C. Ex	NEXBASE® 3080 + SYNATIVE™ DITA (10%)	0.097	0.083	0.071
C. Ex	NEXBASE® 3080 + SYNATIVE™ 2960 (10%)	0.101	0.086	0.075
C. Ex	NEXBASE® 3080 + SYNATIVE™ TMTc (10%)	0.086	0.082	0.072

Table 3. Friction data in a representative polyalphaolefin base oil

Example (Ex) or comparative Example (C.Ex)	Composition	5mm/sec	10mm/sec	20mm/sec
C.Ex	SPECTRASYN™ 8	0.113	0.097	0.075
C.Ex	SPECTRASYN™ 8 + PO/BO-550 (10%)	0.122	0.103	0.086
C.Ex	SPECTRASYN™ 8 + PO/BO-760 (10%)	0.109	0.087	0.069
C.Ex	SPECTRASYN™ 8 + PO/BO-1000 (10%)	0.116	0.085	0.070

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(continued)

Example (Ex) or comparative Example (C.Ex)	Composition	5mm/sec	10mm/sec	20mm/sec
5 C.Ex	SPECTRASYN™ 8 + PO/BO-1300 (10%)	0.100	0.090	0.071
10 C.Ex	SPECTRASYN™ 8 + Monol-BO-600 (10%)	0.122	0.091	0.078
Ex	SPECTRASYN™ 8 + Monol-BO-1000 (10%)	0.076	0.070	0.057
15 C.Ex	SPECTRASYN™ 8 + Monol-BO-2000 (10%)	0.101	0.087	0.069
C.Ex	SPECTRASYN™ 8 + Diol-BO-1000 (10%)	0.123	0.104	0.087
20 C.Ex	SPECTRASYN™ 8 + PO/BO-550 (5%)	0.085	0.074	0.057
C.Ex	SPECTRASYN™ 8 + PO/BO-1300 (5%)	0.102	0.080	0.066
25 Ex	SPECTRASYN™ 8 + Monol-BO-1000 (5%)	0.076	0.065	0.053
30 C.Ex	SPECTRASYN™ 8 + SYNATIVE™ DITA (10%)	0.096	0.088	0.068
C.Ex	SPECTRASYN™ 8 + SYNATIVE™ 2960 (10%)	0.104	0.084	0.068
35 C.Ex	SPECTRASYN™ 8 + SYNATIVE™ TMTc (10%)	0.087	0.078	0.060

[0034] In Tables 2 and 3 friction values are reported at three different speeds (5, 10 and 20 mm/sec). These speeds represent friction in the boundary region of the classical Stribeck curves. NEXBASE® 3080 and SPECTRASYN™ 8 are used as representative hydrocarbon base oils.

[0035] SYNATIVE™ 2960 and DITA (both dicarboxylic acid esters) and SYNATIVE™ TMTc (TMP polyol ester) have been used as friction reducers in hydrocarbon oils for many years. TMP polyol esters are considered to be more favorable but are more expensive. Friction reducers that can provide lower values than these benchmark products are desired. For example friction reducers that offer friction coefficients <0.078 at a speed of 10mm/sec are desired.

[0036] Tables 2 and 3 show data for three different chemical families of oil soluble polymers. Firstly, the propylene oxide/butylene oxide (PO/BO) derived series - these are alcohol (dodecanol) initiated PO/BO (50/50 w/w) random copolymers. Secondly, the monol-BO based series - these are propylene glycol n-butyl ether initiated (butanol + 1 PO) homo-polymers of BO. And thirdly the diol-BO based material (this is an example of a diol initiated BO homo-polymer).

[0037] The data in Tables 2 and 3 shows that the inventive Monol-BO-1000 is clearly differentiated versus the other materials in hydrocarbon base oils and exhibits lower friction values.

[0038] Surprisingly the inventive Monol-BO-1000, and the comparative PO/BO-1300 and Diol-BO-1000 are polymers that have similar viscosities and molecular weights but differ in the polymer architecture and their friction performance. Diol-BO-1000 shows no significant friction reducing behavior in NEXBASE® 3080 or SPECTRASYN™ 8 at a treat level of 10%. PO/BO-1300 shows a mild effect at a treat level of 10%. This polymer has a long chain linear tail (C12) and a mixed PO/BO tail. Inventive Monol-BO-1000 shows a significant friction reducing effect in NEXBASE® 3080 and SPECTRASYN™ 8. Furthermore Monol-BO-600 and Monol-BO-2000, which are lower and higher molecular weight polymers of this family did not exhibit the same friction reducing property.

[0039] FIGs. 1 and 2 illustrate friction profiles for various comparative and inventive polymers in the Mini-Traction

machine experiments. At speeds of <50mm/sec boundary friction can occur and friction reducer additives can be examined for their behavior. As is apparent from the FIGs, the inventive Monol-BO-1000 material exhibits a more favorable friction profile than the comparative materials.

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### Claims

1. A lubricant composition comprising:

10 a hydrocarbon base oil; and  
a friction modifier comprising a polyoxybutylene polymer, the polyoxybutylene polymer having been prepared by polymerizing butylene oxide with a monol initiator, and having a number average molecular weight ranging from 800 g/mol to 1200 g/mol;

15 the polyoxybutylene polymer further characterized as being a propylene glycol n-butyl ether initiated butylene oxide homo-polymer having a kinematic viscosity at 40 °C of 60 mm<sup>2</sup>/s (60 cSt).

2. The lubricant composition of claim 1 comprising up to 10 weight percent of the polyoxybutylene polymer based on the total weight of the hydrocarbon base oil and the polyoxybutylene polymer.

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3. The lubricant composition of claim 1 or 2 wherein the polyoxybutylene polymer has a number average molecular weight of 1000 grams per mole.

4. A method for reducing friction between lubricated surfaces, the method comprising:  
25 lubricating a surface with the lubricant composition of any one of claims 1-3, wherein friction is reduced relative to a composition free of the polyoxybutylene polymer, measured with a Mini-Traction Machine as described in the examples.

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5. A method of lubricating a mechanical device, the method comprising using the lubricant composition of any one of claims 1-3 to lubricate the mechanical device.

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### Patentansprüche

35 1. Eine Schmiermittelzusammensetzung, beinhaltend:

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ein Kohlenwasserstoff-Basisöl; und  
einen Reibungsmodifikator, beinhaltend ein Polyoxybutylenpolymer, wobei das Polyoxybutylenpolymer durch das Polymerisieren von Butylenoxid mit einem Monolinitiator hergestellt worden ist und ein zahlenmittleres Molekulargewicht im Bereich von 800 g/mol bis 1200 g/mol aufweist;

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wobei das Polyoxybutylenpolymer ferner **dadurch gekennzeichnet ist, dass** es ein mit Propylenglykol-n-butylether initiiertes Butylenoxidhomopolymer ist, das eine kinematische Viskosität bei 40 °C von 60 mm<sup>2</sup>/s (60 cSt) aufweist.

45 2. Schmiermittelzusammensetzung gemäß einem Anspruch 1, beinhaltend zu bis zu 10 Gewichtsprozent das Polyoxybutylenpolymer, bezogen auf das Gesamtgewicht des Kohlenwasserstoff-Basisöls und des Polyoxybutylenpolymers.

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3. Schmiermittelzusammensetzung gemäß Anspruch 1 oder 2, wobei das Polyoxybutylenpolymer ein zahlenmittleres Molekulargewicht von 1000 Gramm pro Mol aufweist.

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4. Ein Verfahren zum Reduzieren von Reibung zwischen geschmierten Oberflächen, wobei das Verfahren Folgendes beinhaltet:

Schmieren einer Oberfläche mit der Schmiermittelzusammensetzung gemäß einem der Ansprüche 1-3, wobei relativ zu einer Zusammensetzung, die das Polyoxybutylenpolymer nicht enthält, Reibung reduziert wird, gemessen mit einer Mini-Traction-Machine, wie in den Beispielen beschrieben.

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5. Ein Verfahren zum Schmieren einer mechanischen Vorrichtung, wobei das Verfahren das Verwenden der Schmier-

mittelzusammensetzung gemäß einem der Ansprüche 1-3 zum Schmierem der mechanischen Vorrichtung beinhaltet.

## Revendications

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1. Une composition lubrifiante comprenant :

une huile de base hydrocarbonée ; et

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un modificateur de frottement comprenant un polymère de polyoxybutylène, le polymère de polyoxybutylène ayant été préparé par polymérisation d'oxyde de butylène avec un initiateur monol, et ayant une masse moléculaire moyenne en nombre comprise dans l'intervalle allant de 800 g/mol à 1 200 g/mol ;

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le polymère de polyoxybutylène étant en sus caractérisé comme étant un homo-polymère d'oxyde de butylène initié par un éther n-butylique de propylène glycol ayant une viscosité cinématique à 40 °C de 60 mm<sup>2</sup>/s (60 cSt).

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2. La composition lubrifiante de n'importe quelle revendication 1 comprenant jusqu'à 10 pour cent en poids du polymère de polyoxybutylène rapporté au poids total de l'huile de base hydrocarbonée et du polymère de polyoxybutylène.

3. La composition lubrifiante de la revendication 1 ou de la revendication 2 dans laquelle le polymère de polyoxybutylène a une masse moléculaire moyenne en nombre de 1 000 grammes par mole.

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4. Un procédé destiné à la réduction de frottement entre des surfaces lubrifiées, le procédé comprenant : la lubrification d'une surface avec la composition lubrifiante de n'importe laquelle des revendications 1 à 3, dans lequel le frottement est réduit relativement à une composition exempte du polymère de polyoxybutylène, mesuré avec une machine MTM (Mini-Traction Machine) comme décrit dans les exemples.

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5. Un procédé de lubrification d'un dispositif mécanique, le procédé comprenant l'utilisation de la composition lubrifiante de n'importe laquelle des revendications 1 à 3 afin de lubrifier le dispositif mécanique.

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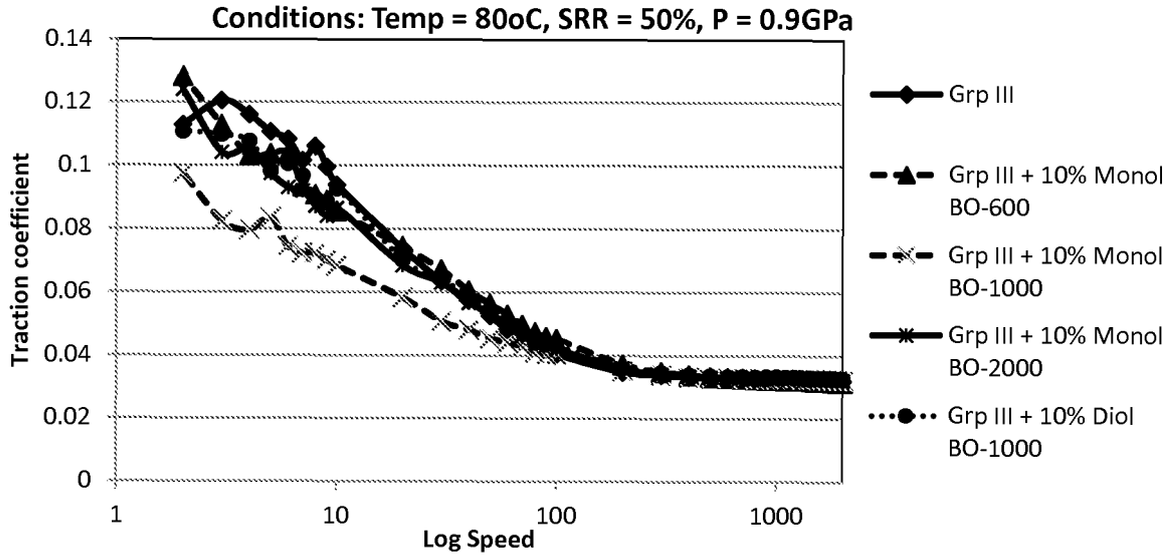


FIG 1

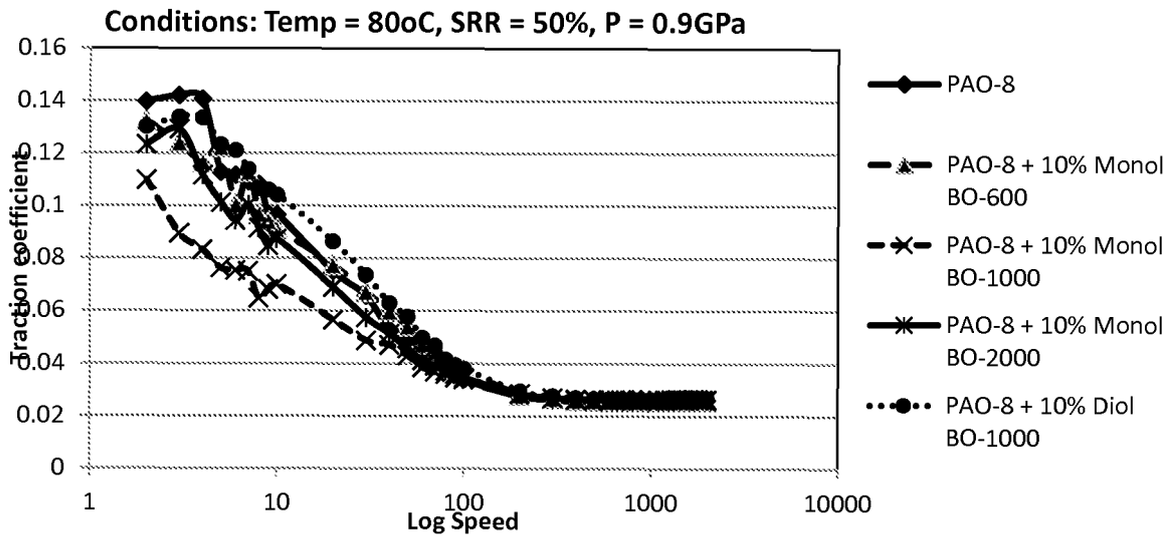


FIG 2

**REFERENCES CITED IN THE DESCRIPTION**

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**Patent documents cited in the description**

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