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(54) POLY(2,6-DIMETHYL-1,4-PHENYLENE-OXIDE) (56) References Cited DERIVATIVES WITH SULFONYL PENDANT GROUPS AND METHODS THEREFOR PUBLICATIONS

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- (51) Int. Cl.

- (52) U.S. Cl. CPC **C08G 65/44** (2013.01); **C08G 65/4056** (2013.01) ; C08G 65/485 (2013.01); C08G
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- Field of Classification Search CPC .. CO8G 65/44 ; CO8G 65/485 ; CO8G 2650/56 ; C08G 65/4056; G01R 33/46 USPC 524/208 See application file for complete search history. (58)

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Hay, Polypheneylene Oxide and Modified Polypheneylene Oxide Membranes, Chowdhury, G. et al., Eds., Springer, 1-26, 2001. (Year: 2001).*

* cited by examiner

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(57) ABSTRACT

A sulfonyl-substituted polyphenylene-ether polymer having improved dielectric properties and methods for making the same. The sulfonyl-substituted PPE include sulfone-contain-(*) Notice: Subject to any disclaimer, the term of this ing polyphenylene oxides ("PPO") polymers having repeat (*)

wherein each of R^1 and R^2 is H or an alkyl group containing
from 1 to 4 carbon atoms, R^3 is an alkylene group containing
from 1 or 2 carbon atoms, R^4 is selected from a group
consisting of an alkyl group conta atoms, an aryl group, and CF_3 , and X is a halogen atom. The subscript n is 0 or 1; the subscript m is 1 or 2, provided that when m is 2, R^2 is H. A degree of polymerization ranges from about 5 to about 1,000, and a ratio of units having
formula (1) to units having formula (2) of the sulfone-
containing PPO polymer ranges from about 10:90 to about $90:10.$

8 Claims, 6 Drawing Sheets

 $FT.1A$

FG . 10

FB , 2A

FG 2B

 \mathbf{FE} .3A

 $F.S.SC$

 $\mathbf{FE.4A}$

 $\texttt{FE.4B}$

FB.5A

FG , SB

FIG.6A

 $\mathbf{FE.6B}$

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POLY(2,6-DIMETHYL-1,4-PHENYLENE-OXIDE) Poly (phenylene ether)s or "PPEs" are a family of high DERIVATIVES WITH SULFONYL PENDANT performance thermoplastics with good high temperature GROUPS AND METHODS THEREFOR properties a

divisional application of prior filed, U.S. application Ser. That that is commercially produced on a large scale. PPO may
No 15/670.724, filed Aug 17, 2017, the disclosure of which be prepared by an oxidative coupling poly No. 15/679,724, filed Aug. 17, 2017, the disclosure of which be prepared by an oxidative coupling por polymerization of the prepared by an oxidative coupling polymerization of 2.6 -dimethylphenol as shown below: is expressly incorporated herein by reference, in its entirety.

RIGHTS OF THE GOVERNMENT

The invention described herein may be manufactured and used by or for the Government of the United States for all governmental purposes without the payment of any royalty.

FIELD OF THE INVENTION 25

The present invention is directed to sulfonyl-substituted poly(phenylene-ether)s having improved dielectric proper- $\frac{1}{20}$ wherein DP is a degree of polymerization.
ties and to methods for making the same.
PPEs have been considered for low-k materials for the

BACKGROUND OF THE INVENTION

rials have received increasing interest recently for various also have a low moisture absorption and are highly resistive
to acids and bases. On the other hand, for these thermoplas-
 potential applications including high energy-density-storage to acids and bases. On the other hand, for these thermoplas-
conceitor acto dialectrics and electrocritics materials. In capacitor, gate dielectrics, and electroactive materials. In
particular, high temperature polymers (those having a melt-
in a point greater than 150° C) with high dielectric constant ing point greater than 150° C.) with high dielectric constant cations, the ε_r value would need to be substantially higher
ord low dielectric loss are critical as embedded conscitors (i.e., greater than 6.0) and tan $\$ and low dielectric loss are critical as embedded capacitors, $(1.6.0)$
gradient metal need to be less than d would need to be less than 1% . which enable microelectronic-system integration to reduce
size without compromising performance or, better still, with Δs Attempts to modify PPEs to achieve the foregoing propthe possibility of enhanced performance in electronic sys-
terms. High-k materials suitable for some applications, such cross-linking, requires a multi-step process, or both, such tems. High-k materials suitable for some applications, such cross-linking, requires a multi-step process, or both, such as film capacitors for power-conditioning , power electronics that conventional methods cannot be used in hybrid electric vehicles, pulsed power, and gate dielectric polymers having a high degree of functionality. For
field-effect transistors, must possess processability, good ⁵⁰ example, Br—PPO reacting with sodium thiom field-effect transistors, must possess processability, good ³⁰ example, Br—PPO reacting with sodium thiomethoxide dielectric properties over a broad frequency range, and be (methylthiolate) via a nucleophilic substitutio dielectric properties over a broad frequency range, and be (methylthiolate) via a nucleophilic substitution reaction thermally stable. It is conventionally believed that no single according to convention methods affords me thermally stable. It is conventionally believed that no single according to convention methods affords methylsulfido-PPO material is able to satisfy all the above-mentioned prereq- ("CH₃S—PPO"). Upon subsequent oxidatio uisites. Therefore, recent years have witnessed an extensive 55 CH₃S—PPO may be quantitatively converted to the sulfonyl exploitation of polymer-nanocomposite strategies. The over-
arching goal of these efforts has be arching goal of these efforts has been to combine the best conditions, Br—PPOs having a higher degree of Br-func-
characteristics of nanofillers with polymer substrates in a tionalization (greater than 25 mol %) fails to c characteristics of nanofillers with polymer substrates in a tionalization (greater than 25 mol %) fails to convert to the synergistic fashion to improve dielectric performance of the CH₃S—PPO product because of extensive synergistic fashion to improve dielectric performance of the CH_3S —PPO product because of extensive gel formation composite materials by maximizing the dielectric constant 60 during the reaction. While wishing to not while managing the dielectric loss to an acceptable level. theory, it is believed that as an effective concentration of
From the material science standpoint, there is clearly an benzylbromide in Br—PPO is increased, the CH increasing need for high-k, nonconducting polymers (i.e., dant reacted faster than methylthiolate nucleophile with the devoid of both intrinsically electronic and ionic conduction) $_{65}$ nearby CH₂—Br pendant, resultin that are processible and compatible with high-k nanopar-
erization mixture via the formation of sulfonium salt crossticles. **links**, which is shown below: as film capacitors for power-conditioning, power electronics

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member of the PPEs family, poly(2,6-dimethyl-1,4-phe-nylene oxide), commonly known as PPO, is a useful mate-Pursuant to 37 C.F.R. § 1.78(a)(4), this application is a $\frac{1}{2}$ nylene oxide), commonly known as PPO, is a useful mate-

lent dielectric properties. For example, the dielectric connext generation of microelectronic devices because of excel stant, ε_r , of PPEs ranges between 2.4 and 2.6 and tan δ High dielectric constant or permittivity, "high-k," mate $\frac{35}{2}$ ranges between 0.002 and 0.003 (at 5 GHz for PPO). PPEs

size without compromising performance or, better still, with 45 Attempts to modify PPEs to achieve the foregoing prop-50 60

manufacture and operation and improved methods for making these polymers.

The present invention overcomes the foregoing problems
and other shortcomings, drawbacks, and challenges of con-
ventional PPEs by providing methods for synthesizing PPEs
having higher dielectric constants and lower tan invention includes all alternatives, modifications, and
equivalents as may be included within the spirit and scope In some embodiments, the units of formula (1) of the
of the present invention.

a sulfonyl-substituted PPE having improved dielectric prop- 45 erties includes sulfone-containing PPO having repeating $\overline{ }$ (3) units of the formulae:

wherein each of R^1 and R^2 is H or an alkyl group containing 65 from 1 to 4 carbon atoms, R^3 is an alkylene group containing from 1 or 2 carbon atoms, R^4 is selected from an alkyl group

 χ is a halogen atom, the subscript n is 0 or 1, the subscript containing from 1 to 4 carbon atoms, an aryl group, and CF_3 , m is 1 or 2 (provided that when m is 2 then \mathbb{R}^2 is H), the degree of polymerization ranges from about 5 to about $5\quad1,000$, and a mole ratio of units having formula (1) to units having formula (2) ranges from about 10:90 to about 90:10.

Yet other embodiments of the present invention are directed to a method for synthesizing a sulfone-containing $H_3C \longrightarrow$ $\begin{matrix} 5 \end{matrix}$ $\begin{matrix} \oplus \text{Br} \\ \text{directed to a method for synthesizing a sulfon-containing} \\ 10 \end{matrix}$ PPO and includes reacting a di-alkyl-substituted polyphenylene oxide polymer with a halogenating agent to form a halogenated PPO containing a major amount benzylic PPO and a minor amount of ring-substituted halogen-containing PPO. The halogenated PPO is then reacted with a sulfinating agent to provide the sulfone-containing

structure (1) is a bromine atom.

In other embodiments, R^1 and R^2 are selected from $-\text{CH}_3$ and $-\text{CH}_2\text{CH}_3$, R^3 is selected from $-\text{CH}_2$ — and $-\text{CH}_2\text{CH}_2$ —, and R^4 is selected from CF_3 and an aryl group.

CH₃ In some embodiments, the aryl group of R^4 contains one,
²⁵ two, or three substituents selected from —CH₃, —OCH₃, a
dimethyl amino group, and a diethylamino group. In other Thus, there remains a need for polymers having suitable dimethyl amino group, and a diethylamino group. In other
electric constants that facilitate microelectronic device embodiments, the aryl group is a para-substituted a dielectric constants that facilitate microelectronic device embodiments, the aryl group is a para-substituted aryl group
manufacture and operation and improved methods for mak-
wherein the substituent is selected from a ha group. In still other embodiments, the aryl group is an SUMMARY OF THE INVENTION ortho-substituted aryl group wherein the substituent is selected from a halide, $-\text{CH}_3$, $-\text{OCH}_3$, a dimethyl amino $- OCH₃$, a dimethyl amino group, and a diethylamino

is not limited to these embodiments. To the contrary, this $\frac{40}{10} \frac{0.40 \text{ mol}}{96}$

the present invention. Sulfone containing polymer include units of the formulas:
According to some embodiments of the present invention,

wherein a molar ratio of units of formula (3) to units of formula (4) ranges from about $64:36$ to about $99:1$.

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In other embodiments, the units of formula (2) include by practice of the invention. The objects and advantages of units of formulas:

wherein a molar ratio of units of formula (5) to units of FIG. 1C is a H-NMR spectrum (taken in CDCl₃) of a formula (6) ranges from about 1:0 to about 5:3.

selected from elemental bromine and a brominated imide 25 invention.

compound. In some embodiments, the halogenating agent is FIG. 2A is a H-NMR spectrum of sulfone-containing PPO

bromo-succinimide in admixture with azotrile). In other embodiments, a mole ratio of di-alkyl-
second with an asterisk, *, are due to protio residues of
substituted PPO to brominating agent ranges from about
deuterated dimethyl sulfoxide ("DMSO-d₆").

from sodium methane sulfinate. In other embodiments, a nylmethyl)benzene, wherein signals denoted by * are due to mole ratio of sodium methane sulfonate salt to halogenated protio residues of DMSO-d₆. PPO ranges from about 1.0:1.0 to about 3.0:1.0. In still other FIGS. 3A-3C are H-NMR spectra of Br—PPO, CH₃S—
embodiments, the sulfinating agent may be selected from but 35 PPO, and CH₃SO₂—PPO in DMSO-d₆, respecti not limited to, the following alkaline salts of trifluorometh-
anesulfinate, benzenesulfinate, 4-fluorophenylsulfinate, effects on a dielectric constant for 25 mol % CH₃SO₂—PPO
4-toluenesulfinate, 4-methoxyphenylsulfin

ethylamino-phenylsulfinate.

Other embodiments include a method for synthesizing a 40 FIG. 4B graphically illustrates temperature-dependent

sulfone-containing PPO by reacting a di-alkyl-substituted effects on dielectric l having a halogenated benzylic PPO content and a ring-
substituted halogen-containing PPO content that is less than FIGS. substituted halogen-containing PPO content that is less than FIGS. 5A-5B graphically illustrate a D-E hysteresis loop
the halogenated benzylic PPO content. The halogenated 45 for the 25 mol % CH₃SO₂—PPO film of FIGS. 4

According to embodiments of the present invention, com-
positions and methods of preparation for a series of high 50 not necessarily to scale, presenting a somewhat simplified
dielectric constant and relatively low loss de containing variable amounts of highly polar sulfonyl pen-
dans exemple, specific dimensions, orientations, locations, and
dans exemplified by methylsulfonyl (CH₃SO₂—), trifluo-
example, specific dimensions, orientatio dants exemplified by methylsulfonyl (CH_3SO_2), trifluo-example, specific dimensions, orientations, locations, and romethylsulfonyl (CF_3SO_2), phenylsulfonyl ($PhSO_2$), 55 shapes of various illustrated components, will be and 4-fluorophenylsulfonyl $(4\text{-FC}_6H_4SO_2)$ moieties are in part by the particular intended application and use envi-
described. The foregoing PPE derivatives are potentially ronment. Certain features of the illustrated described. The foregoing PPE derivatives are potentially ronment. Certain features of the illustrated embodiments useful as neat polymer films matrix polymers or polymer have been enlarged or distorted relative to others t substrates for electronic applications where high permittiv-
ity and clear understanding. In particular, thin
ity and low dielectric loss are desired such as polymer 60 features may be thickened, for example, for clarity o photovoltaics, flexible electronics, polymer capacitors, high illustration.

energy-density hybrid capacitors, and the like, as well as

coatings for electromagnetic shielding applications.

Additional objects, advantages,

Additional objects, advantages, and novel features of the invention will be set forth in part in the description which 65 follows, and in part will become apparent to those skilled in follows, and in part will become apparent to those skilled in Embodiments of the present invention provide sulfonyl-
the art upon examination of the following or may be learned substituted poly(phenylene ether)s ("PPEs") h

the invention may be realized and attained by means of the instrumentalities and combinations particularly pointed out in the appended claims.
(5) $\frac{1}{5}$

BRIEF DESCRIPTION OF THE DRAWINGS

The accompanying drawings, which are incorporated in and constitute a part of this specification, illustrate embodi-10 ments of the present invention and, together with a general description of the invention given above, and the detailed $CH_2SO_2R^4$ description of the invention given above, and the detailed description of the embodiments given below, serve to

CH₂SO₂R⁴ (6) explain the principles of the present invention.

FIG. 1A is a H-NMR spectrum (taken in CDCl₃) of a

¹⁵ sulfone-containing, Br—PPO synthesized in a carbon tetrachloride solvent according to embodiments of the present invention .

FIG. 1B is a H-NMR spectrum (taken in CDCl₃) of a $CH₂SO₂R⁴$ sulfone-containing, Br—PPO synthesized in a phenyl chlo-20 ride solvent according to embodiments of the present inven tion.

formula (6) ranges from about 1:0 to about 5:3. sulfone-containing, Br—PPO synthesized in a tetrachlo-
In other embodiments, the halogenating agent may be roethane solvent according to embodiments of the present

1.0:0.5 to about 1.0:3.0.
In some embodiments, the sulfinating agent is selected in a reaction mixture of benzyl bromide and (methylsulfo-

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improved dielectric properties and methods of synthesizing as polymeric initiators for anionic polymerization of various
the same. According to an exemplary embodiment, poly-
pherylene oxide generally, or poly(2,6-dimethyl

of sites available for chemical modification, namely the PPO, for example, radical halogenation (more specifically as polymer backbone, methyl pendant groups, and endgroups, applied to some embodiments, bromination) which polymer backbone, methyl pendant groups, and endgroups, applied to some embodiments, bromination) which may be may be considered aromatic $(sp^3C \rightarrow \text{the most ubiquitous method used because of the ease in})$

Organic and organometallic reactions may be used as spe cific tools to introduce functional groups into the polymer

making the sulfonyl-substituted PPEs described herein. $\frac{5}{4}$ According to one embodiment of the present invention, a
As can be seen from the PPO structure, below, the nature chemical transformation may be used to struc may be considered aromatic (sp^2C-H), aliphatic (sp^3C the most ubiquitous method used because of the ease in H), and protic (O—H), in that order. halogenated PPO ("X—PPO," where X is the halogen) is a versatile intermediate for introducing a wide range of functional groups to impart desired final properties (such as risk of molecular weight reduction and the possibility of CH_3 converting the linear architecture of PPO to that of a network.

> According to one specific embodiment of the present invention, PPO may be brominated with bromine in 1,1,2, 2-tetrachloroethane ("TCE") and may yield two bromine atoms per repeat unit and a substantial amount of aromatic bromination: 25

50 corresponding to the chemical nature of the sites. Thus, wherein x is a mole fraction of unreacted PPO units, y is a aromatic electrophilic substitution reactions (such as nitra-
mole fraction of benzyl brominated units aromatic electrophilic substitution reactions (such as nitra-
tion, halogenation, Friedel-Crafts acylation and sulfo-
fraction of phenyl brominated units, and the subscript n is a tion, halogenation, Friedel-Crafts acylation and sulfomation of phenyl brominated units, and the subscript n is a nylation, and sulfonation) may be used to generate PPO number of repeat unites in the Br—PPO. The mole fract nylation, and sulfonation) may be used to generate PPO number of repeat unites in the Br—PPO. The mole fractions derivatives with aromatic nitro, halide, or sulfonic acid, 55 of x, y, and z are such that $x+y+z=1$. respectively. Free radical halogenation with chlorine or
bromine at higher temperatures or under irradiation may be an embodiment of the present invention, halogenation may
a membodiment of the present invention, halogenat used to react with the $-CH_3$ pendant groups. The resulting proceed with or without a suitable halogenation agent,
PPO derivative having benzylic halogens may subsequently which is, at least partially, dependent on the ha form neutral pendants or quarternized with tertiary amines. of the present invention, PPO may be brominated with Metalation of alkyl substituted polyethers with organo-

bromine or N-bromosuccinimide ("NBS"). Reaction para Metalation of alkyl substituted polyethers with organo-
lithium compounds occurs initially on the nucleus and, on
standing or at higher temperatures, transfers to the pendant
alkyl group via Li—H exchange. However, with o

on the alkyl group. The metalated PPO derivatives will react PPO; (iii) chlorinated solvents of varied dipole moments, reagents $(Br_2$ versus NBS/azo-2,2'-isobutyronitrile ("AIBN") and NBS/benzoyl peroxide ("BPO"), (ii) a stoi(iv) temperatures, and (v) reaction times. A reaction start, there may be a nucleophilic displacement reaction with sequence with NBS/AIBN in two different solvents and at an appropriate alkylthiolate, which may then be fo

various temperatures may include: \sim oxidation of the sulfide intermediate to a sulfone product

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range of benzyl bromination may be considered as a range for the degree of functionalization resulting from the subsequent reaction to displace bromide in Br—PPO.

In order to synthesize high dielectric constant polymers,
nonpolar methyl pendants may be converted to highly polar
with conventional methods, and particularly when Br—PPO
groups, such as —CH₂SO₂Me, —CH₂SO₂CF₃, groups, such as $-\text{CH}_2\text{SO}_2\text{Me}$, $-\text{CH}_2\text{SO}_2\text{CF}_3$, and the like.
The high polarity of the foregoing groups is expected to
derive from the high polarity exhibited by the parent mol-
do was eliminated by masking or moment, 4.44+0.1 D, gas phase; 4.25 D); methyl trifluo-
romethylsulfone (CH₃SO₂CF₃; dipole moment 3.74 D); and
hope particularly, Br—PPO may be converted to RSO₂—
people moment 4.8 D). PPO by nucleophilic substitu phenyl methyl sulfone $(CH_3SO_2Ph$; dipole moment 4.8 D), PPO by nucleophilic substitution reaction of the benzyl-
which are expected to contribute to the enhanced dielectric as bromo pendants of Br—PPO by sodium organosulfi which are expected to contribute to the enhanced dielectric 45 bromo pendants of Br—PPO by sodium organosulfinate salt
constant and acceptable dielectric loss of sulfonyl-PPO. in a polar aprotic solvent, such as N,N'-di

The PPO pendant may be converted from benzyl bromide $^{(4)}M_A^4$
benzyl sulfonyl (" Bz -SO₂R") using a two-step process. At below: to benzylsulfonyl (" Bz -SO₂R") using a two-step process. At

By following an appropriate combination of the reaction $_{30}$ with an appropriate oxidizing agent (such as hydrogen parameters (i)-(v), benzyl bromination ranging from 10 mol peroxide, sodium periodate, m-chloroperbenzoi

constant and acceptable dielectric loss of sulfonyl-PPO. in a polar aprotic solvent, such as N,N'-dimethylacetamide
The PPO pendant may be converted from benzyl bromide ("DMAc"), at room temperature. Such reaction is illus

RSO2 - PPO

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The illustrative method may be used to achieve any degree $\frac{15}{20}$ Example 4—Bromination of PPO with NBS and of RSO₂-functionalization ranging from about 10 mol % to AIBN (Br—PPO with 52.0 Mol % Functionality) of RSO₂-functionalization ranging from about 10 mol % to about 80 mol %, or more.

25 invention. Furthermore, these are examples of reduction to \tilde{a} a process similar to Example 3. The mixture was heated practice of the present invention and confirmation that the under reflux for 3 hr to provide 6.4 g practice of the present invention and confirmation that the under reflux for 3 hr to provide 6.4 g (79%) of fibers. The principles described in the present invention are therefore ATR-IR peak values $\text{(cm}^{-1})$ were: 3036,

Example 1-Bromination of PPO Via NBS and AIBN (Br—PPO with 25.1 Mol % Functionality) Example 5—Bromination of PPO with NBS and

30 35 40 Commercial PPO $(4.80 \text{ g}, 40.0 \text{ mmol})$ and CCl₄ (100 mL) were added to a 500 mL, 3-necked flask equipped with a were added to a 500 mL, 3-necked flask equipped with a
magnetic stirrer, a nitrogen inlet, and an outlet, and stirred
magnetic stirrer, a nitrogen inlet, and an outlet, and stirred
magnetic stirrer, a nitrogen inlet, and under dry nitrogen at 50° C. until PPO was dissolved. NBS a process similar to Example 1. After the mixture was heated
(3.56.9.20.0 mmol) and AIBN (0.32.9) were then placed in under reflux for 1 hr, a gel formed and becam (3.56 g, 20.0 mmol) and AIBN (0.32 g) were then placed in under reflux the flask. The mixture was heated under reflux for 1 hr. The 35 any solvent. the flask. The mixture was heated under reflux for 1 hr. The viscous solution was diluted with $CCl₄$ and precipitated into methanol to afford white fibers, which was collected by methanol to afford white fibers, which was collected by Example 6—Bromination of PPO with NBS and filtration. The product was dried in an oven and provided AIBN (Br—PPO with 79.2 Mol % Functionality) 5.60 g (88%) of light brown fibers. The ATR-IR peak values (cm^{-1}) were: 3036, 2973, 2918, 2860, 1603, 1462, 1305,

Commercial PPO (4.80 g, 40.0 mmol), CCl₄ (100 mL), C₈H_{6.41}Br_{1.59}O: C, 39.10 RS (7.12 g, 40.0 mmol) and AIBN (0.32 g) were used in 39.01; H, 2.56; Br, 52.47. NBS (7.12 g, 40.0 mmol), and AIBN (0.32 g) were used in a process similar to Example 1, except the mixture was heated under reflux for 2 hr. The product was dried in oven $\frac{50}{2}$ Example 7—Bromination of PPO with NBS and and provided 6.10 g (88%) of light brown fibers (FIG. 1A). AIBN (Br—PPO with 80.1 Mol % Functionality)
The ATR-IR peak values (cm⁻¹) were: 3035, 2973, 2918,
2860, 1603, 1464, 1305, 1187, 1028, 986, 964, 860, 827, Commerc

Commercial PPO (4.80 g, 40.0 mmol), CCl_4 (100 mL), 60 NBS (7.12 g, 40.0 mmol) and AIBN (0.32 g) were used in Example 8—Bromination of PPO with NBS and a process similar to Example 1 except the mixture was AIBN (Br—PPO with 19.2 Mol % Functionality) a process similar to Example 1, except the mixture was heated under reflux for 3 hr. The yield was 6.2 g (85%). The ATR-IR peak values (cm^{-1}) were: 3035, 2980, 2918, 1603, ATR-IR peak values (cm⁻¹) were: 3035, 2980, 2918, 1603, Commercial PPO (4.80 g, 40.0 mmol), CHCl₃ (100 mL), 1464, 1305, 1219, 1189, 1030, 986, 862, 825, 629, 589, 503. 65 NBS (7.12 g, 40.0 mmol), and AIBN (0.32 g) wer Anal. Calc. (%). for $C_8H_{7.18}Br_{0.82}O$: C, 51.98; H, 3.92; Br, 35.44. Found: C, 51.71; H, 3.81; Br, 35.35.

 $RO₂S$

about 80 mol %, or more.

The following examples illustrate particular properties

Commercial PPO (4.80 g, 40.0 mmol), CCl₄ (200 mL),

and advantages of some of the embodiments of the present $_{20}$ NBS (12.82 g, 72.0 m valid but should not be construed as in any way limiting the 1464 , 1305, 1219, 1190, 1030, 986, 862, 826, 629, 589, 503.
scope of the invention.
 25 Anal. Calc (%). for C₈H_{6.94}Br_{1.06}O: C, 47.14; H, 3.44; Br, 41.5

AIBN (Br-PPO)

(cm ⁻) were: 3036, 2973, 2918, 2860, 1603, 1462, 1305,
1188, 1028, 986, 964, 860, 827, 637, 590, 504. (266 mL), NBS (16.64 g, 93.4 mmol), and AIBN (0.32 g) were used. The mixture was heated under reflux for 24 hours
to provide 9.30 g (95%) of fibers. The ATR-IR peak values Example 2—Bromination of PPO with NBS and to provide 9.30 g (95%) of fibers. The ATR-IR peak values AIBN (Br—PPO with 33.9 Mol % Functionality) $45 \text{ (cm}^{-1})$ were: 3029, 2975, 1601, 1454, 1303, 1216, 1190, 1030, 994, 865, 794, 627, 575, 509. Anal. Calc. (%) for $C_8H_{6,41}Br_{1,59}O$: C, 39.10; H, 2.64; Br, 51.72. Found: C,

 (266 mL) , NBS $(21.36 \text{ g}, 120 \text{ mmol})$, and AIBN (0.32 g) ⁵⁵ were used. The mixture was heated under reflux for 24 hr to Example 3—Bromination of PPO with NBS and provide 9.60 g (98%) of fibers. (Example 7 in FIG. 1B). The AIBN (Br—PPO with 40.7 Mol % Functionality) AIR-IR peak values (cm⁻¹) were: 3036, 2975, 1602, 1455, 1304, 1191, 1030, 995, 866, 793, 669, 628, 576, 509.

under reflux for 3 hr to afford 5.3 g $(82%)$ of fibers. The

ATR-IR peak values (cm⁻¹) were: 3036, 2922, 2917, 1601, 1464, 1305, 1219, 1190, 1189, 1022, 986, 862, 826, 634, 586.

Example 9—Bromination of PPO with NBS and $\frac{5}{5}$ BPO (Br—PPO with 9.94 Mol % Functionality)

10 Commercial PPO $(4.80 \text{ g}, 40.0 \text{ mmol})$, 1,2-dichloroethane (100 mL), NBS $(7.12 \text{ g}, 40 \text{ mmol})$, and benzyl peroxide (BPO) (0.48 g) were used in a process similar to Example 3.

The mixture was heated at 70° C. for 3 hr to afford 5.90 g

(91%) of fibers. The ATR-IR peak values (cm⁻¹) were: 3036,

2953, 2919, 1602, 1468, 1424, 1379, 13

Bromine (Br-PPO with 21.9 Mol % Benzylbromide)

Commercial PPO (4.80 g, 40.0 mmol) and 1,1,2,2-tetra-
chloroethane (100 mL) were added to a 500 mL, 3-necked 20 flask equipped with a magnetic stirrer, a nitrogen inlet, and an outlet, and stirred under dry nitrogen at 50° C. until PPO was dissolved. Then Br_2 (3.52 g, 44.0 mmol) was added slowly under reflux for 3 hr. The viscous solution was precipitated into methanol to afford white fibers, which were collected by filtration. The product was dried in oven to provide 7.40 g (97%) of light brown fibers. NMR spectra is shown in FIG. 1C. The ATR-IR peak values $(cm⁻¹)$ were: shown in FIG. IC. The ATR-IR peak values (cm^{-}) were: $3035, 2954, 2921, 1600, 1462, 1400, 1380, 1306, 1188, ₃₀$
1022, 986, 963, 857, 791, 670, 631, 590. 25

Example 11 — Bromination of PPO with 37.7 Mol % Benzylbromide

Commercial PPO (4.80 g, 40.0 mmol), Br_2 (6.40 g, 80.0³⁵ mmol), and 1,1.2.2-tetrachloroethane (100 mL) were used. The mixture was heated under refluxing for 3 hr to afford 7.4 g (96%) of light brown fibers. The ATR-IR peak values (cm⁻¹) were: 3035, 2954, 2921, 1600, 1462, 1400, 1380, 1306, 1188, 1022, 986, 963, 857, 791, 670, 631, 590.
Table 1, below, is a summary of the reaction conditions 40

and results for the above bromination examples.

wherein 1 the molar ratio is bromination agent to PPO, ²the mol % of Methyl-Br and mol % of Aryl-Br were calculated from NMR spectra (FIGS. 1A-1C) and in accordance with the Equations 1 and 2 (below), and ³ gelati

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driven by free-radical chain process, were found to be temperature until Br—PPO was dissolved. Then sodium effective in minimizing the formation of aryl bromide (Ex-
methanesulfinate (0.75 g, 7.3 mmol) was added to the amples 1 to 4), based on the quantitative proton NMR mixture in the flask. The mixture was agitated overnight at comparison as depicted in FIGS. 1A-1C. With reference to 20 room temperature and poured into water/methano comparison as depicted in FIGS. $1A-1C$. With reference to 20 FIGS. $1A-1C$, proton-NMR spectra (CDCl₃) of Gr-PPO in a FIGS. 1A-1C, proton-NMR spectra (CDCl₃) of Gr-PPO in a vent (1:1). The off-white solid was collected by filtration and CCl₄ solvent (FIG. 1A; Example 2), Br—PPO in chloroben-
dried in a vacuum oven to provide 1.38 g (zene solvent (FIG. 1B; Example 7), and Br—PPO in tetra-
choroethylene solvent (FIG. 1C; Example 10) are shown.
The ATR-IR peak values $\text{(cm}^{-1})$ were: 2981, 2929, 2886,
chloroethylene solvent (FIG. 1C; Example 10) are sho aromatic PPO repeat unit) and 3 for CH₂Br (i.e., the desired

Reaction time is important and, under the conditions described above, should be conducted for no more than 3 hr. ³⁰
After this 3 hr reaction time, Br—PPO starts to precipitate After this 3 hr reaction time, Br—PPO starts to precipitate Following the procedure described in Example 12, Br—
out and form an insoluble gel if the reaction is allowed to PPO (Example 2; 1.73 g, 10.0 mmol), methane sodiu out and form an insolution get if the reaction is anowed to
continue after the first sign of precipitation (Example 5).
Because of this constraint, the highest percentage of bro-
mine on methyl group is 52.0 mol % when CC as the solvent. Thus, to keep the Br—PPO in solution, The foregoing Examples 12 and 13 demonstrate a two-
chlorobenzene with greater solvating nower was used and step process for preparing methylsulfonyl-PPO having at chlorobenzene with greater solvating power was used and, step process for preparing methylsulfonyl-PPO having at indeed, the percentage of bromine on the methyl group was least 25.1 mol % benzylsulfide. However, as shown i indeed, the percentage of bromine on the methyl group was raised to about 80 mol % (Examples 6 and 7).

temperature until Br-PPO was dissolved to form a yellow 50 solution. Then, the sodium salt of methane sulfinic acid $(0.35 \text{ g}, 5.0 \text{ mmol})$ was added slowly. The mixture was agitated overnight at room temperature and poured into
methanol. The white powder was collected by filtration and
solvent methanol. The white powder was collected by filtration and dried in an oven to afford 1.34 g (94%) of product. The 55
ATR-IR peak values (cm^{-1}) were: 2982, 2971, 2917, 1598, 1460, 1420, 1397, 1381, 1304, 1183, 1021, 992, 958, 852. PPO

Example 13 — Preparation of Methylsulfonyl - PPO $\left(\text{CH}_3\text{SO}_2\text{—PPO with }25.1 \text{ Mol }\%$ Functionality 60 CO_3 CO_2N $\text{CH}_3\text{SO}_2\text{Na}$

Oxidation of methylsulfide pendants by meta-chlorope-
oxybenzoic acid: CH₃S—PPO (Example 12: 1.0 g, 7.0 mmol) and CHCl₃ (40 mL) were added to a 100 mL, 3-necked flask equipped with a magnetic stirrer, a nitrogen 65 inlet, and an outlet, and stirred under dry nitrogen at room temperature until Br-PPO was dissolved. Then the flask

was cooled to 0° C. and meta-chloroperoxybenzoic acid Intergration Value of Peak 2 addition of mCPBA, a temperature of the reaction mixture
Intergration Value of Peak 2 + was allowed to rise to room temperature for 2 hr, and the was allowed to rise to room temperature for 2 hr, and the mixture was precipitated into methanol. The white precipi-

Br—PPO repeat unit) in FIGS. 1A-1C.

Reaction time is important and under the conditions (CH₃S—PPO with 33.9 Mol % Functionality)

ised to about 80 mol % (Examples 6 and 7). Example 14, for benzylsulfide functionality above 25 mol Functionalization of Br—PPO with a polar group is 40% , crosslinking and gelation occur. To circumvent the prob-Functional methillustrated by the following examples.
 $\frac{1}{1-\theta}$ and $\frac{1}{1-\theta}$ ods, the nucleophilic character of the methylsulfido pendant Example 12—Preparation of Thiomethoxide-PPO

(CH₃S—PPO with 25.1 Mol % Benzylsulfide)

Br—PPO (Example 1: 1.59 g, 10.0 mmol) and dimethy-

accuse organic sulfinate salts (e.g., RSO₂-Na⁺) react easily

with alkyl or

wherein R is selected from the group consisting of $-CH_3$, 1292, 1235, 1194, 1152, 1086, 1031, 988, 839, 820, 740.
 $-CF_3$, $- C_6H_5$, $- C_6H_4CH_3$, and $-C_6H_4F$, as illustrated, 15 In order to confirm the feasibility of

mL) were added to a 100 mL, 3-necked flask equipped with proton NMR spectrum of the reaction mixture, shown in
a magnetic stirrer a nitrogen inlet and an outlet and stirred FIG. 2B, indicates the complete disappearance of a magnetic stirrer, a nitrogen inlet, and an outlet, and stirred
in the start-
ing benzyl bromide benzyl proton signal, the concomitant
inder dry nitrogen at room temperature until Br—PPO was
ing benzyl bromide benzyl prot under dry nitrogen at room temperature until Br—PPO was ing benzyl bromide benzyl proton signal, the concomitant
dissolved. Sodium methanesulfinate (0.32 g, 2.7 mmol) was appearance of benzyl proton signal of the sulfone p dissolved. Sodium methanesulfinate (0.32 g, 2.7 mmol) was appearance of benzyl proton signal of the sulfone product at added to the mixture in the flask. After the mixture was 25 lower chemical shift, and a new —CH₃ p agitated at room temperature overnight it was poured into
water/methanol co-solvent (1:1). The brown solid was col-
mixture of benzyl bromide (FIG. 2A) and methylsulfonyl-
 lected by filtration and dried in a vacuum oven to provide

0.41 g (82%) of product. The ATR-IR peak values (cm^{-1})

converted into (methylsulfonylmethyl)benzene at room tem-0.41 g (82%) of product. The ATR-IR peak values $\text{(cm}^{-1})$ converted into (methylsulfonylmethyl) benzene at room tem-
were: 3015, 2930, 1602, 1464, 1301, 1248, 1192, 1160, 30 perature in DMSO-d₆. A small amount of be were: 3015, 2930, 1602, 1464, 1301, 1248, 1192, 1160, 30
1118, 1030, 989, 963, 897, 757, 491, 450.

Example 18—Benzenesulfonyl-PPO $(C_6H_5SO_2$ and % .

PPO with 50 Mol % Functionality and the polymers described herein Example 20, except

were: 3063 , 1606 , 1466 , 148 , 1307 , 1194 , 1156 , 1132 , The molecular weight and polydispersity ("PDI") of the 1085, 1028, 988, 877, 737.

used in a process similar to Example 15 to provide 0.70 g
 $\binom{CH_3}{(cm^{-1})}$ (100%) of a light-yellow powder. The ATR-IR peak values
 $\binom{cm^{-1}}{(cm^{-1})}$ were: 2926, 1600, 1465, 1319, 1303, 1245, 1194.

 $(4-FC₆H₅SO₂—PPO with 50 Mol % Functionality)$

Br—PPO (Example 4; 0.50 g, 2.5 mmol), sodium 4-fluo-10 robenzenesulfonate (0.90 g, 4.9 mmol), and DMAc (20 mL) RSO₂-PPO were used in a process similar to Example 15 to provide 0.63 g (91%) of an off-white powder. The ATR-IR peak values (cm^{-1}) were: 3074, 2928, 1591, 1493, 1466, 1405, 1322,

Example 15—Preparation of Methylsulfonyl-PPO

(CH₃SO₂—PPO with 50 Mol % Functionality)

with sodium methanesulfinate (MeSO₂Na) at room tempera-

ture in DMSO. An instantaneous and clean reaction to

PDO (F₁ 1 A 0. Br—PPO (Example 4; 0.50 g, 2.5 mmol) and DMAc (20 20 produce (methylsulfonylmethyl)benzene was observed. The proton NMR spectrum of the reaction mixture, shown in $(-CH2- at d 3.35 and OH at d 4.70, denoted by $*$) was also formed from the reaction of excess benzyl bromide with$

Example 16—Methylsulfonyl-PPO (CH₃SO₂—PPO) water present in DMSO-d₆.
with 80 Mol % Functionality, 4-3) Thus, in the similar reaction, chemically modified PPO
and 2.0 Clare 2.0 minutes in the similar reaction, chemic Br—PPO (Example 7; 0.50 g, 2.0 mmol), sodium meth-
anesulfinate (0.64 g, 5.4 mmol), and DMAc (20 mL) were
used in a process similar to Example 15 to provide 0.46 g
(0.2%) of a heavy neurlar The ATP IP pools volume (om-1) (92%) of a brown powder. The ATR-IR peak values cm^{-1} the comparison of the proton NMR spectra of the starting
ware: 2081, 2080, 1602, 1460, 1405, 1203, 1250, 1104 Br—PPO and the isolated CH₃SO₂—PPO, together wit were: 2981, 2930, 1602, 1460, 1405, 1293, 1250, 1194, Br—PPO and the isolated CH₃SO₂—PPO, together with the 1157, 1114, 991, 961, 884, 756, 685, 448. and benzylmethylsulfone, has validated the quantitative
Example 17—Trifluoromethylsulfonyl-PPO
CF-SO₂—PPO with 50 Mol % Functionality) extended to RSO₂Na with R other than methyl, especially (CF₃SO₂—PPO with 50 Mol % Functionality)
Br—PPO (Example 4; 0.50 g, 2.5 mmol), sodium trifluo-
Br—PPO (Example 4; 0.50 g, 2.5 mmol), sodium trifluo-
onethanesulfinate (0.50 g, 3.2 mmol), and DMAc (20 mL)
of Br—PPO (FI were used in a process similar to Example 15 to provide 0.62 7), CH₃S—PPO (FIG. 3B, spectrum taking in CDC1₃; g (98%) of a light-yellow powder. The ATR-IR peak values Examples 12 and 13), and CH₃SO₂—PPO (FIG. 3C (cm^{-1}) were: 2937, 1605, 1369, 1307, 1196, 1119, 1030, trum taking in DMSO-d₆; Example 16). Both Br—PPO and 988, 870, 769, 729. $CH₃SO₂$ —PPO had the degree of functionalization of 80 mol %. $CH₃S$ —PPO had the degree of functionalization of

 $CH₃SO₂$ —PPO containing 80 mol % functionality, are soluble in CHCl₃, chlorobenzene, tetrahydrofuran ("THF"), Br—PPO (Example 4; 0.50 g, 2.5 mmol), sodium benze- 55 soluble in CHCl₃, chlorobenzene, tetrahydrofuran ("THF"), nesulfonate (0.85 g, 5.2 mmol), and DMAc (20 mL) were DMAc and DMSO. Highly polar CH₃SO₂—PPO containin

Example 19—4-Toluenesulfonyl-PPO

Example 19—4-Toluenesulfonyl-PPO

(P-MEC₆H₄SO₂—PPO (with 50 Mol % Functional-

ited comparable GPC results to a high molecular weight ity)

ity) PPO sample (HMW PPO) (Table 2). However, Br—PPO

65 and CH₃S—PPO polymers exhibited exceptionally high

enesulfonate (0.85 g, 4.4 mmol), and DMAc (20 mL) were

unmodified PPO) and narrow PDIs (decreasing from unmodified PPO) and narrow PDIs (decreasing from 3 to

aggregated in THF solution. The molecular weight of films for use as high energy density capacitors, the dielectric CH₂SO₂—PPO was only slightly higher than PPO, as properties of the films were determined using broadba $CH₃SO₂$ —PPO was only slightly higher than PPO, as shown in Table 2.

PPO MOLECULAR WEIGHTS AND POLYDISPERSITY								
Sample No	Sample Name	M.,	M.,	PDI	10			
Commercial $PPO-1$	$PPO-1$	36,500	112,000	3.06				
Commercial $PPO-2$	$PPO-2$	21,700	68,600	3.20				
Example 2	$Br-PPO$ (34 mol %)	240,000	333,000	1.39				
Example 4	Br-PPO (52 mol %)	338,000	466,000	1.38	15			
Example 6	Br-PPO (79 mol %)	210,000	304,000	1.52				
Example 12	$CH3S-PPO (25 mol %)$	254,000	469,000	1.84				
Example 13	$CH3SO2-PPO$ (25 mol %)	247,000	472,000	1.91				
Example 16	CH ₃ S-PPO (80 mol %)	61,200	120,000	1.96				

groups prevent the PPO main chains from rotating freely. I_g 30 during the temperature ramping and recorded the resulting values for CH₃SO₂—PPOs were equal to or higher than T_g current. Similarly, the aluminum-coat The commercially-available PPO possessed a glass tran-
sition temperature (" T_g ") at 212° C. and good thermal
stability under inert atmosphere. Incorporation of bromine
(less than 52 mol %) and thiomethoxide resulted in CH₃SO₂—PPO, respectively since the modified side groups
would increase the free volume and impart plasticizing
elimearly -140° C. to 150° C., at a rate of 2° C./min. The BDS
effect. The T_e of Br—PPO containing 80 mo effect. The T_g of Br—PPO containing 80 mol % of func-
tionality increased to 250° C. since the bulky bromide from 107 Hz to 1 Hz across a sample film every 14 sec tionality increased to 250° C. since the bulky bromide from 107 Hz to 1 Hz across a sample film every 14 sec groups prevent the PPO main chains from rotating freely. T_g 30 during the temperature ramping and recorded the values of PPO due to the high polarity of side chains, and the used in D-E loop measurements performed at 23° C., 100°
thermally stabilities of all were all substantially reduced. A C., and 190° C. The D-E summary of the thermal properties of the PPO is provided in ferroelectric analyzer by applying two consecutive sine
Table 3.
Table 3.

THERMAL PROPERTIES OF PPO AND ITS DERIVATIVES						ments are summarized and compare				
	Example No.	Sample Name	Tg ¹ $(^{\circ}$ C.)	$T_{d5\%}^2$ $(^{\circ}$ C.) in air	$T_{d5\%}^2$ $(^{\circ}$ C.) in nitrogen	40	for polymer dielectrics, <i>i.e.</i> , biaxia ene ("BOPP") in Table 4.			
	Commercial	$PPO-2$	212	390	423					TABLE 4
	$PPO-2$ 2 $\overline{4}$ 6	Br-PPO Br-PPO Br-PPO Br-PPO	208 204 201 250	281 277 268 248	285 277 270 234	45	Sample ID	\in . $(1$ kHz. 25° C.)	€." $(1$ kHz. 25° C.)	$tan \delta$ $(1$ kHz, 25° C.
	12	$CH3S-PPO$	181	283	287					
	13	$(25 \text{ mol } \%)^3$ $CH3SO2-PPO$ $(25 \text{ mol } \%)^4$	211	332	339	50	BOPP PPO PPO-SMe	2.2 $2.7 - 2.8$ 3.5	< 0.028 0.01	0.0002 < 0.001 0.0028
	15	$CH3SO2-PPO$ $(50 \text{ mol } \%)^5$	228	327	336		25 mol % PPO-SO ₂ Me	6.0	0.06	0.008
	16	$CH3SO2-PPO$ $(80 \text{ mol } \%)^5$	262	331	340		$(25 \text{ mol } %$ PPO-SO ₂ Me	8.0	0.065	0.008

wherein ¹ T_g was measured from mid-point of change in slope on DSC thermogram obtained with a heating rate of 10° C./min, ² temperature at which 5% weight loss recorded 10° C./min, ² temperature at which 5% weight loss recorded
on TGA thermogram obtained with a heating rate of 10° $_{60}$ FIGS. 4A and 4B include present BDS spectra of real ϵ ,
C./min, ³ prepared from Br—PPO (Example nearly identical thermal properties were observed for CH_3SO_2 —PPO (Example 13). One obvious transition could CH₃SO₂—PPO prepared by nucleophilic substitution of be identified between –140 and 150°C, with an electric CH₃SO₂—PPO prepared by nucleophilic substitution of be identified between -140 and 150° C. with an electric field Br—PPO (25 mol %) with sodium methanesulfinate, and ⁵ 65 at 100 Hz, that is, at about -90° C. Above prepared from the respective Br—PPO via nucleophilic reached a peak value of 6.0. The foregoing transition may be substitution with sodium methanesulfinate. 60

less than 2), which indicates a likelihood that the polymers In order to evaluate the suitability of the RSO_2 —PPO aggregated in THF solution. The molecular weight of films for use as high energy density capacitors, t dielectric spectroscopy ("BDS") and dielectric displace-
5 ment-electric field (D-E) loop measurements. BDS measurement-electric field (D-E) loop measurements. BDS measurements of the γ transition reveal how much dipole switching TABLE 2

PPO MOLECULAR WEIGHTS AND POLYDISPERSITY

PPO MOLECULAR WEIGHTS AND POLYDISPERSITY

M_n PDI

M_n PDI

10 Feld performance of the CH₃SO₂—PPO films, to measure the dielectric constant at high field strengths, to estimate lossiness of the material based on the hysteresis, and to determine the breakdown strength. Discharge energy and efficiency were determined from D-E loop data.

> In order to determine the dielectric properties of the polymer samples, the samples were dried in a vacuum oven for 24 hr at 130 $^{\circ}$ C. The samples were then silver coated (100 nm), on both sides, by physical vapor deposition to improve electrical contact. The coated samples were kept in 20 a desiccator filled with desiccant until measurements were

C., and 190° C. The D-E loops were measured with a ferroelectric analyzer by applying two consecutive sine sample. The peak amplitude of the voltage was increased TABLE 3 every two cycles beginning from field strengths of 50 MV/m and with 316 MV/m. The results of the dielectric measurements are summarized and compared with a current standard
40 for polymer dielectrics, i.e., biaxially oriented polypropylfor polymer dielectrics, i.e., biaxially oriented polypropylene ("BOPP") in Table 4.

1 1 U – 4	---	<i>.,,</i> ,	᠇∠ၪ							
Br-PPO Br-PPO Br-PPO Br-PPO CH ₃ S-PPO	208 204 201 250 181	281 277 268 248 283	285 277 270 234 287	45	Sample ID	\in . $(1$ kHz, 25° C.)	€." (1 kHz) 25° C.)	$tan \delta$ $(1$ kHz, 25° C.)	Released Energy Density (J/cm ²)	Maxima Breakdown Field Strength (MV/m)
$(25 \text{ mol } \%)^3$					BOPP	2.2		0.0002	4.8	730
CH ₃ SO ₂ -PPO	211	332	339	50	PPO.	$2.7 - 2.8$	< 0.028	< 0.001		
$(25 \mod 96)^4$					PPO-SMe	3.5	0.01	0.0028	冰	家
$CH3SO2-PPO$	228	327	336		25 mol %					
$(50 \text{ mol } \%)^5$					PPO-SO ₂ Me	6.0	0.06	0.008	30	900
$CH3SO2-PPO$	262	331	340		$(25 \text{ mol } %$				(at 200 MV/m)	
$(80 \text{ mol } \%)^5$					PPO-SO ₂ Me	8.0	0.065	0.008	28	800
				55	$(50 \text{ mol } %$				(at 240 MV/m)	
					PPO-SO ₂ Me	$-8.0*$	宋	冰	宋	冰
s measured from mid-point of change in					$(80 \text{ mol } %$					

assigned as a γ transition resulting from the rotation of the

sulfone groups, which can not only increase dielectric con-
stant, but also maintain a dielectric loss as low as ε_r ^{"=0.02}. (Example 13) had a U_{e,discharged} of 18 J/cm³ and a relatively There was a slight decrease in ε_r starting from about 0° C. high discharge efficiency about 91% (see FIG. 5B). The Slight decrease in ε_r may be due to the fact that as discharge efficiency of 50 mol % CH₃SO₂—P temperature increases, dipoles gained more thermal energy 5 15) at about 635 MV/m was about 85% and dramatically
and the amplitude of random thermal motion increases, decreased with increased the electric field, as show transition was observed in the spectrum of FIG. 4B, which Accordingly, increasing the amount of sulfone groups
may be assigned as a weak β transition associated with the 10 attached on the side chain of the polymer may may be assigned as a weak β transition associated with the 10 attached on the side chain of the polymer may not neces-
waggling of ether groups along a polymer backbone. When sarily result in a better dipolar glass pol absorbed moisture and impurity ions in the polymer matrix. these embodiments have been described in considerable
While even 25 mol % CH₃SO₂—PPO (Example 13) dem-15 detail, they are not intended to restrict or in any wa While even 25 mol % CH_3SO_2 —PPO (Example 13) dem- 15 onstrated strong dipoles, CH_3SO_2 —PPO was not sensitive onstrated strong dipoles, CH_3SO_2 —PPO was not sensitive scope of the appended claims to such detail. Additional to moisture and was capable of keeping a low dielectric loss advantages and modifications will readily appea to moisture and was capable of keeping a low dielectric loss advantages and modifications will readily appear to those up to 150° C. These properties are very attractive for skilled in the art. The invention in its broader

hysteresis loop measurements. At room temperature and at described. Accordingly, departures may be made from such 1000 Hz, 25 mol % CH₃SO₂—PPO (Example 13) exhibited details without departing from the scope of the gene 1000 Hz, 25 mol % CH₃SO₂—PPO (Example 13) exhibited details without departing from the scope of the general narrow bipolar linear loops at different electric fields ranging inventive concept. from 105 MV/m to 650 MV/m, as shown in FIG. 5A. The
apparent k was determined to be 6.2 via k= $D/(e_0E)$, which 25
was consistent with the result obtained from BDS data. The
linear nature of the loops suggests that sulfonu linear nature of the loops suggests that sulfonyl groups phenylene oxide (PPO) polymer comprising.

reacting a di-alkyl-substituted PPO polymer with a halorotated easily without dipole coupling under a high electric reacting a di-alkyl-substituted PPO polymer with a halo-
field When the electric field was further increased such as field. When the electric field was further increased, such as genating agent to form a natiogenated PPO polymer
to 800 MV/m diological loss became natiogenable (measured 20 to 800 MV/m, dielectric loss became noticeable (measured 30 having a halogenated benzylic PPO polymer content
havit be englosed area inside the uningler loops in FIG. ED. by the enclosed area inside the unipolar loops in FIG. 5B).
The degree of disloctic loss increased with field hasove of mer content that is less than the halogenated benzylic The degree of dielectric loss increased with field because of mer content that is less than the halogenated benzylic content that is less than the halogenated benzylic content that is less than the halogenated benzylic con either increased ionic conduction or electronic conduction at PPO polymer content; and
high field At the highest testing field (25 mol %) the reacting the halogenated PPO polymer content with a high field. At the highest testing field (25 mol %), the reacting the halogenated PPO polymer content with a dialectric of CH SO ppO (Example 13) reached as bigh as sulfinating agent to provide the sulfone-containing PPO dielectric of CH_3SO_2 —PPO (Example 13) reached as high 35 sulfinating agent to provide the sulfone-containing PPO as 913 MV/m, above which breakdown occurred. Such high polymer by nuclear polymer at an angle $\frac{p}{p}$ breakdown strength may be attributed to: (1) the rigid PPO $\frac{1}{2}$. The method of claim 1, wherein the halogenating agent backbone containing aromatic rings suppressed segmental 2. The method of claim 1, wherein the halogenating agent is selected from the group consisting of elemental bromine motion and thus weakened charge hopping or tunneling is selected from the group consisting under a high electric field: (2) the coulombic interaction ϕ and a brominated imide compound. under a high electric field; (2) the coulombic interaction 40 and a bromhladed imide compound.

between dipoles and charge carriers caused extra scattering

in addition to phonon-electron scattering, which reduced the

com conduction current and prevented dielectric breakdown by $\frac{dy}{dx}$. The method of claim 2, wherein a mole ratio of stabilizing the electron energy; (3) the polar groups served as the polar groups served as the polar group as traps of charge carriers; and (4) a uniform film with a 45 di-alkyl-substituted PPO polymer to smooth film surface lowers surface-defects levels.

The discharge energy density ($U_{e,discharge}$) and discharge

The discharge energy density ($U_{e,discharge}$) and discharge

efficiency were calculated from the electric displacement-

electric displacement

electric displacement

of $CH₃SO₂$ -PPO (Example 13) and 50 mol % $CH₃SO₂$ so methane-surfonate sait to halogenated PPO polymer ranges
PPO (Example 15) provided in FIGS. 5 and 6, respectively.
With increasing external electric f tially increased up to 22 J/cm³ and 24 J/cm³ for 25 mol % comprises a compound selected from the group consisting of CH₃SO₂—PPO (Example 13) and 50 mol % CH₃SO₂—
PO (Example 15), respectively. The discharge energy den-
sity, $U_{e,discharge}$ of 50 mol % CH₃SO₂—PPO (Example 15)
sity, $U_{e,discharge}$ of 50 mol % CH₃SO₂—PPO (Example 1 increased faster than 25 mol % CH₂SO2 \geq PPO (Example 15) **8**. A sulfone-containing PPO polymer made by the pro-
13), likely because 50 mol % CH₂SO₂ - PPO (Example 15) cess of claim 1, wherein the sulfone-contai has a higher dielectric constant. However, 25 mol % cess of claim 1, wherein the sulfone-containing PPO poly-
CH SO BPO (Example 12) oxidiated lower dialectric loss 6. mer comprises from about 10 mol % to about 90 mol % $CH₃SO₂$ -PPO (Example 13) exhibited lower dielectric loss 60 mer comprises for about 90 mol $\frac{100}{90}$ mol $\frac{100}{90}$ mol $\frac{100}{90}$ mol and higher discharge efficiency (defined as 100 $(1-U_{e,discharged}/U_{e,stored})\%)$. For example, upon exposure to $CH₃SO₂$ -PPO (Example 13) and 50 mol % $CH₃SO₂$ 50

description of one or more embodiments thereof and while these embodiments have been described in considerable practical dielectric applications. therefore not limited to the specific details, representative
High-field dielectric properties were explored by D-E 20 apparatus and method, and illustrative examples shown and
hysteresis

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-