General Disclaimer

One or more of the Following Statements may affect this Document

- This document has been reproduced from the best copy furnished by the organizational source. It is being released in the interest of making available as much information as possible.
- This document may contain data, which exceeds the sheet parameters. It was furnished in this condition by the organizational source and is the best copy available.
- This document may contain tone-on-tone or color graphs, charts and/or pictures, which have been reproduced in black and white.
- This document is paginated as submitted by the original source.
- Portions of this document are not fully legible due to the historical nature of some
 of the material. However, it is the best reproduction available from the original
 submission.

Produced by the NASA Center for Aerospace Information (CASI)

AIAA

OK

NASA Technical Memorandum 82834

'NASA-TM-82834) THERMAL CXICATIVE JEGRADATION REACTIONS OF LINEAR PERFLUOROALKY LETHERS (NASA) 24 PHC A02/MF A01

N82-26468

CSCL 07C G3/27

Unclas 28101

Thermal Oxidative Degradation Reactions of Linear Perfluoroalkylethers

William R. Jones, Jr. Lewis Research Center Cleveland, Ohio

and

K. J. L. Paciorek, T. I. Ito, and R. H. Kratzer Ultrasystems, Inc. Irvine, California



Prepared for the Fluoropolymers Symposium sponsored by the American Chemical Society Las Vegas, Nevada, March 29-April 2, 1982



William R. Jones, Jr. Lewis Research Center Cleveland, Ohio

and

K. J. L. Paciorek, T. I. Ito, and R. H. Kratzer Ultrasystems, Inc. Irvine, California

SUMMARY

Thermal and thermal oxidative stability studies have been performed on linear perfluoroalkylether fluids. The effect on degradation by metal catalysts (M-50 steel and Ti(4 Al, 4 Mn) alloys) and degradation inhibitors (a monophospha-s-triazine and a perfluorophenyl phosphine) are reported.

The linear perfluoroalkylethers are inherently unstable at 316°C in an oxidizing atmosphere. This instability is not due to the presence of hydrogen chain termination or peroxide linkages. The metal catalysts greatly increased the rate of degradation in oxidizing atmospheres. In the presence of these metals in an oxidizing atmosphere, the degradation inhibitors were highly effective in arresting degradation at 288°C. However, the inhibitors had only limited effectiveness at 316°C. The metals promote degradation by chain scission.

Based on elemental analysis and oxygen consumption data, the linear perfluoroalkylether fluids have a structural arrangement based on difluoroformyl (-CF₂O-) and tetrafluoroethylene oxide (-CF₂CF₂O-) units, with the

former predominating.

INTRODUCTION

Perfluoroalkylethers are a class of fluids which exhibit excellent thermal and oxidative stability (refs. 1 and 2). Combined with good viscosity characteristics (ref. 3), good elastohydrodynamic film forming capabilities (ref. 4), good boundary lubricating ability (refs. 3 and 5), and nonflamability properties (ref. 6) make these fluids promising candidates for

high temperature lubricant applications.

Basically, there are two types of perfluoroalkylethers, a linear (ref. 7) and a branched class (ref. 1). The general structures of these classes are shown in figure 1. The most important representatives of the branched materials (fig. 1(a)) are based on the polymerization of hexafluor-opropylene oxide (HFPO). These compositions suffer some deficiencies. In order to satisfy low temperature fluidity, volatility problems are encountered (ref. 8). In addition, the poly(hexafluoropropene oxides) were found to exhibit poor compatibility with ferrous and titanium alloys above 260° C (ref. 3).

The mechanism responsible for the low temperature (<316°C) incompatibility has been studied by Paciorek et al. (ref. 9) and was found to be due to the presence of ~3 percent unstable chains. These chains have been subsequently shown to be hydrogen (rather than fluorine) terminated. Exposure of HFPO fluids at 343°C in oxygen resulted in removal of these chains by volatilization. The resultant fluid was not degraded by oxygen at 343°C

nor by M-50 and Ti(4 Al, 4 Mn) alloys at 316° C. However, degradation did occur with these alloys at 343° C.

A new class of perfluoroalkylethers based on the photo-oxidation of fluoro-olefins (ref. 7) has been developed. This class of materials, whose general chemical structure appears in figure 1(b), has a linear structure. These fluids have better viscosity-temperature properties than the branched (HFPO) class (ref. 8). However, the linear fluid class has exhibited lower thermal oxidative stability compared to the HFPO fluids (ref. 8). This is surprising since the chemical bonding in both classes is very similar. In fact, it has been shown (ref. 10) that tertiary carbon-fluorine bonds are less stable than those involving reimary or secondary carbon atoms. This would lead one to conclude that the HFPO fluids (which contain tertiary carbon atoms) should be less stable than the linear fluids.

Therefore, the objective of this work was to conduct a study of the thermal oxidative degradation processes of the linear class of perfluoro-alkylether fluids. Tests were conducted with the neat fluid and also in the presence of M-50 or Ti(4 Al, 4 Mn) alloys. Test atmospheres included air, oxygen, and nitrogen. Test temperatures ranged from 288°C to 343°C. Selected tests were conducted in the presence of a degradation inhibitor (a phospha-s -triazine or a perfluorophenyl-phosphine).

EXPERIMENTAL FLUIDS

Two types of perfluoroalkylether fluids were employed in this study. One class is believed to have a linear structure and is manufactured by Montecatini Edison under the trade name Fomblin Z. Two different batches of this class were studied which are designated MLO-72-22 and MLO-79-129. The second class has a branched structure and is manufactured by E. I. Dupont de Nemours and Co. under the trade name Krytox (MLO-71-6). These are Air Force designations where the first two numbers refer to the year the fluid was received.

DEGRADATION INHIBITORS

Two different degradation inhibitors were used in these studies. One was a perfluoroalkylether substituted perfluorophenyl phosphine (fig. 2(a)). Results with this additive have been previously reported (ref. 11) where it was designated as P-3. The second additive was a perfluoroalkylether substituted monophospha-s-triazine (fig. 2(b)) (ref. 12). This additive will be designated as C_2PN_3 for convenience.

OXIDATION - CORROSION APPARATUS

The micro-oxidation corrosion apparatus is a modified version of the type reported by Snyder and Dolle (ref. 3). The decomposition tube configuration and the rod assembly for holding the metal corrosion catalysts are schematically shown in figures 3 and 4, respectively. The metal catalysts (M-50 steel or Ti(4 Al, 4 Mn) alloys) were 9.5 mm (3/8 in.) OD and 3.2 mm (1/8 in.) ID. These were obtained from Metaspec Co., San Antonio, Texas. For heating sample tubes in a vertical position, a modified Lindberg Heavy-Duty box furnace, Type 51232 (fig. 5) was used. In this arrangement, 180 mm of the 420 mm of tube length were at test temperature; the fluid occupied, at most, the lower 75 mm (fig. 3). The extra gas reservoir was in the ambient environment.

Prior to testing the metal catalysts were polished using first, Norton No-Fil Durite finishing paper Type 4 220A. This was followed by open coat Silicon Carbide papers grades 400A and 500A, respectively. Subsequently, the catalysts were washed with Freon 113, dried, weighed, and suspended in the test apparatus (see fig. 4). After the completion of a given experiment, the metal catalysts were washed with Freon 113, dried in an inert atmosphere chamber, then weighed and visually inspected. The used catalysts were subsequently sealed in Mylar envelopes.

In a typical experiment, the fluid was introduced into the decomposition tube (see fig. 2) which was then evacuated and filled to a known pressure at a known temperature with a selected gas (air, N2, or O2). Inasmuch as the apparatus was calibrated and the fluid volume measured accurately, the quantity of gas thus introduced was exactly known. The decomposition tube was then inserted into the preheated box furnace and kept there for a specified period of time; throughout this exposure the temperature was continuously recorded. After removal from the furnace, the tube was allowed to cool to room temperature, attached to the vacuum line, and opened. The liquid nitrogen noncondensibles were collected quantitatively, measured, and analyzed by gas chromatography and infrared spectroscopy. The liquid nitrogen condensibles, which were volatile at room temperature, were measured, weighed, and analyzed by infrared spectroscopy and mass spectrometry. The fluid residue itself was weighed and subjected to infrared spectral analysis; in selected instances, molecular weight and nuclear magnetic resonance (NMR) determinations were performed.

The degradation rate is calculated from the amount of liquid nitrogen condensibles formed and is reported as milligrams of condensible product per gram of original fluid per hour.

THERMAL DECOMPOSITION APPARATUS (TENSIMETER)

The tensimeter apparatus is shown in figure 6. The basic function of the tensimeter consists of heating a liquid sample and of plotting either the vapor pressure or the isothermal time rate of increase of vapor pressure due to thermal decomposition as a function of temperature.

The sample cell is a glass bulb having a 5-milliliter $(5x10^{-6} \text{ m}^3)$ volume with a stem extending through the oven wall to a valve and pressure transducer mounted outside the oven. Three to four milliliters $(3x10^{-6} \text{ to } 4x10^{-6} \text{ m}^3)$ of test fluid are placed in the sample cell. The cell is attached to the cell assembly and the fluid is degassed and refluxed under vacuum. The cell assembly is then placed in the oven. The sample is heated to an initial temperature about 50°C below the suspected decomposition temperature. After a 5-minute stabilization period, the increase in vapor pressure, if any, is recorded as a vertical bar during a fixed time interval. Then the programmer raises the sample temperature by a preset amount, usually 5°C, and the previous process is repeated.

This yields a plot of the logarithm of the isothermal rate of vapor pressure increase as a function of reciprocal absolute temperature. A straight line is drawn connecting the tops of the recorded bars at higher temperatures. The intersection with the temperature reference axis is the thermal decomposition temperature (T_d) . The temperature axis corresponds to a pressure rise of 50 torr per hour, which defines the T_d .

RESULTS AND DISCUSSION

This investigation involves mainly the linear class of perfluo kylether fluids. This class of materials is prepared from tetrafluorouchylene and oxygen uncer ultraviolet radiation (ref. 7). This process introduces peroxide linkages which are then removed by subsequent heat treatment.

Some limited data is presented for the branched class of fluids. These are prepared by telomerization of hexafluoropropylene oxide (HFPO) (ref. 10). The termination reaction is well documented (ref. 10) and theoretically should yield only perfluoroalkyl end groups. No such information is available for the linear fluids, although based on published data (ref. 13) it can be concluded that the grouping is -CF2H. The nature of these end-groups and the presence of structural irregularities are factors which limit the inherent stability of the periluoroalkylether systems. Thus, any study of thermal oxidative behavior must take into account structural considerations.

DEGRADATION STUDIES

Thermal and thermal oxidative studies performed on the two batches of linear perfluoroalkylethers are summarized in tables I (MLO-72-22) and II (MLO-79-196). The effect of metal catalysts and degradation inhibitors on fluid M $_2$ O-72-22 appear in tables III and IV, respectively. The distribution of condensible volatiles appears in table V.

Nonoxidizing atmospheres. - The presence of heat "removable" species was determined in tests 1 (MLO-72-22) and 2, 3, and 7 (MLO-79-196) which were performed in nitrogen atmospheres. Tests 1 and 2 were run under identical test conditions (3 hr at 150°C, 2 hr at 200°C, and finally 2 hr at 250°C). The total condensible products for test 2 was about 50 times that of test 1. This indicates that batch MLO-72-22 is thermally more stable than MLO-79-196. However, further heat treatment at 250°C of the MLO-79-196 residue from test 2 yields a much lower degradation rate (test 3). One can conclude that some unstable species were removed during the initial heat treatment. These data are illustrated in figure 7.

Thermal tests on MLO-79-196 using another device (the tensimeter) were performed. A typical result is shown in figure 8. Early evolution of volatiles is evident well below the thermal decomposition temperature (T_d) of 361°C. A rerun of the residue from this test appears in figure 9. There is less lower temperature decomposition and the T_d is elevated (365°C). This supports the prior statement that some unstable species are removed after exposure at high temperatures in the absence of oxygen.

In contrast, the thermal decomposition data (tensimeter) for a HFPO fluid (Krytox PR 143AB) appears in figure 10. There is almost no low temperature decomposition and the T_d is almost 30°C higher than the linear fluid.

Oxidizing atmospheres. - In oxidizing atmospheres at 316°C, initial exposure of the MLO-72-22 fluid (test 5) promoted considerable degradation. However, subsequent exposures of the residue from test 5 at 316°C, results in less degradation (tests 6, 9, and 10). This is illustrated in figure 11. These rates are more than an order of magnitude greater than a comparable test of the MLO-79-196 fluid in nitrogen (test 7).

At higher temperatures in oxygen, the degradation rate increases about 1.5 times for each 10° C rise in temperature. This is illustrated in figure 12. A similar temperature rate increase (1.3 times per 10° C rise) occurred in nitrogen (tests 3 and 7).

It became evident that sample size was affecting the degradation rate in some of the tests. For example, tests 17 and 19 run with less than 3 grams of fluid yielded rates of approximately 2.6 to 2.7 mg/g-hr. In contrast, tests 21, 30, and 39 run with large samples (>19 g) yielded considerably lower rates (0.72 to 0.90 mg/g-hr). A plot of rate of condensible product formation for MLO-72-22 as a function of sample size appears in figure 13. The reduction in rate at large sample sizes is not due to oxygen depletion since calculations show that only 5 percent of the available oxygen was consumed in test 21 and 12 percent in test 39. One explanation could be that the degradation is diffusion limited for large sample sizes. This effect is not a factor for small samples (~3g) since halving the fluid quantity (tests 17 and 19) did not alter the rate.

The residue from large sample test 21 was used for small sample tests 31 and 40. Both of these tests yielded rates which appear anomalously high compared to figure 12. It seems likely that the pretreatment (test 21) was not complete. As a point of interest, previous tests on HFPO fluids in the absence of metal catalysts indicated that sample size did not affect degra-

dation rates (ref. 9).

The thermal oxidative degradation behavior of MLO-79-196 fluid (table II) was similar to that of MLO-72-22 (table I). The degradation rate of MLO-79-196 as a function of temperature appears in figure 14. A reference curve for MLO-72-22 (from fig. 12) is shown for comparison. The rates of degradation for the two batches are very similar. The nature and relative concentration of the condensible volatiles (table V) formed by these two fluids are also similar.

Metal Catalysis. - Only fluid MLO-72-22 was evaluated in the presence of metal catalysts and degradation inhibiting additives. The test series involving the metal catalysts M-50 steel and Ti(4 Al, 4 Mn) alloys is compiled in table III.

The presence of the metal catalysts greatly accelerated fluid degradation in O_2 . At 288°C, M-50 and Ti(4 Al, 4 Mn) alloys increased the degradation rate by factors of 28 and 73, respectively. A test with M-50 at 316°C yielded a rate about 30 times that of the uncatalyzed fluid. However, in the absence of oxygen, neither catalyst promoted fluid degradation

(tests 43, 44, and 45). These data are illustrated in figure 15.

Degradation Inhibitors. – The effect of inhibitors on the degradation rate are compiled in table IV. Both a monophospha-s-triazine (C_2PN_3) and a perfluorophenyl phosphine (P-3) were studied at a concentration of I weight percent. Both additives were effective in reducing the degradation processes in the presence of metal catalysts. At 288° C, rates were reduced by factors of 700 to 2000. At 316° C the inhibitors were still effective but to a much lesser degree (less than an order of magnitude). It appears that on a weight basis the C_2PN_3 additive is about twice as effective as the P-3 additive. The data for M-50 are illustrated in figure 16.

Earlier work (ref. 9) on HFPO fluids showed that inhibitors also arrested fluid degradation in the absence of metal catalysis. A similar result was obtained for a linear fluid (test 33) where the degradation rate

was reduced by a factor of 8.

Nuclear magnetic resonance (NMR) measurements. — The 19 F NMR spectra of the fluid residue from test 19 appears in figure 17. Expanded sections of figure 17 labeled A, B, C, and D appear in figure 18. Proton NMR spectra for the residues from tests 19 and 21 appear in figure 19. The CCl₃F reference peak and the peaks associated with the solvent, C_6F_6 , are identified in figure 17.

The fluid appears to contain CF30-, -0CF2, -0CF2CF2-, and -0CF2CF2CF2CF2-groups arranged randomly. Area "A" may be assigned to CF30-, -0CF20-CF20-, -0CF20CF2CF2, etc.; areas "B" and "C" may be assigned to -0CF2CF20-, -0CF2CF2CF2- groups depending on what is attached to the oxygen, e.g., -0CF2CF20CF20 is different from -0CF2CF20CF2CF20-. Area "D" is due to a -CF2-surrounded by other CF2 groups, e.g., -0CF2CF2CF20-.

No peaks were found in the 0 to +100 ppm region (not shown) indicating an absence of -COF groups. The spectra are different from those reported by Sianesi, et al. (ref. 13). Significantly, there are no doublets with 50-70 Hz splitting characteristic of -OCF₂H or -OCF₄H groups, etc. In other words, the ¹⁹F NMR is consistent with the ¹H NMR (fig. 19) indi-

cating the absence of protons.

The absence of protons in the pretreated fluid is in agreement with the experimental data (i.e., the lack of stabilization after exposure to oxidizing atmospheres for prolonged periods of time). The experimental degradation data thus indicates that either all the chains are hydrogen-terminated, which was not revealed by proton NMR, or such termination is insignificantly low and is not responsible for the observed thermal oxidative behavior of the fluids. It is unlikely that residual peroxide linkages are causing the observed thermal oxidative instability since the rate of degradation failed to decrease after prolonged exposure at elevated temperatures. Metals, in the absence of oxygen, did not cause any fluid degradation, supporting further that there are no residual peroxide linkages. Consequently, it must be concluded that the arrangement of the units in the chains or the end-groups is responsible for the observed degradation behavior.

Elemental analysis. - Elemental analyses of the fluid, both "lightly" thermally stressed (test 21) and after extensive exposure to oxygen at elevated temperature (test 19), differ somewhat as evident from the tabulation given in table VI. On the other hand, the variation in carbon content is within acceptable limits and it has been usually found that in these type of compositions, fluorine values are generally lower than those actually present. Thus, it is unknown whether the calculated increased oxygen content in the residue of test 19 is real. However, the C:O ratio which is lower for both of the samples than the theoretical 2:1 value for (CF2CF2O)x polymer is definitely real. Consequently, it would appear from these data that the ratio of CF2O entities to CF2CF2O groups greater than 1:1. This would further imply that the concentration of $-O(CF_2)_XOunits$ wherein x is greater than 2 is low. All these stipulations do not contradict the NMR spectral findings; unfortunately the latter do not provide information regarding the concentration of specific groups, i.e., $-OCF_2O_+$, $-O(CF_2)_2O_-$, $-O(CF_2)_4O_-$, etc.

concentration of specific groups, i.e., $-0CF_2O_-$, $-0(CF_2)_2O_-$, $-0(CF_2)_4O_-$, etc. From the calculations of the ratios of fluid volatilized to oxygen atoms consumed (given in table VII) for the different segments, it appears that the content of $-0CF_2O_-$ units is higher than that denoted by $-CF_2OCF_2CF_2O_-$ arrangement. Tests 16, 19, 21, and 31 were carried out in the absence of metal catalysts. In these tests only traces of materials non-volatile at -78° C were formed. Based on the nature of the collected volatiles, it can be deduced that COF_2 was the initial product formed. SiF4, CO_2 , and BF3 are the products of COF_2 interactions with constituents of glass at the elevated temperatures. Since some COF_2 survived the treatment, one would expect to see CF_3COF or related acid fluorides if there was substantial content of $-O(CF_2)_XO_-$ units where x is greater than 2. It should be noted that CF_3COF was found in the volatiles formed by HFPO derived polyethers (ref. 9), but none was detected in the linear fluid investigations.

Metals definitely promoted scission of the chain bonds as shown by the increased ratio of the volatilized fluid to oxygen consumed and by the type of products formed. This is especially significant in the case of Ti(4 Al, 4 Mn) alloy wherein at 288°C almost half of the products were in the form of relatively nonvolatile, predominantly -COF terminated chains as determined by infrared spectral analysis.

SUMMARY OF RESULTS

Based on thermal-oxidative degradation studies of linear perfluoroalkylether fluids the following results were obtained.

- 1. Linear perfluoroalkylether fluids are inherently unstable at 316°C in oxidizing atmospheres. This instability is not due to hydrogen chain termination or residual peroxide linkages.
- 2. In the presence of M-50 steel or Ti(4 Al, 4 Mn) catalysts at 316°C in oxidizing atmospheres, the linear perfluoroalkylether fluids exhibit much greater degradation than in uncatalyzed tests. These catalysts do not promote degradation at 316°C in inert (nitrogen) atmospheres.
- 3. Two inhibitors, a monophospha-s-triazine and a perfluorophenyl phosphine were highly effective in arresting degradation at 288° C, but had only limited effectiveness at 316° C.
 - 4. The metal catalysts promote degradation by chain scission.
- 5. Based on elemental analysis and oxygen consumption data, the linear perfluoroalkylether fluids have a structural arrangement based on difluoroformyl (-CF₂O-) and tetrafluoroethylene oxide (-CF₂CF₂O-) units with the former predominating.

ACKNOWLEDGEMENT

The authors would like to acknowledge the assistance of Mr. C. E. Snyder c: the Air Force Wright Aeronautical Laboratories, Fluids, Lubricants, and Elastomers Branch. We would also like to express our indebtedness to Dr. Paul R. Resnick of E. I. Dupont de Nemours and Company for the measurement and interpretation of the NMR spectra.

REFERENCES

- 1. Gumprecht, W. H.: "PR-143-A New Class of High Temperature Fluids," ASLE Trans. 9, p. 24 (1966).
- 2. Sianesi, D., Zamboni, V., Fontanelli, R. and Binaghi, M.: "Perfluoro-polyethers: Their Physical Properties and Behavior at High and Low Temperatures," Wear, 18, p. 85 (1971).
- 3. Snyder, C. E., Jr. and Dolle, R. E., Jr.: "Development of Polyper-fluoroalkylethers as High Temperature Lubricants and Hydraulic Fluids," ASLE Trans. 19, p. 171, (1976).

- 4. Jones, W. R., Jr., Johnson, R. L., Winer, W. O. and Sanborn, D. M.: "Pressure-Viscosity Measurements for Several Lubricants to 5.5x108 Newtons Per Square Meter (8x104 psi) and 149° C (300° F)," ASLE Trans, 18, 4, p. 249 (1975).
- 5. Jones, W. R. Jr. and Snyder, C. E., Jr.: "Boundary Lubrication, Thermal and Oxidative Stability of a Fluorinated Polyether and a Perfluoropolyether Triazine," ASLE Trans. 23, 3, p. 253 (1980).
- 6. Snyder, C. E. Jr., Gschwender, L. J. and Campbell, W. B.: Development and Mechanical Evaluation of Nonflammable Aerospace (-54° C to 135° C) Hydraulic Fluids, Lubr. Engr. 38, 1, p. 41 (1982).
- 7. Sianesi, D. et al., "Perfluoropolyethers by Photo-Oxidation of Fluoro-olefins," Chemica e l' Industria (Milan), 15, p. 208 (1973).
- 8. Snyder, C. E. Jr., Gschwender, L. J. and Tamborski, C.: "Linear Polyperfluoroalkylether Based Wide-Liquid-Range High-Temperature Fluids and Lubricants," Lubr. Engr. 37, 6, p. 344 (1981).
- 9. Paciorek, K. J. L., Kratzer, R. H., Kaufman, J. and Nakahara, J. H.: "Thermal Oxidative Studies of Poly (Hexafluoropropene Oxide) Fluids," J. of Applied Polymer Sci, 24, p. 1397 (1979).
- 10. Gumprecht, W. H.: "The Preparation and Thermal Behavior of Hexafluoro-propylene Epoxide Polymers," Fourth International Symp. on Fluorine Chemistry, Amer. Chem. Soc. of Fluorine Chem, Estes Park, Colorado (July 1967).
- 11. Snyder, C. E. Jr., Tamborski, C., Gopal, H. and Svisco, C. A.: "Synthesis and Development of Improved High-Temperature Additives for Perfluoroalkylether Lubricants and Hydraulic Fluid, Lubr. Engr. 35, 8, p. 451 (1979).
- 12. Kratzer, R. H., Paciorek, K. J. L., Kaufman, J., and Ito, T. I.: "Fhospha-s-Triazines. I. Syntheses and Properties of Mono (Diarylphospha)-s-Triazines," J. of Fluorine Chem., 10, p. 231 (1977).
- 13. Sianesi, D., Pasetti, A., and Belardinelli, G.: U.S. Patent 3, 715, 378, Feb. 6, 1973.

TABLE I. - THERMAL AND THERMAL OXIDATIVE BEHAVIOR OF LINEAR PERFLUOROALKYLETHER (MLO 72-22)

Test	Sample size, g	Atmosphere	Temperature, °C	Degradation rate, mg/g-hr
1 5 6a 9a 10a 17a 17a 21b 30 39 31c 40c	26.33 3.44 3.26 3.16 3.07 3.02 2.95 1.76 19.51 24.15 54.54 4.27 3.74	N ₂ Air O ₂	316 316 332 343 316 288	0.005 2.0 .93 1.2 .81 1.1 2.7 2.6 .90 .75 .72 2.7

^aInvolatile residue from previous test. ^bInvolatile residue from test 1. ^cCombined residue from tests 21 and 30.

TABLE II. - THERMAL AND THERMAL OXIDATIVE BEHAVIOR OF LINEAR PERFLUOROALKYLETHER (MLO 79-196)

Test	Sample size, 9	Atmosphere	Temperature, °C	Degradation rate, mg/g-hr
2 3a 7a 8a 11a 13a 14a 15a	35.29 32.71 5.03 5.03 3.90 3.82 3.77 3.71 3.46	N ₂ Air Air O ₂	150,200,250 250 288 316 288 316 316 316 316 332 332	0.24 .003 .017 .027 .013 .67 .66 .70 1.39 1.63

^aInvolatile residue from previous test.

TABLE III. - EFFECT OF METAL CATALYSTS ON THE THERMAL AND THERMAL OXIDATIVE BEHAVIOR OF LINEAR PERFLUOROALKYLETHER (MLO 72-22)

Testa	Atmosphere	Atmosphere Temperature, Catalyst		Degradation rate, mg/g-hr
40	0 ₂	288	None	1.0
38	0 ₂	288	M-50	28
41	0 ₂	288	Ti(4 Al, 4 Mn)	73
31	0 ₂	316	None	2.7
32	0 ₂	316	M-50	80
43	N 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	316	None	0.093
44		316	M-50	.095
45		316	Ti(4 Al, 4 Mn)	.099

 $^{^{\}mathrm{a}}$ All fluids were pretreated at 343° C in $^{\mathrm{o}}$ C for 24 hours.

TABLE IV. - EFFECT OF INHIBITORS ON THE THERMAL OXIDATIVE BEHAVIOR OF LINEAR PERFLUOROALKYLETHER (MLO 72-22)

IN AN OXYGEN ATMOSPHERE

Testa	⁷ emperature, °C	Catalyst	Inhibitor 1% wt	Degradation rate, mg/g-hr
36	288	M-50	C2PN3 b	0.014
37	288	M-50	P-3C	.040
42	288	Ti(4 Al, 4 Mn)	P-3	.047
33	316	None	P-3	0.12
34	316	M-50	P-3	12
35	316	M-50	C ₂ PN ₃	6.6

 $[^]a All\ fluids$ were pretreated at 343° C in 0_2 for 24 hours. $^b Monophospha-s-triazine. <math display="inline">^c Perfluorophenyl\ phosphine.$

TABLE V. - PRODUCT DISTRIBUTION OF THERMAL OXIDATIVE DEGRADATION OF LINEAR PERFLUOROALKYLETHERS

Test	Fluid	Fluid Volatile products, p			percent	
		CO2	COF2	S1F4	BF 3	
5 6 9 10 12 17 19 21 31 32a 33b 34c 38a 40 41d	MLO 72-22	65 60 63 60 65 55 66 61 47 49 67 70	3.1 3.7 2.2 2.0 2.2 4 1.5 6.6 19.7 13.6 13.9 14.9 23.3	22 26 28 22 36 34 27 25 21 52 30 13 9	9.5 9.8 9.9 10.7 9.6 11.0 6.0 2.5 9.8 4.0	
13 14 15 16	MLO 79-196	68 73 53 46	5.2 4.0 .9 1.3	16 11 41 46	10.6 12.0 5.0 6.8	

a_M-50 catalyst. bp-3 inhibitor. c_M-50 ard P-3. d_{Ti}(4 Al, 4 Mn).

TABLE VI. - ELEMENTAL ANALYSIS OF PERFLUOROALKYL ETHERS

: 62

Material	С,%	F,%	C:F:0			
MLO 72-22 (test 21)	20.03	61.21	2.85:5.50:2			
MLO 72-22 (test 19)	19.63	60.09	3.86:7.48:3			
(CF ₂ CF ₂ O) _x	20.69	65.52	2:4:1			
(CF ₂ OCF ₂ OCF ₂ CF ₂ O) _x	19.37	61.29	4:8:3			
(CF ₂ OCF ₂ CF ₂ O) _x	19.80	62.64	3:6:2			

TABLE VII. - INFLUENCE OF METAL CATALYSTS ON RATIOS OF VOLATILIZED FLUID TO OXYGEN CONSUMED

Test	Atmosphere	Temperature,	Catalyst	Products,		fluid lost is consumed
					Segm,116ª	Segm,182 ^b
16 19 21 31 32 38 41	02	332 343 343 316 316 288 288	None M-50 M-50 Ti(4 Al, 4 Mn)	272 216 426 279 (4) ^C 2666 (10) 1117 (22) 2235 (47)	2.1:1 2.6:1 2.1:1 2.0:1 2.6:1 3.7:1 4.6:1	1.3:1 1.5:1 1.3:1 1.2:1 1.7:1 2.3:1 2.9:1

aSegment -CF₂CF₂O-.
bSegment -CF₂OCF₂CF₂O-.
cValues in brackets correspond to the percentage of products which were involatile at -78°C.

(a) Hexafluoropropylene oxide (HFPO) based fluids.

 ${\rm R}_f {\rm O(CF_2CF_2O)_m (CF_2O)_n R_f}$ (b) Linear perfluoroalkylether fluids.

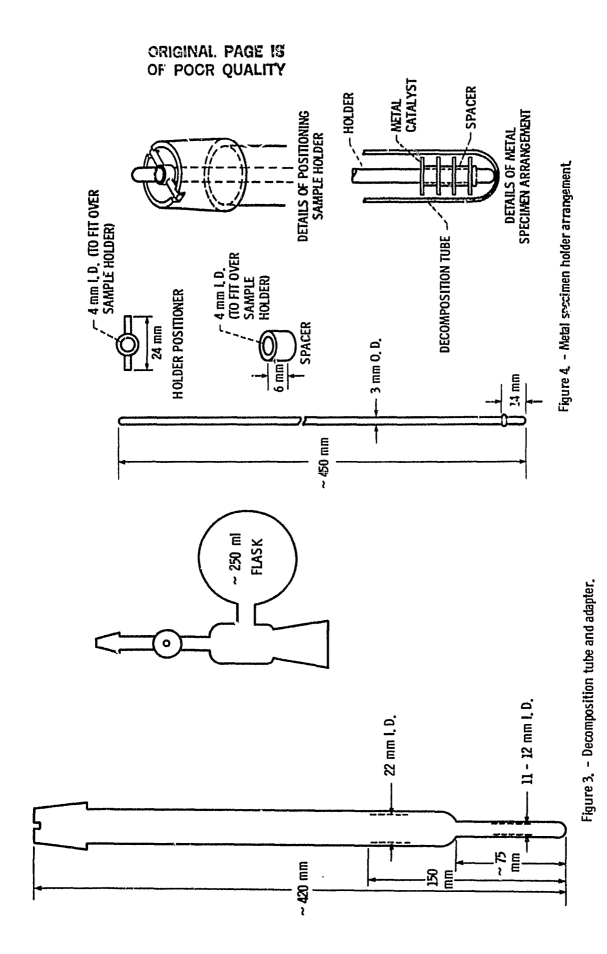
Figure 1. - Chemical structures of perfluoroalkylether fiulds ($R_f = CF_3$ or C_2F_5).

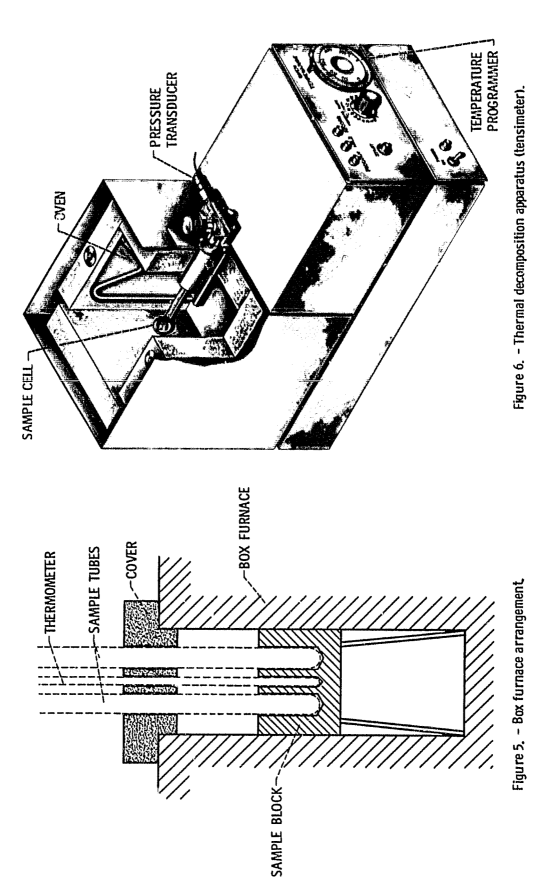
$$[\mathsf{C}_3\mathsf{F}_7\mathsf{OCF}(\mathsf{CF}_3)\mathsf{CF}_2\mathsf{OCF}(\mathsf{CF}_3)\mathsf{CF}_2 - \underbrace{ \left(\mathsf{F} \right) }_{=}]_3\mathsf{P}$$

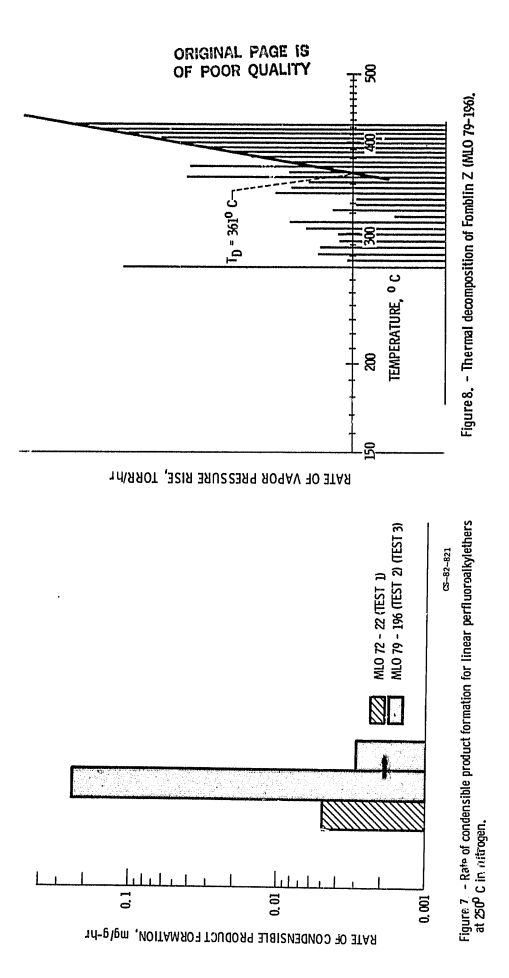
(a) Perfluoroalkylether perfluorophenyl phosphine,

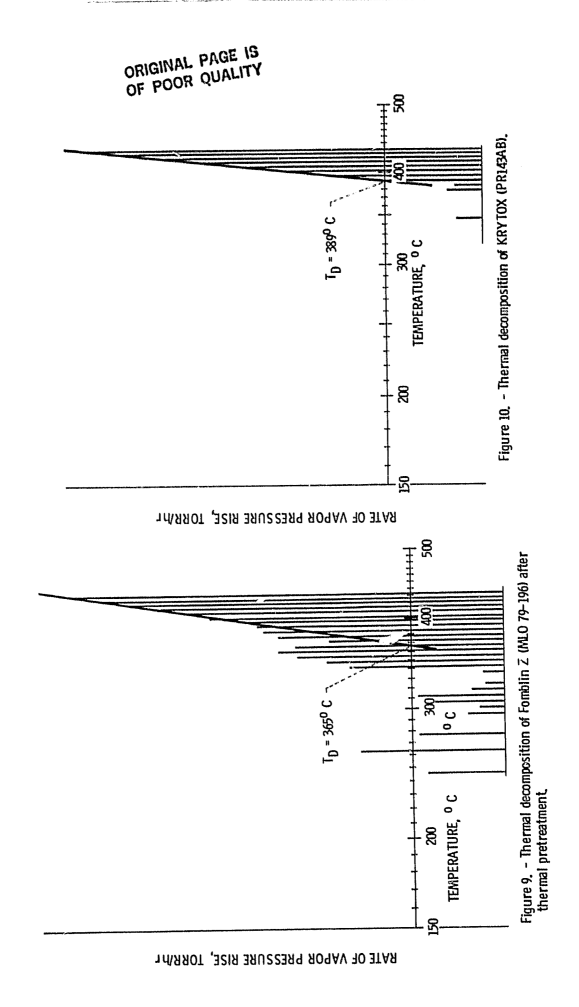
$$\begin{array}{c|c} (C_6H_5)_2\\ P\\ N\\ N\\ I\\ C_3F_7OCF(CF_3)CF_2O(CF_3)CFC\\ N\\ (b) \ Monophospina~s-triazine. \end{array}$$

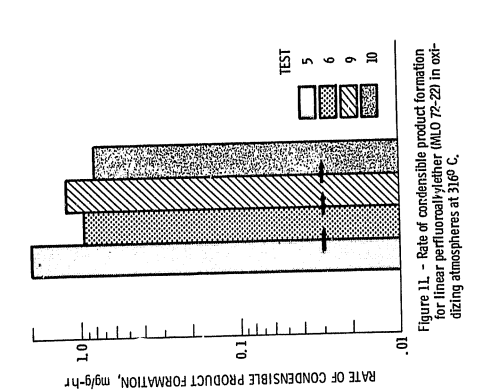
Figure 2. - Chemical structures of degradation inhibitors.











RATE OF CONDENSIBLE PRODUCT FORMATION, mg/g-hr

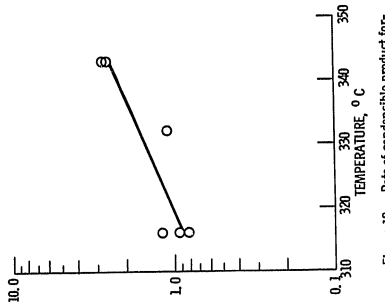
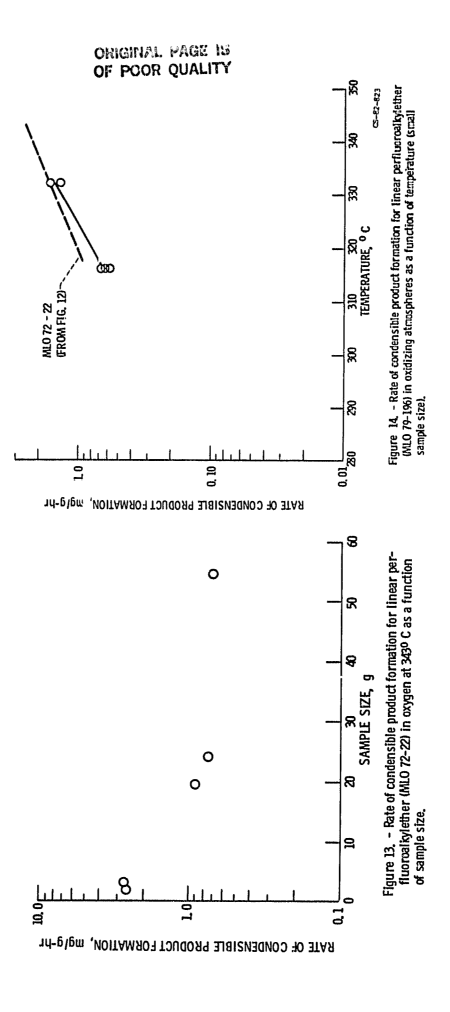
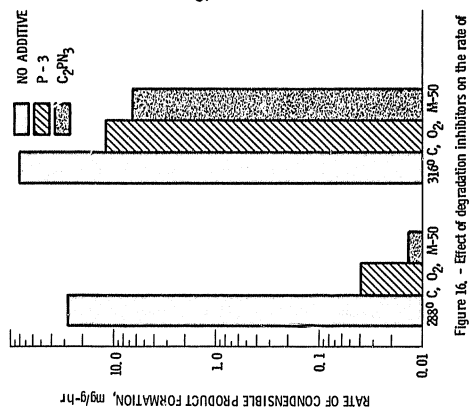
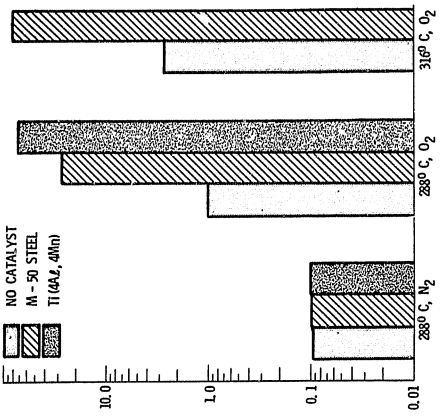


Figure 12, - Rate of condensible product formation for linear perfluoroalkylether (MLO-72-22) in oxygen as a function of temperature (small sample size).



ORIGINAL PAGE IS OF POOR QUALITY





RATE OF CONDESIBLE PRODUCT FORMATION, mg/g-hr

Figure 15. - Rates of condensible product formation for linear perfluoroalkylether (MLO 72-22) in the presence of metal catalysts.

condensible product formation of linear perfluoroalkylether (MLO 72-22).

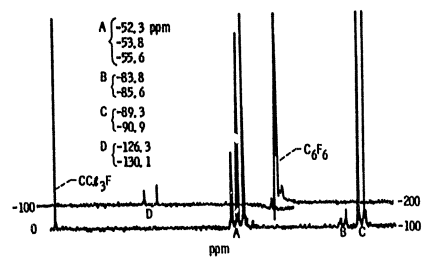


Figure 17. - ¹⁹F NMR spectrum of linear perfluoralkylether fluid MLO-72-22,

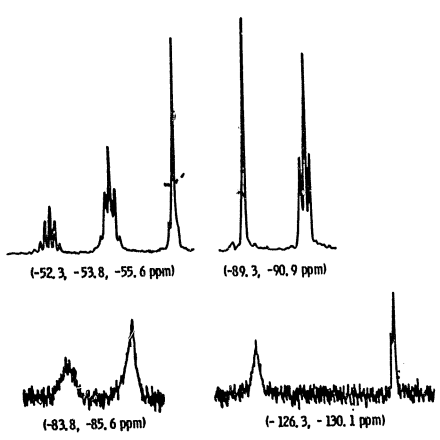


Figure 18. – ¹⁹F NMR spectrum of fluid MLO-72-22, expanded sections A, B, C, and D.

ORIGINAL PAGE IS OF POOR QUALITY

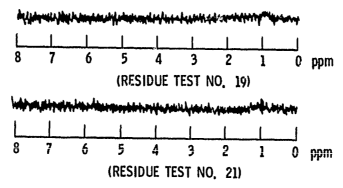


Figure 19. - Proton NMR spectra of fluid MLO 72-22.