

*OST!
Rec'd
9/24/99*

DOE F 1332.16 (10-84)
(Formerly RA-427)

OMB Approval
No. 1910-1400

U. S. DEPARTMENT OF ENERGY

UNIVERSITY CONTRACTOR, GRANTEE, AND COOPERATIVE AGREEMENT.
RECOMMENDATIONS FOR ANNOUNCEMENT AND DISTRIBUTION OF DOCUMENTS

See Instructions on Reverse Side

1. DOE Report No. DOE/ER/12119-2		3. Title SYNTHESIS OF NEW HIGH PERFORMANCE LUBRICANTS AND SOLID LUBRICANTS	
2. DOE Contract No. DE-FG05-91ER12119			
4. Type of Document ("x" one)			
<input checked="" type="checkbox"/> a. Scientific and technical report			
<input type="checkbox"/> b. Conference paper: Title of conference _____			
Date of conference _____			
Exact location of conference _____			
Sponsoring organization _____			
<input type="checkbox"/> c. Other (Specify) _____			
5. Recommended Announcement and Distribution ("x" one)			
<input checked="" type="checkbox"/> a. Unrestricted unlimited distribution.			
<input type="checkbox"/> b. Make available only within DOE and to DOE contractors and other U. S. Government agencies and their contractors.			
<input type="checkbox"/> c. Other (Specify) _____			
6. Reason for Recommended Restrictions _____			
7. Patent and Copyright Information:			
Does this information product disclose any new equipment, process, or material? <input checked="" type="checkbox"/> No <input type="checkbox"/> Yes If so, identify page nos. _____			
Has an invention disclosure been submitted to DOE covering any aspect of this information product? <input checked="" type="checkbox"/> No <input type="checkbox"/> Yes			
If so, identify the DOE (or other) disclosure number and to whom the disclosure was submitted. _____			
Are there any patent-related objections to the release of this information product? <input checked="" type="checkbox"/> No <input type="checkbox"/> Yes If so, state these objections. _____			
Does this information product contain copyrighted material? <input checked="" type="checkbox"/> No <input type="checkbox"/> Yes			
If so, identify the page numbers _____ and attach the license or other authority for the government to reproduce.			
8. Submitted by Professor Richard J. Lagow		Name and Position (Please print or type)	
Organization The University of Texas at Austin			
Signature <i>Richard J. Lagow</i>		Phone (512) 471-1032	Date 4/8/93

FOR DOE OR OTHER AUTHORIZED
USE ONLY

9. Patent Clearance ("x" one)
- a. DOE patent clearance has been granted by responsible DOE patent group.
 - b. Report has been sent to responsible DOE patent group for clearance.

APPROVED FOR RELEASE OR
PUBLICATION. INTEL. PROP. GP.
OFC. OF CHIEF COUNSEL, DOE/ORO
By *Edwin* Date *9/23/99*

DISCLAIMER

This report was prepared as an account of work sponsored by an agency of the United States Government. Neither the United States Government nor any agency thereof, nor any of their employees, make any warranty, express or implied, or assumes any legal liability or responsibility for the accuracy, completeness, or usefulness of any information, apparatus, product, or process disclosed, or represents that its use would not infringe privately owned rights. Reference herein to any specific commercial product, process, or service by trade name, trademark, manufacturer, or otherwise does not necessarily constitute or imply its endorsement, recommendation, or favoring by the United States Government or any agency thereof. The views and opinions of authors expressed herein do not necessarily state or reflect those of the United States Government or any agency thereof.

DISCLAIMER

Portions of this document may be illegible in electronic image products. Images are produced from the best available original document.

DOE/ER/12119-2

SYNTHESIS OF NEW HIGH PERFORMANCE LUBRICANTS
AND SOLID LUBRICANTS

Progress Report

April 1992 - March 1993

RECEIVED
SEP 24 1999
UST

Professor Richard J. Lagow

Department of Chemistry

The University of Texas at Austin

Austin, Texas 78712-1167

April 1993

PREPARED FOR THE U.S. DEPARTMENT OF ENERGY
UNDER GRANT NUMBER DE-FG05-91ER12119

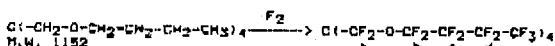
SYNTHESIS OF NEW HIGH PERFORMANCE LUBRICANTS AND SOLID LUBRICANTS

Technical Progress Report

In our second year of funding we began the testing phase of a number of new classes of lubricants. Three different testing collaborations have already begun and a fourth one is in the works with Dr. Stephen Hsu of the National Institute of Standards and Technology with whom we had established a working relationship after meeting at the Automotive and Technology Development Coordination Meeting held in November 2-5, 1992 in Dearborn, Michigan. Dr. Hsu also plans to test some of the same materials for us that Shell Development is studying.

With Dr. Bill Jones of NASA, we are studying the effects of branching on high temperature lubricant properties in perfluoropolyethers. Initially Bill Jones is comparing the lubrication and physical properties of perfluoro-tetraglyme and the following two spherical perfluoropolyethers. Note that one contains a fluorocarbon chain and the other one contains a fluorocarbon ether chain. The synthesis of these was reported in the last progress report.

Perfluoro(pentaerythrityl tetraethyl ether)



17 N.M.R. data: ppm, $\delta(CFCl_3)$

CF ₃	(a) -65.3
CF ₂	(b) -82.9
CF ₂	(c) -126.2
CF ₂	(d) -126.7
CF ₂	(e) -81.8

Mass Spectral data:

(P-F)	m/m	1133
(P-C ₄ F ₉ O)		917
C ₁₃ F ₃₃ O ₄		895
C ₁₃ F ₂₉ O ₃		678
C ₆ F ₁₅ O		397
C ₄ F ₉		219
C ₄ F ₇		181
C ₃ F ₅		131 (base peak)
C ₂ F ₅		119
C ₂ F ₄		100

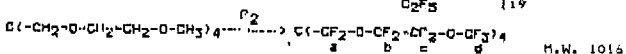
Elemental Analysis:
not available

Perfluoro(pentaerythrityl tetraethyl ether)

18 N.M.R. data: ppm, $\delta(CFCl_3)$

(a) -66.3
(b) -89.0
(c) -71.2
(d) -56.3

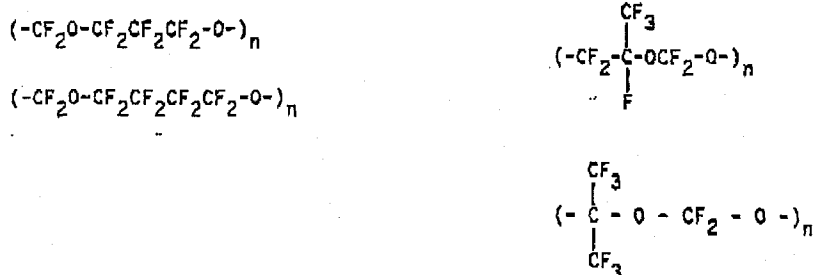
Elemental Analysis:
not available



Mass Spectral data

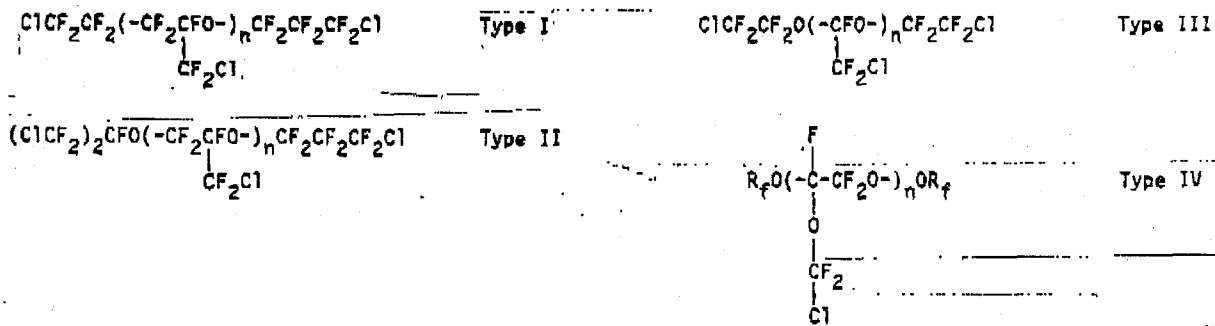
(P-F)	m/m	797 (base)
(P-C ₂ F ₅ O)		681
(P-C ₃ F ₇ O ₂)		813
C ₁₄ F ₂₇ O ₇		793
C ₁₃ F ₂₅ O ₆		727
C ₁₁ F ₂₁ O ₅		611
C ₇ F ₁₁ O ₃		341
C ₃ F ₇ O ₂		201
C ₃ F ₇ O		185
C ₂ F ₅		119

With Professor Patricia Thiel of Iowa State University, we are working on studies of perfluoromethylene oxide ethers and have prepared a series of four of these polyethers to study in collaboration with her research group. These are model compounds which correspond to structures proposed on the bottom of page 17 and top of page 18 of our research proposal:

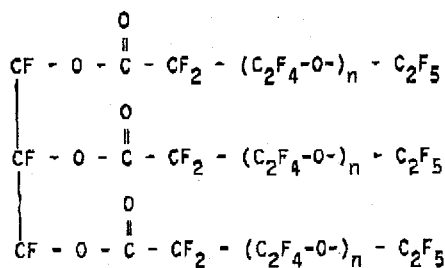


These perfluoromethylene oxide ethers have the best low temperature properties of any known lubricants. Thiel's group is studying their interactions with metals under extreme conditions.

Thirdly, we have also begun an interaction with Dr. August Birke of Shell Development Company in Houston for whom we have already prepared samples of the chlorine-substituted fluorocarbon polyether lubricants whose structures appear on page 54 of our research proposal. Each of these four structures is thought to have potential as lubricant additives to motor oils. Each of the following structures is completely soluble in hydrocarbon motor oils and hydrocarbon polyalphaolefins.



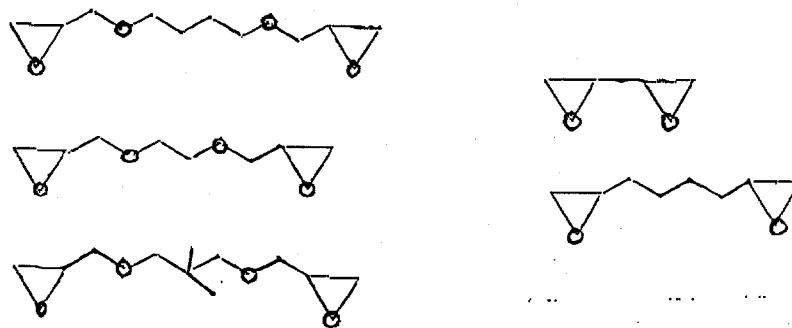
We also have underway syntheses of other fluorine-containing branched ether lubricants. These new materials which are also promising as antifriction additives for motor oils appear ahead of the perfluoro additives as Appendix I to the progress report. Additionally for Birke and Shell Development we have at their request prepared the novel compound perfluoro salicylic acid. This synthesis was suggested by the Shell staff who thought that esters of perfluoro salicylic acid might be an excellent antifriction additive for motor oil fuels. One of the best additives currently used in motor oils is the hydrocarbon ester of salicylic acid.



where n is varied to produce fluids and solids with different properties

Dr. Kuangsen Sung of our research group has succeeded in preparing the first example of glycerin-based perfluoropolyester structures and specifically has succeeded in preparing the glyceride ester of perfluoro stearic acid, $\text{CF}_3(\text{CF}_2)_{16}\text{COOH}$. We shall be submitting this new class of branched perfluoro-carbon esters for testing shortly.

We also have achieved success with synthesis of perfluoro epoxy ether chains, a class of compounds that have never been previously prepared:



Additionally with Dr. Bill Jones of NASA we are testing a new class of antifriction additives for perfluoropolyether lubricants, the perfluorophosphoranes. We have made quite a number of these and will shortly be submitted these to Dr. Jones for screening. The first structure appears below and the rest of the new structures constitute Appendix II.

^{19}F NMR Chemical Shifts

a - 47.5 ppm (d of mult.)

b -110.3 ppm (d of t)

c -125.7 (s)

d - 82.2 (s)

Coupling Constants

$^1J_{\text{PF}}=1041$ Hz

$^2J_{\text{FCP}}=124$ Hz



^{31}P NMR Chemical Shift

-41.8 ppm (t of sept.)

Low Resolution Mass

Spectrum Fragments

Fragment	m/z
$\text{FP}(\text{C}_3\text{F}_7)_3^+$	557
$\text{F}_2\text{P}(\text{C}_3\text{F}_7)_2^+$	407
$\text{F}_3\text{P}(\text{C}_3\text{F}_7)^+$	257
F_4P^+	107
C_3F_7^+	169

High Resolution Mass Analysis

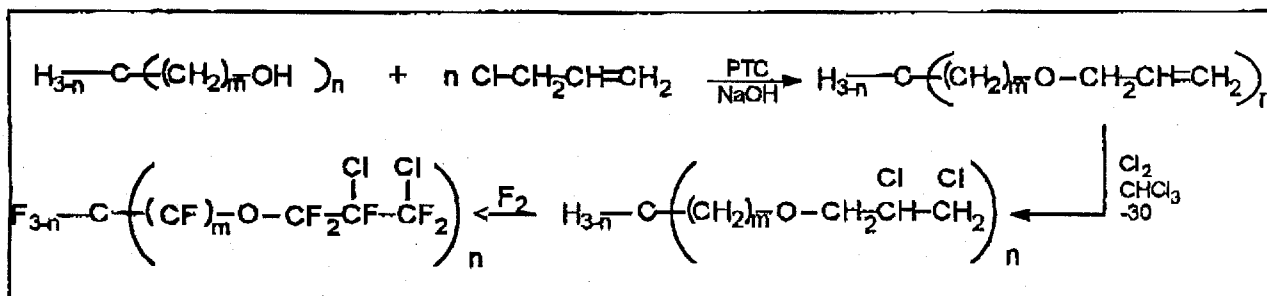
$\text{C}_9\text{F}_{22}\text{P}^+$ Calculated: 556.938635

Observed : 556.937623

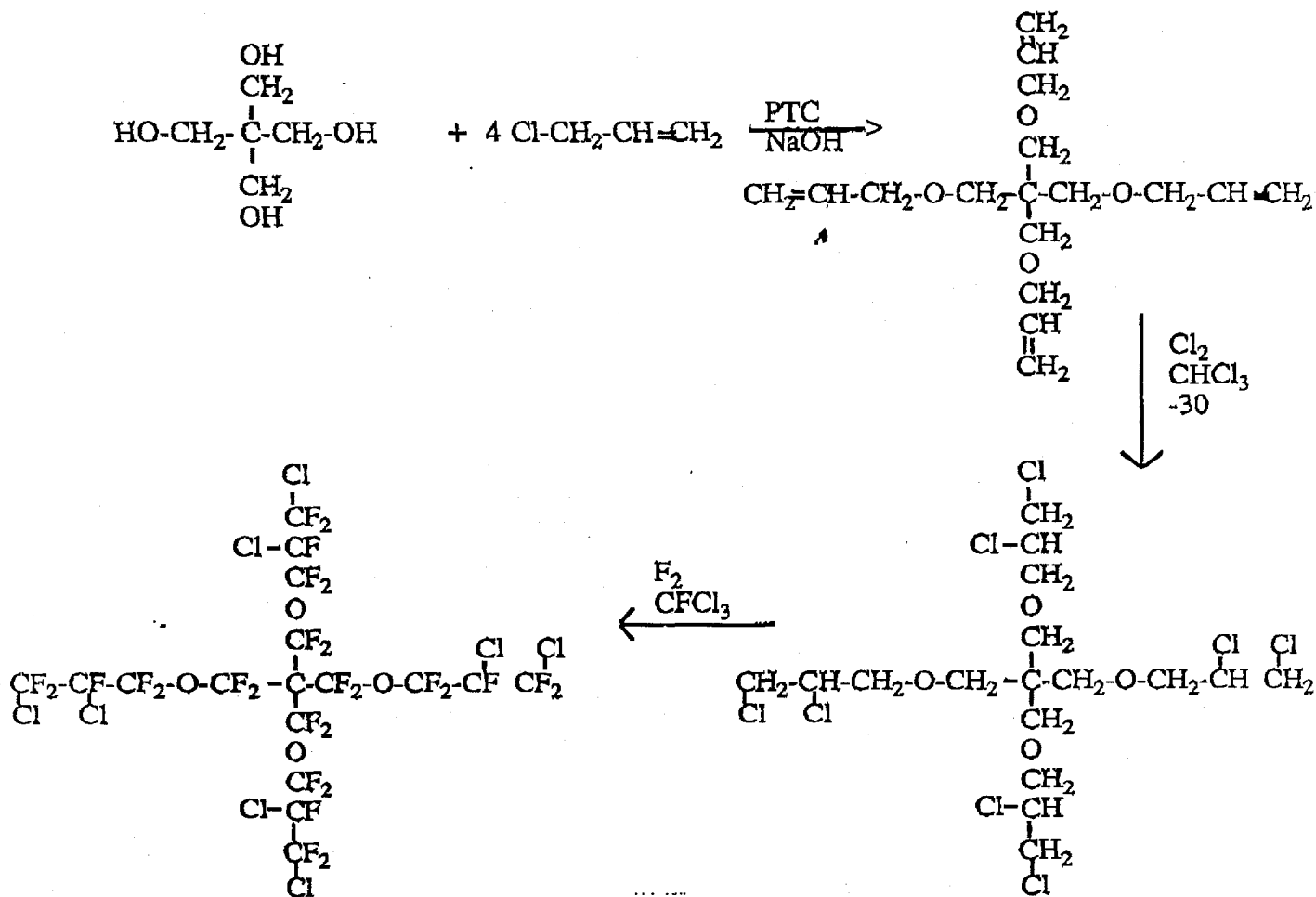
There are successes in many other areas to report but we lack the space here to do so. This has been a very successful year in our program.

APPENDIX I

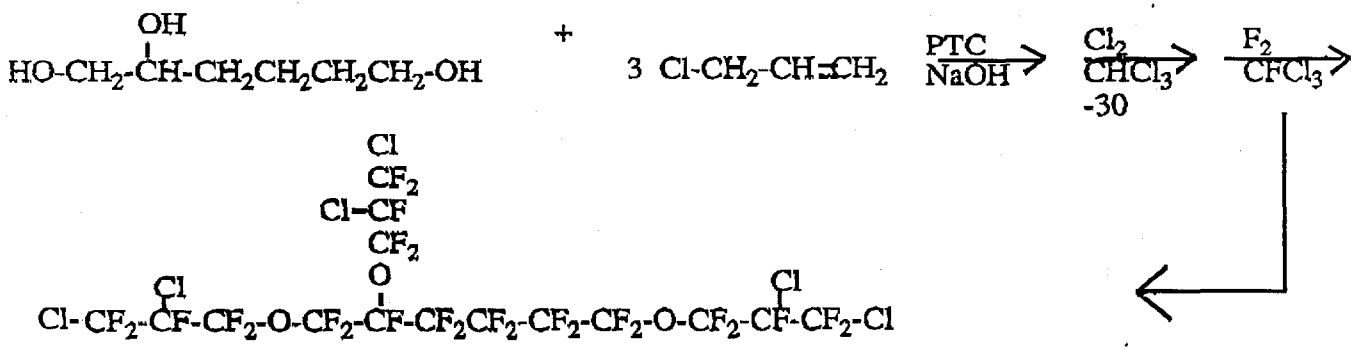
General Scheme



Representative examples:

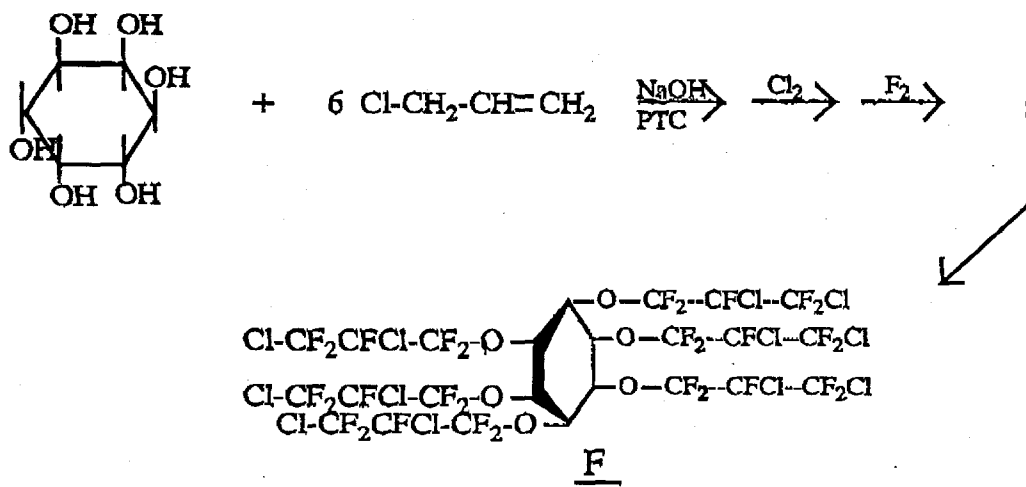
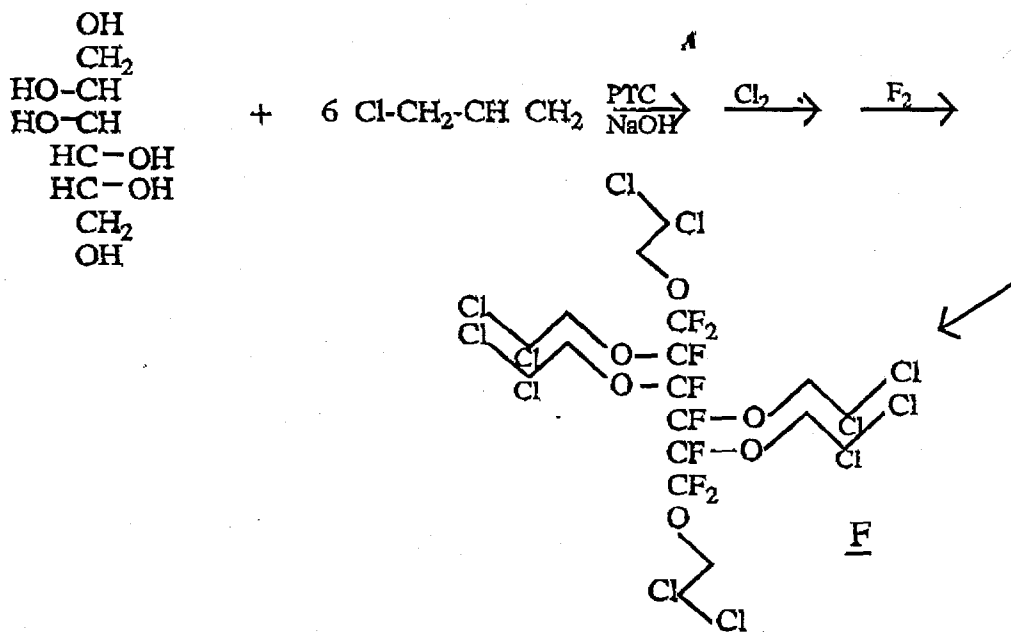


C.I. Mass Spec. Parent -F m/e = 1065



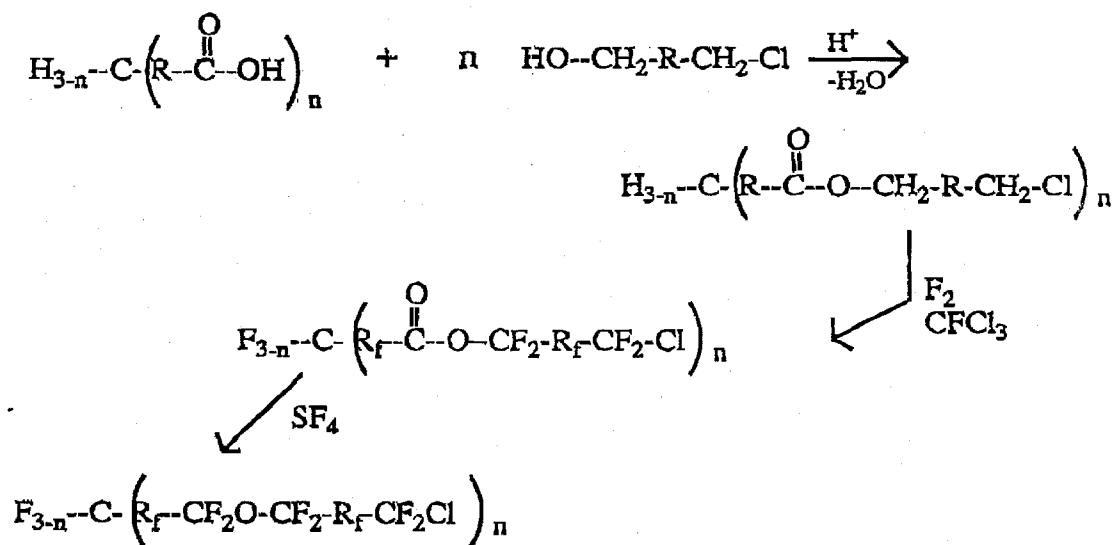
C.I. Mass. Spec. Parent -F m/e = 916

Variations in the polyhydrolic starting materials will provide a variety of chlorofluorocarbon ether structures. Synthesis of the following compounds is underway.

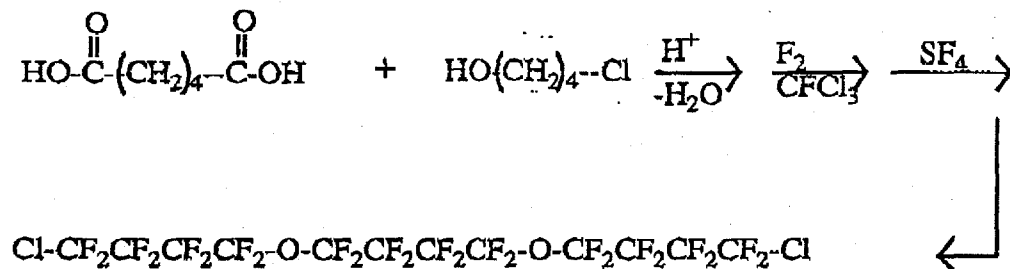


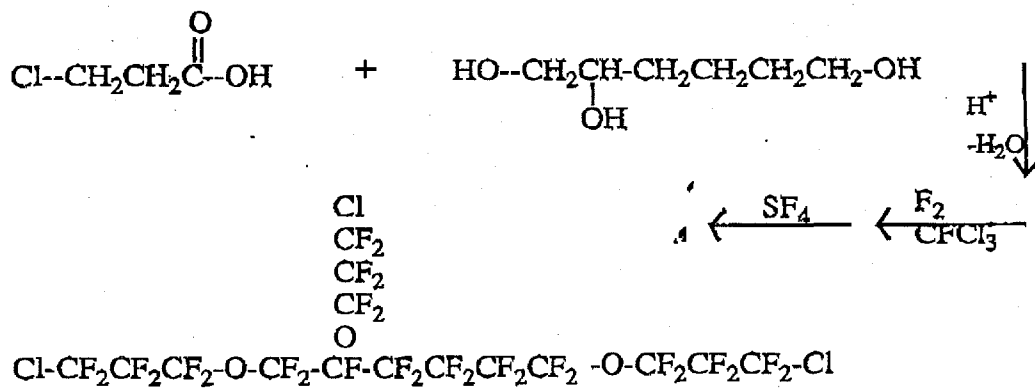
Approach II: Conversion of Perfluoroesters to Perfluoroethers using SF₄ is well known. Synthesis of chlorine containing fluoroesters followed by their conversion to ethers will provide for a variety of chlorinated fluoroethers, and will also provide a great deal of control over chlorine content and location. The synthesis of such molecules is underway.

General Scheme



Representative examples:





APPENDIX II

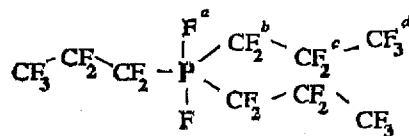
¹⁹F NMR Chemical Shifts

a - 47.5 ppm (d of mult.)

b -110.3 ppm (d of t)

c -125.7 (s)

d - 82.2 (s)

Coupling Constants¹J_{PF}=1041 Hz²J_{PCP}= 124 Hz³¹P NMR Chemical Shift

-41.8 ppm (t of sept.)

Low Resolution MassSpectrum Fragments

Fragment m/z

FP(C₃F₇)₃⁺ 557F₂P(C₃F₇)₂⁺ 407F₃P(C₃F₇)⁺ 257F₄P⁺ 107C₃F₇⁺ 169High Resolution Mass AnalysisC₉F₂₂P⁺ Calculated: 556.938635

Observed : 556.937623

¹⁹F NMR Chemical Shifts

a - 46.8 ppm (d of mult.)

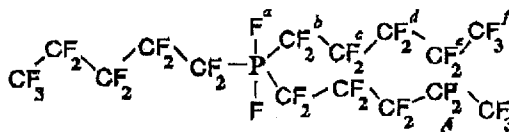
b -108.8 ppm (d)

c -120.7 (s)

d - 123.4 (s)

e -127.4 (s)

f - 82.5 (s)

Coupling Constants¹J_{PF}=1050 Hz²J_{PCP}= 126 Hz³¹P NMR Chemical Shift

-37.6 ppm (t of sept.)

Low Resolution MassSpectrum Fragments

Fragment m/z

FP(C₅F₁₁)₃⁺ 857F₂P(C₅F₁₁)₂⁺ 607F₃P(C₅F₁₁)⁺ 357F₄P⁺ 107C₅F₁₁⁺ 269High Resolution Mass AnalysisC₁₅F₃₄P⁺ Calculated: 856.919474

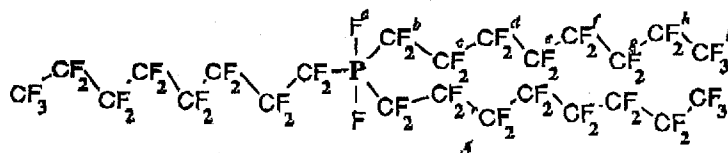
Observed : 856.918693

¹⁹F NMR Chemical Shifts

- a - 46.9 ppm (d of mult.)
- b -109.1 ppm (d)
- c -120.8 (s)
- d - 122.8 (s)
- e -123.1 (s)
- f -123.4 (s)
- g -124.3 (s)
- h -128.0 (s)
- i - 83.3 (s)

Coupling Constants

- ¹J_{PF}=1060 Hz
- ²J_{FCP}= 123 Hz



³¹P NMR Chemical Shift

-38.0 ppm (t of sept.)

Low Resolution Mass

Spectrum Fragments

Fragment	m/z
F ₂ P(C ₈ F ₁₇) ₃ ⁻	1326
FP(C ₈ F ₁₇) ₃ ⁺	1307
F ₂ P(C ₈ F ₁₇) ₂ ⁺	907
F ₃ P(C ₈ F ₁₇) ⁺	507
F ₄ P ⁺	107
C ₈ F ₁₇ ⁺	419

High Resolution Mass Analysis

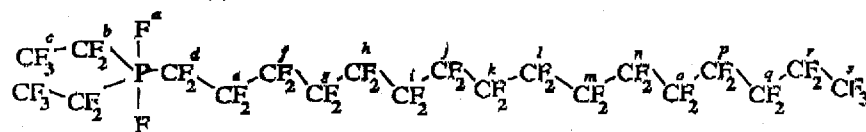
C₂₄F₅₃P⁻ Calculated: 1325.889136
Observed : 1325.888968

¹⁹F NMR Chemical Shifts

- a - 48.6 ppm (d of mult.)
- b -113.3 ppm (d)
- c - 81.5 (s)
- d - 117.2 (d)
- e-p -121.9 (s)
- q -123.0 (s)
- r -126.6 (s)
- s - 81.7 (s)

Coupling Constants

- ¹J_{PP}=1030 Hz
- ²J_{FCP}= 112 Hz
- ²J_{FCP}= 91 Hz



³¹P NMR Chemical Shift

-43.8 ppm (t of sept.)

Low Resolution Mass

Spectrum Fragments

Fragment	m/z
FP(C ₂ F ₅) ₂ (C ₁₆ F ₃₃) ⁺	1107
F ₂ P(C ₂ F ₅)(C ₁₆ F ₃₃) ⁺	1007
F ₂ P(C ₂ F ₅) ₂ ⁺	307
F ₃ P(C ₂ F ₅) ⁺	207
F ₃ P(C ₁₆ F ₃₃) ⁺	907
F ₄ P ⁺	107
C ₁₆ F ₃₃ ⁺	819
C ₂ F ₅ ⁺	119

High Resolution Mass Analysis

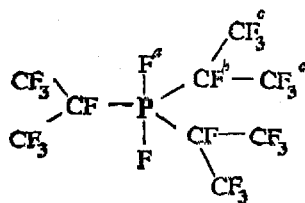
C₂₀F₄₄P⁺ Calculated: 1106.903506
Observed : 1106.906923

¹⁹F NMR Chemical Shifts

- a - 18.7 ppm (d of mult.)
- b -176.7 ppm (d)
- c - 71.0 (s)

Coupling Constants

- ¹J_{PF} = 1015 Hz
- ²J_{FCP} = 99 Hz



Low Resolution Mass

Spectrum Fragments

Fragment	m/z
FP(C ₃ F ₇) ₃ ⁺	557
F ₂ P(C ₃ F ₇) ₂ ⁺	407
F ₃ P(C ₃ F ₇) ⁺	257
F ₄ P ⁺	107
C ₃ F ₇ ⁺	169

³¹P NMR Chemical Shift

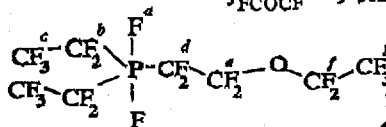
-29.0 ppm (t of q)

¹⁹F NMR Chemical Shifts

- a - 50.2 ppm (d of mult.)
- b -114.7 ppm (d)
- c - 83.1 (s)
- d - 114.1 (d)
- e - 84.9 (s)
- f - 90.0 (t)
- g - 88.9 (s)

Coupling Constants

- ¹J_{PF} = 1007 Hz
- ²J_{FCP} = 122 Hz
- ²J_{FCP} = 112 Hz
- ⁴J_{FCOCP} = 9 Hz



Low Resolution Mass

Spectrum Fragments

Fragment	m/z
F ₂ P(C ₂ F ₅) ₂ (C ₄ F ₉ O) ⁺	542
FP(C ₂ F ₅) ₂ (C ₄ F ₉ O) ⁺	523
F ₂ P(C ₂ F ₅)(C ₄ F ₉ O) ⁺	423
F ₂ P(C ₂ F ₅) ₂ ⁺	307
F ₃ P(C ₂ F ₅) ⁺	207
F ₃ P(C ₄ F ₉ O) ⁺	323
F ₄ P ⁺	107
C ₄ F ₉ O ⁺	235
C ₂ F ₅ ⁺	119

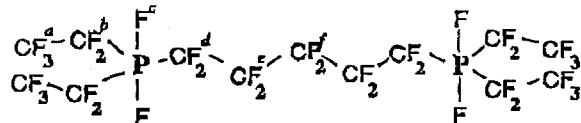
³¹P NMR Chemical Shift

-45.3 ppm (t of sept.)

High Resolution Mass Analysis

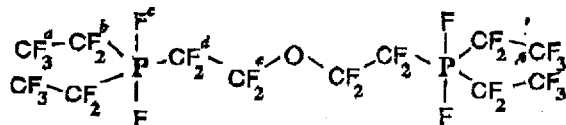
C₈F₂₀OP⁺ Calculated: 522.936743
Observed : 522.935822

¹⁹ F NMR Chemical Shifts	Coupling Constants
a - 83.6 ppm (s)	² J _{FCF} = 122 Hz
b - 114.9 ppm (d)	¹ J _{PP} = 1019 Hz
c - 49.8 (d of mult.)	² J _{FCP} = 125 Hz
d - 110.0 (d)	
e - 120.6 (s)	
f - 128.2 (s)	

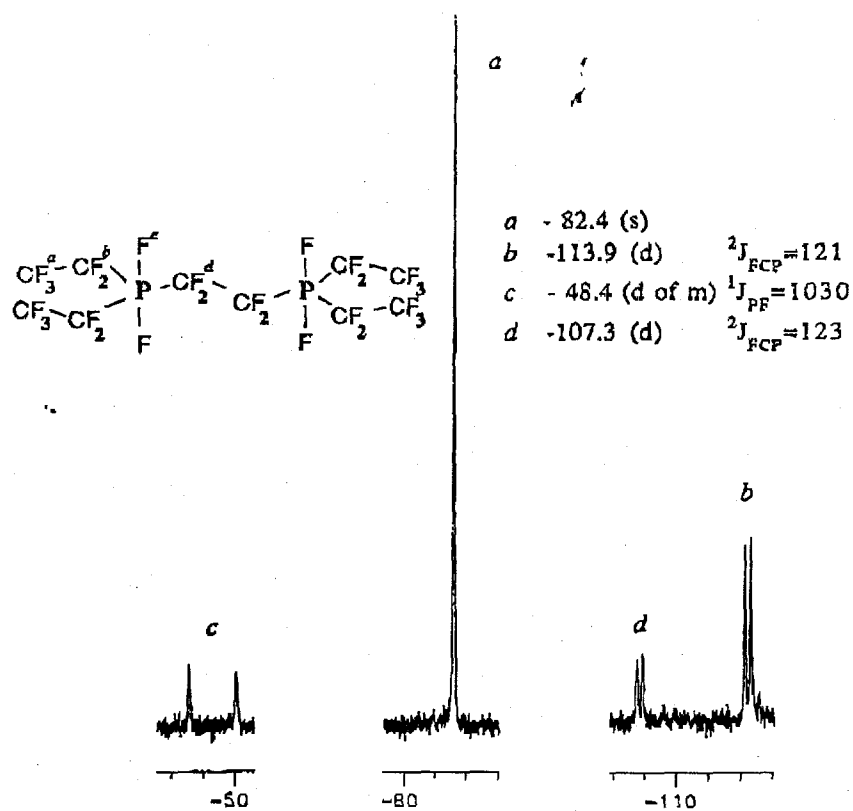


³¹ P NMR Chemical Shift	High Resolution Mass Analysis
-44.0 ppm (t of sept.)	C ₁₃ F ₃₃ P ₂ ⁺ Calculated: 844.894834 Observed : 844.894530

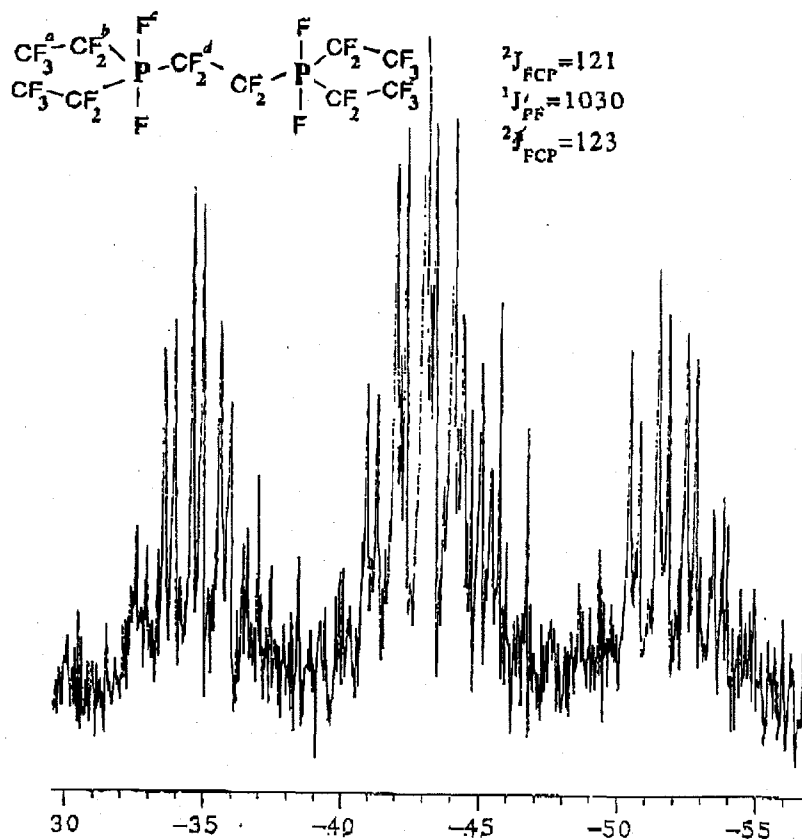
¹⁹ F NMR Chemical Shifts	Coupling Constants
a - 83.6 ppm (s)	² J _{FCP} = 122 Hz
b - 115.0 ppm (d)	¹ J _{PP} = 1008 Hz
c - 50.5 (d of mult.)	² J _{FCP} = 123 Hz
d - 114.0 (d)	
e - 84.3 (s)	



³¹ P NMR Chemical Shift	High Resolution Mass Analysis
-45.2 ppm (t of sept.)	C ₁₂ F ₃₂ OP ₂ ⁺ Calculated: 829.891345 Observed : 829.893040



^{19}F NMR of 1,2-bis(difluorobis(pentafluoroethyl)-
phosphorano)tetrafluoroethane



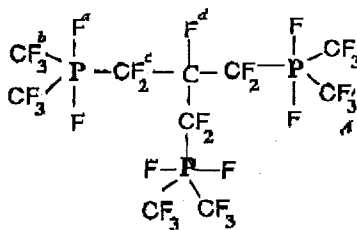
${}^{31}\text{P}$ NMR of 1,2-bis(difluorobis(pentafluoroethyl)-phosphorano)tetrafluoroethane

${}^{19}\text{F}$ NMR Chemical Shifts

- a - 55.2 ppm (d of mult.)
- b - 64.7 ppm (d)
- c - 109.7 (d)
- d - 177.0 (s)

Coupling Constants

- ${}^1J_{PF} = 1068$ Hz
- ${}^2J_{FCP} = 162$ Hz
- ${}^2J_{FCP} = 122$ Hz



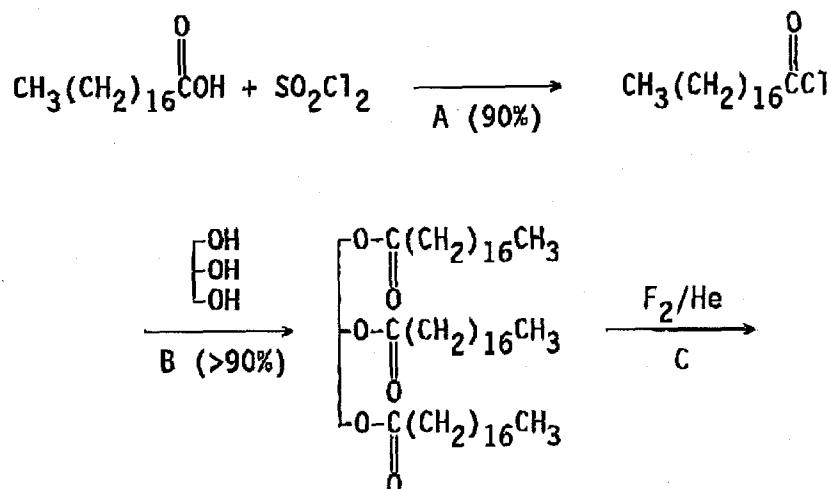
${}^{31}\text{P}$ NMR Chemical Shift

-49.5 ppm (t of mult.)

High Resolution Mass Analysis

$\text{C}_{10}\text{F}_{30}\text{P}_3$ Calculated: 782.873388
Observed: 782.869965

APPENDIX III

Preparation of perfluorinated triglyceride

Procedure A: Thionyl chloride (50 g, 0.42 mole) was added to a one-neck 500 ml flask. Stearic acid (100 g, 0.35 mole) was dissolved in 50 ml of CH_2Cl_2 . The solution was added to the thionyl chloride slowly. After complete addition, the mixture was refluxed for two hours. After pumping off unreacted SO_2Cl_2 , the residue was sublimated at 50 °C to get pure acid chloride in 90% yield.

Procedure B: 80 g of $\text{CH}_3(\text{CH}_2)_{16}\overset{\text{O}}{\parallel}\text{CCl}$ was added slowly to a mixture of 6.7 g glycerin and 80 ml pyridine at room temperature under an argon atmosphere. Two hours after addition of the acid chloride, the mixture was acidified by 10% of H_2SO_4 . After filtration, the precipitate was washed with saturated $\text{NaHCO}_3(\text{aq})$ until it is neutral. The solid triglyceride was dried at 80 °C under high vacuum for 12 hours. The yield is higher than 90%.