DECAY SEQUENCES - ION SERIES OF MASS SPECTRA OF POLYOXAPERFLUOROALKANES AND POLYOXAPERFLUOROALKILS WITH TERMINAL HALIDE ATOMS

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Abstract: This report presents the ionic series of mass spectra of polyoxaperfluoroalkanes, as well as their derivatives with terminal halide atoms. In comparison with ionic series of linear perfluoroalkanes, as well as its derivatives with terminal halide atoms. Fragmentation changes due to oxygen atoms of the chain, as well as the structure of terminal groups, are discussed.

Keywords: ion series of mass spectra, regular and irregular fragment groups, OCF₂ emission, rearrangements with formation of terminal fluorocarbonyl group, double series of rearranging ions, double series of fragmentation of molecular bromine-containing ions.

Experimental part

Samples of polyoxaperfluoroalkanes and polyoxaperfluoroalkils with terminal halide atoms, the spectra of which are not available in NIST libraries, were provided by NPO PiM-Invest CJSC. Electron ionization mass spectra were recorded via Finnigan Polaris Q chromatomass spectrometer (with ion trap, range 14-1000 Da, energy 70 eV and with Rtx-5MS capillary column (5% diphenyl/95% dimethyl polysiloxane 30 m long, 0.25mm-inner diameter, initial column temperature 30°C, isotherm 10 min; heating with a speed of 10°/min up to 250°C)). Since this report also contains spectra from NIST libraries obtained via magnetic or quadrupole apparatus, all presented mass spectra contain information about the type of apparatus on which this mass spectrum was recorded.

Introduction

In processes of electrosynthesis by Kolbe, the aliphatic products of radical dimerization were isolated and interpreted by NMR¹⁹F and chromatomass spectrometry methods [1]. For the first time,
the homologues of a large number of polyoxaperfluoroalkanes and polyoxaperfluoroalkils with terminal halide atoms and with mass numbers from 500 to 1900 Da were synthesized and their mass spectra were recorded.

It was of interest to find out how oxygen chain links change fragmentation and ion series, in comparison with ion series of mass spectra of perfluoroalkanes and perfluoroalkyl halides. Fragmentation of linear \( n \)-alkanes and \( n \)-perfluoroalkanes always begins with terminal groups of \( \text{CH}_3 \) and, respectively, with \( \text{CF}_3 \). This is not only because two terminal groups with three hydrogen or fluorine atoms (compared to \( \text{CH}_2 \) or \( \text{CF}_2 \) groups in chain) are a more likely target for electron attack, but also because they are terminal groups.

There are two ion series in mass spectra of linear perfluoroalkanes \([2,3]\): the series of perfluoroalkyl ions \([\text{C}_n\text{F}_{2n+1}]^+\) and the series of perfluoroalkenyl ions \([\text{C}_n\text{F}_{2n-2}]^+\). The perfluoroalkyl series, terminating with \({}^+\text{C}_3\text{F}_7\), \({}^+\text{C}_2\text{F}_5\) and \({}^+\text{CF}_3\) ions, occurs as a result of primary detachment of fluorine atom and subsequent emissions of difluorocarbene. The series of perfluoroalkenyl ions \([\text{M}-3\text{F}]\) begins with detachment of two fluorine atoms and formation of rearrangement cation radical with terminal perfluorovinyl group protecting one of the «flanks of the chain» from fragmentation. The excited rearranging cation radical emits the third fluorine atom. As a result, the opposite terminal group \( \text{CF}_3 \) becomes the group \(+\text{CF}_2\) \((1)\). Subsequent fragmentation with detachments of difluorocarbene is completed by a series of perfluoroallyl ions. Detachment of two fluorine atoms with formation of perfluorovinyl group, compared with detachment of one fluorine atom, may lead to a gain in energy required for detachment of third fluorine atom. The formation of two ion series can also be the result of difference in removal energies of one or three fluorine atoms. The reason for this difference in excitation energy \(+.\text{M}\) can be two variants of energy transfer that occur when an electron is removed - minimum and maximum. Along with values of radical detachment energies \(\text{CF}_2=\text{CF}\) and \(\text{F}\). a certain “detachments symmetry” also plays a role in fragmentation processes, since the detachment of perfluorovinyl radical would completely exclude the possibility of perfluoroalkenyl series formation.

\[
[M]^+ - 2\cdot \text{F} \rightarrow \text{CF}_3(\text{CF}_2)_n^+ \cdot \text{CF-CF}_2^- \cdot \text{F} \rightarrow {}^+\text{CF}_2(\text{CF}_2)_n\text{CF=CF}_2
\]

(1)

Detachment of \( \text{M-57} \) (\( \text{M-3F} \)) is usually accompanied by emission of \( \text{CF}_2 \), so that instead of \( \text{M-57} \) it corresponds to detachment of \( \text{M-107} \). It is in this way occurs the formation of first fragment ion with \( m/z \) 931 0.5% (M-107) perfluoroeicosane \( \text{C}_{20}\text{F}_{41} \) \([4]\) MW 1038 NIST#: 239239 ID#: 36518 DB: mainlib.
Ionic series of mass spectra of α,ω-dihalogenperfluoroalkanes were also studied, in particular, 1,2-dibromo-1,1,2,2-tetrafluoroethane, 1,4-diiodoperfluorobutane C₆F₃I₂, 1,6-dichloroperfluorohexane C₆Cl₂F₁₂ and 1,7-dibromoperfluoroheptane C₇Br₂F₁₄ [4].

Mass spectra of α,ω-dihalogenperfluoroalkanes include four ionic series. These series terminate with following ions: (⁺CF₃ m/z 69 - perfluoroalkyl series), (⁺C₃F₅ m/z 131 - perfluoroalkenyl), (⁺CF₂Hal - halofluoroalkyl) and (⁺C₃F₄Hal - halofluoroalkenyl series).

The number of ion series probably corresponds to number of electrons being removed, as well as - to number of electrons whose detachments as a result of «optimal» excitation energy transfer options leads to increased energy +.M and their «additional» fragmentation detachments.

Perfluoroalkanes and perfluoroalkiles with terminal halide atoms contain regular fragment groups (CF₂)ₙ, which greatly simplifies the interpretation of their ion series. In contrast, the chains of polyoxaperfluoroalkanes molecules are heterogeneous because they contain fragments with different masses: O (m/z 16), CF₂ (m/z 50) и CF(CF₃) (m/z 100).

In mass spectra of polyoxaperfluoroalkanes with +.M ≥ 700 Da, the primary radical detachments occur, as a rule, for oxygen atom. In this case, the emitted radical may contain one or more oxygen atoms.

Despite the standard value of ionization energy, the mass spectra of polyoxaperfluoroalkanes obtained via mass spectrometer with mass analyzer («ion trap») differ from spectra taken under standard recording conditions - via magnetic and quadrupole devices.

The difference in mass spectra taken via ion trap is due not only to probability of protonation of molecular ions and possibility of ion-molecular reactions, but also - to distortion of peak intensities of molecular and fragment ions. The main reason for distortion of these spectra recorded via device with ion trap is ion separation time.

The time spent by ions and their separation in an ion trap is longer than in a magnetic sector or in a quadrupole filter. Due to increase in ion separation time, the fragmentation in ion trap occurs with a shift in intensity of peaks of successively fragmenting ions - from peaks with higher masses to peaks with lower masses. As a result, the peak intensities of molecular and primary fragment ions decrease, and the peak intensities of secondary and final ions increase (but the general nature of fragmentation does not change).

Fig. 1 shows the mass spectra of 1,1,1,2,3,4,4,4-octafluoro-2,3-bis(perfluoropropoxy)butane C₁₀F₂₂O₂ (MW 570) obtained via magnetic equipment (VG 7070E) and ion trap (Polaris Q).
The weak peak $^{+}\text{M-CF}_3$ m/z 501, which is present in magnetic spectrum, does not appear in spectrum recorded via equipment with ion trap.

These spectra differ in masses of base peaks: m/z 169 (VG) and 69 (Polaris Q). In addition, they also differ in that there is intense peak $^{+}\text{M-C}_3\text{F}_7$, $^{+}\text{M-2F}$ (48%) for rearranging ion with m/z 363 in spectrum recorded via equipment with ion trap, which does not manifest itself in magnetic spectrum. The ion with m/z 363 is the parent ion in series: 263, 197, 147, 97, 47, formed both in magnetic spectrum and spectrum recorded via equipment with ion trap.

Ion with m/z 363 appears to have the structure $^{+}\text{CF}_2(\text{CF}_2)_2\text{O-CF(CF}_3)-\text{CF}_2=\text{O}$. Another difference between compared mass spectra is that the peak of M/2 ion with m/z 285 (5.2%) appears only in spectrum recorded via equipment with ion trap (VG). Although the mass spectra recorded via equipment with ion trap do not correspond to standard conditions for record of spectra via magnetic and quadrupole equipment, they include all main fragment ions.

This makes it possible to establish their ion series and (then compared with magnetic spectra) obtain the additional information confirming the fragmentation pathways.

**Ionic series of polyoxaperfluoroalkanes and general patterns of their fragmentation**

In mass spectra of polyoxaperfluoroalkanes, as well as of polyoxaperfluoroalkyl halides presented in this report, there are peaks of ions with m/z 147, m/z 97, m/z 47, the occurrence of which
is due to simultaneous emission of two fluorine atoms and subsequent rearrangement leading to terminal fluorocarbonyl (acetylfluoride) group (see Fig. 2). When a fluorocarbonyl group occurs and one of terminal groups stabilizes, the detachment of radical $F$ occurs from opposite terminal group and subsequent fragmentation leading to a series of ions with $m/z$ 197, 147, 97.47.

Due to ease of primary detachment of radical from oxygen atom or with oxygen atom, the molecular ions do not appear in mass spectra of polyoxaperfluoroalkanes.

Example of mass spectrum of oxaperfluoroalkane with one oxygen atom is spectrum of 1,1-oxybis(pentafluoroethane). Mass spectrum of compound (see Fig. 2) consists of two series: for oxaperfluoroalkyl, leading to formation of perfluoroalkyl ions with $m/z$ 119 and 69 (marked in red), and for rearrangement series (marked in blue), terminating with peaks of ions with $m/z$ 147, 97 and 47.

![Figure 2. Two ion series of mass spectrum of 1,1'-oxybis(pentafluoroethane) $C_4F_{10}O$ MW:254
Ion Trap (Polaris Q).](image)

When two fluorine atoms are detached from $^+\text{CF}_2\text{OC}_2\text{F}_3$ ion with $m/z$ 185, the rearrangement ion (having $m/z$ 147) with terminal fluorocarbonyl group occurs. It emits the difluorocarbene to form fragment ions with $m/z$ 97 and 47 containing a fluorocarbonyl group.
Terminal perfluorooxaallyl group $\text{OCF}_2\text{CF}=\text{CF}_2$ stabilizes one of the flanks of $^+\cdot\text{M}$ chain, excluding its fragmentation. It is thanks to it that molecular ion appears in mass spectrum. Both ion series of spectrum (see Fig.3): for oxaperfluoroalkyl (marked in red) and for oxaperfluoroallyl (marked in blue) begin with detachments of $\text{OC}_3\text{F}_6$ and $\text{OC}_3\text{F}_7$ radical from opposite terminal group $\text{OC}_3\text{F}_7$. Rearrangements occur in both ion series (rearrangement ions are marked in bold).

![Figure 3. Two ion series of C₈F₁₆O₂ MW:432 mass spectrum:432 Ion Trap (Polaris Q).](image)

When $\cdot\text{CF}=\text{O}$ radical is detaching from ion with m/z 266 (series marked in red), the rearrangement ion $^+\text{C}_4\text{F}_9$ with m/z 219 arises, that fragmenting with formation of a series of perfluoroalkyl ions. As a result of successive detachments of $\cdot\text{F}$, $\cdot\text{OC}_3\text{F}_6$, $\cdot\text{CF}_2$, $\cdot\text{CF}_2$ the oxy-perfluoroallyl ion $^+\text{OCF}_2\text{CF}=\text{CF}_2$ with m/z 147 (series marked in blue) appears.

When oxygen atom is detached from ion with m/z 147 (or fluorine atom - from $^+\text{C}_3\text{F}_6$ ion with m/z 150) (see Fig.3), the base ion of C₈F₁₆O₂ spectrum becomes $\text{CF}_2=\text{C}=\text{O}$ ion with m/z 131.

$\text{CF}_2=\text{CF}=\text{CF}_2+\cdot\text{O}$ ion with m/z 147 does not contain any fluorocarbonyl group. When two fluorine atoms are detached from it and rearranged into $\text{CF}_2=\text{C}=\text{O}$ ion with m/z 109, any fluorocarbonyl group is also not formed. However, the final fragment ion $^+\text{CF}=\text{O}$ m/z 47 contains a
fluorocarbonyl group. Detaching of oxygen atom from ion with m/z 109 is completed by rearrangement of \(^\text{+C}_3\text{F}_3\) ion with m/z 93.

Fig. 4 shows two ion series of mass spectrum \(\text{C}_{10}\text{F}_{22}\text{O}_2\text{MW}:570\).

\[\text{Figure 4. Two ion series of } \text{C}_{10}\text{F}_{22}\text{O}_2 \text{MW:570 mass spectrum:570 Ion Trap (Polaris Q)}.\]

Despite the fact that this compound is symmetrical, there is no decay of M/2. Perfluoroalkyl ion series (marked in red) begins with detachment of \(^\text{OC}_3\text{F}_7\) (-185) radical, then - with detaching of \(\text{C}_2\text{F}_4\) molecule and subsequent emission of \(\text{O=CF}_2\) molecule, with formation of a rearrangement ion \(^\text{+CF}_2\text{C}_3\text{F}_7\) with m/z 219, i.e. group that is not present in original molecule. The rearrangement series of ions (marked in blue) begins with detachment of \(^\text{C}_3\text{F}_7\) radical (without oxygen) as well as - with emission of two fluorine atoms from two \(\text{CF}_3\) groups (the opposite terminal and one of central \(\text{F}_3\text{CCF}\) groups (summary -207). As a result, there is the rearrangement ion with m/z 363 with terminal fluorocarbonyl group. Fragmentation of ion with m/z 363 results in a low-intensity series of fluorocarbonyl ions with m/z 197, 147, 97 and 47. Since there is no series of ions with terminal perfluorovinyl group in spectrum (see Fig. 4), the appearance of intense peak (55%) of perfluoroallyl ion \(\text{CF}_2=\text{CF}^+\text{CF}_2\) with m/z 131 is probably the result of detaching of oxygen atom from the ion with m/z 147.

Fig. 5 shows two ion series of \(\text{C}_{12}\text{F}_{26}\text{O}_3\) mass spectrum. Intense series of ions ending with perfluoroalkyl ions (marked in red), and its rearrangement series (marked in blue), terminates with ions of fluorocarbonyl group. In presented mass spectrum, the number of peaks of recorded ions is only thirteen. Probably due to insufficient amplification or low sensitivity of spectrometer, the low-
intensity ion peaks did not appear in this spectrum (in particular, the peak of ion with m/z 147 (0%) and the peak of ion with m/z 47).

**Figure 5.** Two ion series of mass spectrum of perfluoro2,8-trimethyl-3,6,9-trioxadodecane C\textsubscript{12}F\textsubscript{26}O\textsubscript{3} MW:686 NIST#: 98335 ID#: 36521 DB: mainlib.

Of the two possible options for starting fragmentation [C\textsubscript{12}F\textsubscript{26}O\textsubscript{3}]\textsuperscript{+}, it does not start with \textsuperscript{12}C(CF\textsubscript{3})\textsubscript{3} group (having m/z=219) with a tertiary carbon atom, and with perfluoropropyl group \textsuperscript{13}C\textsubscript{3}F\textsubscript{7} (having m/z=169). After detachment of trifluoromethyl radical (-F, -CF\textsubscript{2}), another emission of fluorine atom occurs (with formation of rearranging cation radical with m/z 598).

Subsequent fragmentation, including detachment of rearrangement \textsuperscript{12}C\textsubscript{3}F\textsubscript{2}OC=O -97 radical, emission of OC\textsubscript{3}F\textsubscript{6} -166 and detachment of CF\textsubscript{2}, leads to ion \textsuperscript{12}C\textsubscript{3}F\textsubscript{2}OC(CF\textsubscript{3})\textsubscript{3} with m/z 285. Before formation of ion with m/z 285, the group \textsuperscript{12}C(CF\textsubscript{3})\textsubscript{3} with tertiary carbon atom does not participate in fragmentation. \textsuperscript{13}C\textsubscript{3}F\textsubscript{2}OC(CF\textsubscript{3})\textsubscript{3} 8% ion with m/z 285 fragments in two ways: with emission of CF\textsubscript{2}=O and formation of perfluoroalkyl ion \textsuperscript{13}C(CF\textsubscript{3})\textsubscript{3} with m/z 219 (11%) (series in red), and with emission of hexafluoroethane molecule and formation of rearrangement ion \textsuperscript{13}C\textsubscript{3}F\textsubscript{2}OC=CF\textsubscript{2} with m/z 147 (0%) and its fragmented ion \textsuperscript{13}C\textsubscript{2}F\textsubscript{2}OC=O with m/z 97 (4%) (series in blue).

Fig. 6 shows two ionic series of polyoxaperfluoroalkane C\textsubscript{12}F\textsubscript{26}O\textsubscript{4} MW=702, the molecule of which has a plane with mirror symmetry, as well as a vertical axis of symmetry passing in the plane of symmetry (between two central groups F-C-CF\textsubscript{3}). There is no ion peak M/2 =351 in mass spectrum of this compound. Primary radical detachments occur at one of central oxygen atoms.
Two ion series (see Fig. 6.) of mass spectrum (marked in red and blue) begin with rearranging ions \{with \textit{m/z} 385 and \textit{m/z} 429\}. Ion with \textit{m/z} 385, occurs when O=CF$_2$ is detached from terminal group CF$_3$O for ion with \textit{m/z} 451. As a result, its terminal group becomes OC$_3$F$_7$ group. Ion with \textit{m/z} 429 occurs after terminal group C$_3$F$_6$OCF$_3$ is emitted and two fluorine atoms are detached (marked in blue dotted line in structural formula). As a result, its terminal group becomes the fluorocarbonyl group FC=O.

The rearrangement series (marked in blue) terminates with ions having \textit{m/z} 197, 147, 97 and 47 and containing a fluorocarbonyl group. Ion $^+$CF$_2$OC$_3$F$_7$ with \textit{m/z} 235 (3\%), occurs in this series (marked in red) and terminates with perfluoroalkyl ions. $^+$CF$_2$OC$_3$F$_7$ ion loses the oxygen atom to form $^+$C$_4$F$_9$ ion with \textit{m/z} 219 (13\%), i.e. a group that is not present in original M$^+$. The emission of oxygen atom confirms the energy benefits of formation of homogeneous perfluoroalkyl rearrangement ion $^+$C$_4$F$_9$.

Fig. 7 shows two ionic series of polyoxaperfluoroalkane C$_{17}$F$_{34}$O$_5$ with central carbonyl group. In series terminating with perftoalkyl ions (marked in red), the detachment of C$_3$F$_7$OCF(CF$_3$)CF$_2$OCF(CF$_3$)-C=O [- C$_9$F$_{17}$ O$_3$ - \textit{m/z} 479] is accompanied by emission of another oxygen atom. As a result of these two detachments (- \textit{m/z} 495) the rearrangement ion C$_3$F$_7$OCF(CF$_3$)CF$_2^+$CF(CF$_3$) with \textit{m/z} 435 arises. Its further fragmentation includes three CF$_2$ detachments with formation of C$_3$F$_7$O$^+$CF(CF$_3$) ion having \textit{m/z} 285.
Final emission of F₂C=O molecule leads to rearrangement of perfluoroalkyl ion C₃F₇⁺CF₂ with m/z 219 (i.e. the group that is not present in original compound).

In the absence of perfluoroallyl ion series, the appearance of weak peaks of ions with m/z 231 and 181 is the result of detachment of oxygen atom from the ion with m/z 247, as well as from the ion with m/z 197.

Two ion series of mass spectrum of polyoxaperfluoroalkane C₃₄F₇₀O₁₀ (MW 1898) are presented (see Fig.8).

In spectrum, recorded within the range of 1000 Da, the first recorded peak is the peak of ion with mass of 667. The peak-free zone for mass spectrum within the range 1898 - 667 at 1231 Da is the result of instability of molecular cation radical with m/z 1898. Decay of +.M can occur at detachment of radical M/2 (m/z 949) + CF(CF₃)OCF₂CF(CF₃)O (m/z 282) with formation of the stable ion [C₁₂F₂₅O₃⁺ C₃₄F₇₀O₁₀⁻OCF₂O⁻CF₂OCF(CF₃)CF₂OCF(CF₃)⁻CF₂ (9%) with m/z 667. As a result, the mass of detached radical (1231 Da) exceeds the mass of resulting ion by 564 Da Given that C₃₄F₇₀O₁₀ molecule is symmetric, there is reason to believe that the decay of M⁺ can also occur with emission of neutral OCF(CF₃) molecule OCF₂OCF(CF₃)⁻CF₂OCF(CF₃)⁻CF₂OCF(CF₃)⁻CF₂⁻CF₂ with a mass of 564 Da from central part of molecular cation-radicl (with formation of symmetric cation-radical [C₂₄F₅₀O₆⁺], having m/z 1334. The symmetric cation-radical further decays into ion and radical with the same masses equal to 667Da In this case, the mass of resulting ion [C₁₂F₂₅O₃⁺ with m/z 667 is greater than the mass of detached neutral molecule C₁₀F₂₀O₄ by 103 Da. Both pathways
of fragmentation of cation $^{+}C_{12}F_{25}O_{3}$ m/z 667 (9%) formed from $C_{34}F_{70}O_{10}$ (Polaris Q, Ion Trap) (see Fig. 8) lead to the same ion with m/z 285.

One pathway is the path of four consecutive, low-intensity detachments, and other way - two detachments with a total mass of -382.

**Figure 8.** Two ion series of mass spectrum ($C_{17}F_{35}O_{5}$)$_2$ MW:1898 Ion Trap (Polaris Q).

Emission of 285 - 66 [CF$_2$O] = 219 (series in red) leads to rearrangement of $^{+}C_4$F$_9$ ion with m/z 219 (27%) – to a group that is not present in original molecule. Detachment of difluorocarbene and two fluorine atoms 285 - 50, -38 = 197 (8%) (series in blue) to rearrangement ion O=CFC$_2$F$_4$*CF$_2$ with m/z 197 (8%) to fluorocarbonyl group. Ion with m/z 197 fragments to form the intense peak of ion with m/z 147 (95%) and its fragmented ion 97 (18%). Despite the absence of perfluoroallyl ion series in spectrum, the intensity of peak for CF$_2$=CF-$^+$CF$_2$ ion with m/z 131 is high (62%). Possible pathways to occurrence of ion with m/z 131 can be either detachment of oxygen atom from ion O=CFCF$_2$*CF$_2$ with m/z 147 (blue dotted line) or detachment of fluorine atom from ion $^{+}C_3$F$_6$ (red dotted line).

**Ionic series of polyoxaperfluoroalkyl with a terminal chlorine atom**

Fig. 9 shows the ion series of mass spectrum of 1-(2-chloro-1,1,2,2-tetrafluoroethoxy)-pentadecaftorgeptan - compounds with one oxygen atom.
Decays of molecular cation radicals with formation of perfluoroalkyl series (in Fig. 9 marked in red) and also - chloroperfluoroalkyl series (marked in green), occur for oxygen atom - the only «heterogeneity» of chain. After detachments of oxygen - containing radicals $\text{OC}_7\text{F}_{15}$ (series in green) and $\text{OC}_2\text{F}_4\text{Cl}$ (series in red) generate $\text{ClCF}_2^+\text{CF}_2$ with $m/z$ 135 and $\text{CF}_3(\text{CF}_2)_5^+\text{CF}_2$ with $m/z$ 369 ions.

Detachment of $\text{CF}_2\text{CF}_2\text{Cl}$ radical without oxygen atom is accompanied by additional emission of two more fluorine atoms (series in blue), detached from opposite terminal pentfluoroethyl group. As a result of these detachments, the rearrangement ion $\text{CF}_2=\text{CF}(\text{CF}_2)_5\text{O}^+$ ($m/z$ 347 (1%)), with terminal perfluorovinyl group, is formed. It emits $\text{O}=\text{CF}_2$ molecule, forming a series of intense peaks of ions with $m/z$ 281, 231, 181, culminating in formation of a perfluoroallyl ion with $m/z$ 131(53%) (marked in blue). Another pathway for ion fragmentation with $m/z$ 347 terminates with a low-intensity series of ions with a fluorocarbonyl group (marked with blue dotted line). Formation of series for ions with $m/z$ 197, 147, 97 and 47 confirms that the ion with $m/z$ 347 is also rearranged to form a terminal fluorocarbonyl group (see Fig. 9).

Three ion series of $\text{C}_9\text{ClF}_{19}\text{O}_2$ mass spectrum are shown in Fig.10.
Replacement of terminal trifluoromethyl group $\text{C}_9\text{ClF}_9\text{O}$ (see Fig. 9) with heptafluoropropyl group $\text{CF}$(CF$_3$)$_2$ (see Fig. 10), with tertiary carbon atom and increase in a number of oxygen atoms in chain by another atom leads to a significant change in fragmentation of $\text{C}_9\text{ClF}_9\text{O}_2$. The series of allyl ions with m/z 281, 231, 181 is not formed. Compared with trifluoromethyl group (see Fig. 9), the heptafluoropropyl group with m/z 219 (see Fig. 10) fragments with detachment of fluorine atom, and then - with emission of hexafluoroacetone. This reduces the excess excitation energy of sequentially fragmenting chlorine-containing ions (marked in green), the number of which in series, compared with the spectrum of $\text{C}_9\text{ClF}_9\text{O}$ (see Fig. 9) increases from two to five (see Fig. 10). The presence of two oxygen atoms in chain increases the likelihood of a series of ions containing a fluorocarbonyl group and eliminates the possibility of formation of perfluoroallyl series terminating with ion $\text{CF}_2=\text{CF}^-\text{CF}_2$ having m/z 131. Intensity of ion peak with m/z 147 increases to 29% and intensity of ion peak with m/z 131 in spectrum of $\text{C}_9\text{ClF}_9\text{O}$ (in Fig. 10 not represented) is only 3%. Given the high intensity of ion peak with m/z 147, the ion $^+\text{C}_3\text{F}_5$ with m/z 131 may be formed by detachment from ion $\text{O}=\text{CFCF}_2^+\text{CF}_2$ with m/z 147.

Fig. 11 shows three ionic series of mass spectrum of polyoxaperfluoroalkyl chloride $\text{C}_{12}\text{ClF}_{25}\text{O}_3$ MW:702. Compared with ion series of mass spectrum of polyoxaperfluoroalkane $\text{C}_{12}\text{F}_{26}\text{O}_4$ (with the same molecular weight MW:702) (see Fig. 6), during fragmentation of which the primary detachment of $\text{OC}_3\text{F}_6\text{OFC}_3$ and $\text{C}_3\text{F}_6\text{OFC}_3 + 2\text{F}$ radicals occurs, with masses of -251 and -273 Da, in spectrum of polyoxaperfluoroalkyl chloride (see Fig. 11) the mass of primary radical emissions: atom $\text{F}^-\text{19}$ and $\text{OC}_3\text{F}_6\text{Cl}^-\text{201}$ Da are significantly smaller.
The terminal chlorine atom (with mass of 35/37) stabilizes the perfluoropolyalkyl chloride molecule, resulting in a minimum detachment of fluorine atom by mass, but with subsequent emission of -C(CF₃)₂OC₃F₆O radical with mass of 332, there is a chlorine-containing ion ClC(F₆O)₂CF₂ with m/z 351 (2%) and mass of M/2.

When increasing the number of oxygen atoms in the chain of polyoxaperfluoroalkyl chlorides, there is a noticeable increasing in intensity of peaks containing the fluorocarbonyl group (197, 147, 97, 47). Appearance of series of perfluoroallyl ions becomes impossible, however a weak peak of perfluoroallyl ion with m/z 131 (3%) is formed when detachment of oxygen atom from the ion with m/z 147 (see Fig. 11).

In mass spectra of α,ω-dichloropolyoxaperfluoroalkanes: [Cl(C₃F₆O)₂CFCF₃]₂ MW:934, [Cl(C₃F₆O)₃CFCF₃]₂ MW:1266, [Cl(C₃F₆O)₄CFCF₃]₂ MW:1598 large peak-free zones occur (up to M/2). This is probably due to the fact that stabilizing effect of two terminal chlorine atoms is directed in opposite direction. In C₁₆Cl₂F₃₂O₄ - C₂₈Cl₂F₅₆O₈ mass spectra the peaks of primary ions have the same or similar m/z values and similar intensity values. Thus, the first chlorine-containing ions of their mass spectra are the following:

C₁₆Cl₂F₃₂O₄ MW:934 - (M/2) + C₂F₄ = [C₁₀ClF₂₀O₂] m/z 567 (11%);
C₂₂Cl₂F₄₄O₆ MW:1266 - (M/2) - OCF₂ = [C₁₀ClF₂₀O₂] m/z 567 (6%);
C₂₈Cl₂F₅₆O₈ MW:1598 - (M/2) - OCF(CF₃) = [C₁₂ClF₂₄O₃] m/z 683 (8%);
In mass spectra of these three compounds (with two terminal chlorine atoms) the intensities of perfluoroallyl ion $\text{CF}_2\text{CFCF}_2$ with m/z 131 and the ion with fluorocarbonyl group $\text{^+CFCF}_2\text{CFCF}_2\text{=O}$ with m/z 147 are quite high (about 50-80%). However, if the sequence of a series of ions with fluorocarbonyl group 197, 147, 97 and 47 is well manifested in a spectra, then the intense perfluorallyl ion (46%) is not the allyl series ion, but is formed from $\text{^+CFCF}_2\text{CFCF}_2\text{=O}$ ion with m/z 147 when oxygen atom or $\text{^+C}_3\text{F}_6$ ion is separated from it (when the fluorine atom is detached from it).

Fig. 12 shows three ion series of $\text{C}_{16}\text{Cl}_2\text{F}_{32}\text{O}_4$ mass spectrum. The presence of two terminal chlorine atoms in $\text{C}_{16}\text{Cl}_2\text{F}_{32}\text{O}_4$ molecule does not lead to appearance of molecular ion in spectrum. The mass of primary radical, detached from chlorine-containing series (marked in green), is 367 Da. Compared with ionic series of monochloride (see Fig.11) in dichloride series (see Fig.12), the peak-free zones increase to 367 and 383 Da.

The first rearrangement ion $\text{^+CF}_3\text{CF}_2\text{CF(CF}_3\text{)OCF}_2\text{OC}_2\text{F}_4\text{=O}$ with m/z 551 Da (series marked in red in Fig. 12) occurs after detachment of radicals $\text{ClC}_3\text{F}_6\text{OCF}_2\text{CF}_2$ (- 301) (- 301) and $\text{C}_3\text{F}_6\text{C}=\text{O}$ (-47), as well as after emission of second chlorine atom. Ion with m/z 551 contains the terminal rearrangement group $\text{C}_4\text{F}_9$, which is not present in original molecule.

Fragmentation of this series (marked in red) leads to perfluoroalkyl ions with m/z 219, 169, 119 and 69. Rearrangement ion with m/z 551 also fragments with emission of $\text{C}_3\text{F}_6$ molecule and two fluorine atoms, forming a series of ions with fluorocarbonyl terminal group (marked in blue dotted and solid line) with m/z 197, 147, 97 and 47.

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**Figure 12.** Three ion series of mass spectrum $\text{C}_{16}\text{Cl}_2\text{F}_{32}\text{O}_4$ MW: 934 Ion Trap (Polaris Q).
The intense peak (71%) of perfluoroallyl ion $\text{CF}_2=\text{CF}^+\text{CF}_2$ with m/z 131 is not allyl series ion, but is formed when the oxygen atom detaches from ion $^+\text{CF}_2\text{CF}_2\text{FC}=\text{O}$ (m/z 147), or the fluorine atom - from ion $^+\text{C}_3\text{F}_6$ (m/z 150).

**Ionic series of polyoxaperfluoroalkyl with terminal bromine atom**

Fig. 13 shows four ion series of $\text{C}_7\text{BrF}_{15}\text{O}_2$ mass spectrum.

![Figure 13](image)

**Figure 13.** Four ion series of mass spectrum $\text{C}_7\text{BrF}_{15}\text{O}_2$ MW:480 Ion Trap (Polaris Q).

Two bromine-containing ion series is marked in green. One series of perfluoroalkyl is marked in red and one series of ions containing a fluorocarbonyl group - is marked in blue.

In bromine-containing series of ions with more intense ion peaks, after emission of fluorine atom, the detachment of $\text{C}_2\text{F}_4$ and two atoms $\text{F}$ occurs with formation of rearrangement ion $\text{BrC}_2\text{F}_4\text{OCF}(^+\text{CF}_2)\text{FC}=\text{O}$ with m/z 323.

Ion with m/z 323 emits the dicarbonyl difluoroethane molecule $\text{O}=\text{CF}-\text{CF}=\text{O}$, turning into ion $\text{BrC}_2\text{F}_4^+\text{CF}_2$ with m/z 229. In a less intense bromine-containing series (after detachment of $\text{OC}_2\text{F}_5$ and formation of $\text{BrC}_2\text{F}_4\text{OCF}(\text{CF}_3)^+\text{CF}_2$ ion with m/z 345), the emission of $\text{C}_2\text{F}_4$ and two fluorine atoms results in rearrangement ion $\text{BrC}_2\text{F}_4\text{C}=\text{O}^+\text{CF}_2$ with m/z 207.

The subsequent detachment of $\text{CF}_2$ with formation of $\text{BrC}_2\text{F}_2^+\text{C}=\text{O}$ ion with m/z 157 (5%) and emission of CO (-28) completes $\text{Br}^+\text{CF}_2$ ion with m/z 129 (47%).

A series starting with emission of $\text{Br}^+\text{CF}_2$ and detachment of two fluorine atoms (marked in blue), terminates with ions having m/z 197,147, 97 and 47 and containing a fluorocarbonyl group.
A series in which the detachment of BrC$_2$F$_4$O with maximum mass occurs (~195 Da) (marked in red) terminates with perfluoroalkyl ions. All four series of ions (see Fig. 13) are rearranging. In three series, additional, simultaneous detachments of two fluorine atoms occurs, and in one series (marked in red) - emission of OCF$_2$ molecule from the middle of chain.

Fig. 14 shows three ion series of C$_{11}$BrF$_{23}$O$_3$ mass spectrum.

![Figure 14. Three ion series of mass spectrum C$_{11}$BrF$_{23}$O$_3$ MW:696 Ion Trap (Polaris Q).](image)

Of the three ionic series of C$_{11}$BrF$_{23}$O$_3$ spectrum, presented in Fig.14, in two series of detachments of bromine-containing radicals occurs: in BrC$_2$F$_4$O series marked in red and in BrC$_2$F$_4$ + 2F series marked in blue. The series marked in red terminates with perfluoroalkyl ions. The first fragment ion in blue series with m/z 479 is a rearrangement ion with terminal fluorocarbonyl group. As a result of detachments from it - C$_3$F$_6$O (marked by blue dotted line), or two detachments (-F-C=O и -C$_2$F$_4$, -F)) (marked by two solid blue lines), the ion with m/z 313 arises. It fragments to form a series of ions: 147, 97 and 47 with terminal fluorocarbonyl group.

In bromine-containing series (marked in green), the first bromine-containing ion *C$_3$F$_6$OC$_2$F$_4$Br with a mass of m/z 345 (7%) fragments with emission of OC$_2$F$_4$ and formation of rearrangement ion *BrC$_2$F$_4$+CF$_2$ with m/z 229 (2%) - a group that is not present in original molecule. Perfluoroallyl ion, whose intensity does not exceed 6%, is not indicated in Fig. 14.
Ionic series of two structural isomers $\text{C}_{11}\text{BrF}_{23}\text{O}_3$ with different terminal groups (see Fig. 14 and Fig. 15) differ in intensities of most peaks, as well as in number of arising bromine-containing ions.

Detachment of M-F from isomer with terminal group $\text{CF}_3\text{CF}_2\text{CF}_2\text{O}$ (see Fig. 14) does not occur. In its bromine-containing series (see Fig. 14) the number of bromine-containing ions is less than in bromine-containing series of isomer with terminal group $\text{CF}_3\text{FC}$ (see Fig. 15). In particular, there are no ions with $m/z$ 677, 561 and 395.

Figure 15. Three ion series of mass spectrum $\text{C}_{11}\text{BrF}_{23}\text{O}_3$ MW:696 Ion Trap (Polaris Q).

In another isomer the fluorine atom is detached from terminal group $(\text{CF}_3)_2\text{FCO}$ (see Fig. 15). The sequence of detachments in this bromine-containing series (marked in green) allows us to conclude that next detachment $-\text{OC}_2\text{F}_4$ comes from opposite terminal group $\text{OC}_2\text{F}_4\text{Br}$. As a result, a rearrangement ion with terminal group $\text{OCF(CF}_3)\text{CF}_2\text{Br}$ with $m/z$ 561 arises (see Fig. 15).

Fig. 16 shows three ion series of $\text{C}_{16}\text{BrF}_{33}\text{O}_5$ mass spectrum.
Of the three ion series, shown in Fig. 16, only non-arrangement series is the bromine-containing series (marked in green). It includes seven bromine-containing ions. In most intense polyoxaperfluoroalkyl ion series (marked in red), there are consecutive detachments Br, C$_2$F$_4$O and C$_3$F$_6$O are terminated by ion with m/z 617. Detachment of OC$_2$F$_4$ from opposite terminal group OCF$_2$CF$_3$ leads to a rearrangement ion with m/z 501, with terminal group CF$_3$:CF(CF$_3$)O, which is not present in original molecule. A series of fluorocarbonyl ions with m/z 147, 97, 47 (marked in blue) begins with detachment of C$_2$F$_4$ and two fluorine atoms. There is a rearrangement ion (m/z 761) with terminal fluorocarbonyl group. It fragments with detachment of two radicals OC$_3$F$_6$, emissions of - C$_2$F$_4$, -OCF$_2$, - C$_2$F$_4$, -OCF$_2$, loss of fluorocarbonyl group and its formation again (ions with m/z 147.97, 47).

Fig. 17 shows the ion series of C$_{14}$Br$_2$F$_{28}$O$_4$ mass spectrum.
During fragmentation of dibromide $\text{C}_{14}\text{Br}_2\text{F}_{28}\text{O}_4$ (see Fig. 17), as a result of two different primary detachments of bromine-containing radicals (as well as result of additional emission of two fluorine atoms in one of series) two «parallel» series of bromine-containing ions are formed (marked in green). The primary detachments in these two $\text{C}_{14}\text{Br}_2\text{F}_{28}\text{O}_4$ series differ in that in series with a more intense peaks the detachment of $-\text{BrC}_2\text{F}_4$ (-195) occurs, and in series with a less intense series - detachments of $-\text{BrC}_2\text{F}_4$ and $-2\text{F}$ (-217) from opposite terminal group $\text{BrCF}_2\text{CF}_2\text{O}$. In a less intense series as a result of primary detachments, the rearrangement ion with m/z 705 containing $\text{BrCF}_2\text{C}=\text{O}$ group is formed.

The next, «extraordinary release» $-\text{OC}_4\text{F}_8$ in this series comes from the middle of chain. The presence of carbonyl group in ions with m/z 705, 489 and 207 confirms the detachment of molecule $\text{O=CCF}_2$ (-78) from ion with m/z 229, leading to formation of $\text{Br}^+\text{CF}_2$ ion.

Two different fragmentation pathways $^+\text{M}$ ($\text{C}_{14}\text{Br}_2\text{F}_{28}\text{O}_4$ with formation of two series of bromine-containing ions are either the result of two different excitation energies $^+\text{M}$, or the result of removal of two electrons from molecule that are topologically different with respect to bromine atom.

The series of polyoxaperfluoroalkyl ions (marked in red) begins with emission of bromine atom, as well as with detachment of rearrangement molecule $\text{BrCF}_2\text{FC}=\text{O}$ to form a rearrangement ion with terminal group $\text{CF}_3$ having m/z 667. After a series of detachments of oxygen-containing fragments, the ion with m/z 219 arises, that fragments to form a series of perfluoroalkyl ions.
Another series (marked in blue) begins with formation of rearrangement ion (m/z 646) with terminal FC=CF$_2$ group. This ion series terminates with formation of ions with fluorocarbonyl group (m/z 147, 97 and 47).

**Ionic series of polyoxaperfluoroalkyl with terminal iodine atoms**

In mass spectra of polyoxaperfluoroalkils with terminal chlorine or bromine atom the molecular cation radicals do not appear due to their instability. In some cases, the first fragment ion in their spectra is M-F ion. Compared with other halides, the terminal iodine atom contributes to maximum stabilization of molecular ion, especially if the linear chain of compound is not very large.

In all mass spectra of polyoxaperfluoroalkyliodides presented in this report, there are molecular ions. However, their intensity decreases as molecular weight increases and proportion of iodine atom mass to mass of chain decreases.

Ion series of C$_5$F$_{11}$IO mass spectrum are shown in Fig. 18.

![Figure 18. Three ion series of mass spectrum C$_5$F$_{11}$IO MW:412 Ion Trap (Polaris Q).](image)

Intensity of peak of molecular ion C$_5$F$_{11}$IO with m/z 412 is 17% (see Fig. 18). Using examples of mass spectra of perfluoroalkyl homologues with terminal iodine atom, it was found that when the iodine atom is detached, a base ion [M-I]$^+$ arises if the ratio of stabilized chain mass to iodine mass is 2:1 or 3:1[4]. This rule seems to be (see Fig. 18). It is also preserved for lower homologues of perfluoroalkyl iodides.
Since the chain mass, stabilized by iodine atom, is 285 Da (412-127 = 285) and the peak mass ratio 285:127 = 2.24 for ion *M-I* with m/z = 285 is the base.

In iodine-containing series (marked in green), the detachment and emission from the middle of chain of group -OC2F4 with formation of rearrangement ion I*+C(CF3)2 (with m/z 277) deserves attention.

Ion series of C₈F₁₇IO₂ MW:578 mass spectrum are shown in Fig. 19.

![Figure 19. Three ion series of mass spectrum C₈F₁₇IO₂ MW:578 Ion Trap (Polaris Q).](image)

As a result of increase in mass of iodine-stabilized chain by 166 Da, the intensity of molecular ion peak with m/z 578 (see Fig. 19) compared with the peak intensity of molecular ion with m/z 412 (see Fig. 18) decreases from 17% to 2%. Emission of one iodine atom in spectrum does not manifest itself. Iodine is either emitted together with fluorine atom (with formation of terminal oxaperfluorovinyl group) (series marked by blue), or as part of terminal group OC₂F₄I (series marked in red).

In «blue series» the rearrangements occur with formation of ions (m/z 313 and 147) with terminal fluorocarbonyl groups. When the ion with m/z 313 occurs, the oxaperfluorovinyl group of ion with m/z 413 is rearranged into a fluorocarbonyl group. Subsequent detachment of fluorocarbonyl group (ion with m/z 266), emissions of C₂F₄ and atom F, again culminate in formation of ions with fluorocarbonyl groups having m/z 147, 97 and 47.

Fig. 20 shows the ion series of mass spectrum C₈F₁₇IO₂ MW:578. The intensity of molecular ion peak, whose mass is 910 Da, is only 1%.
When the iodine atom is detached, the intensity of M-I ion peak (with m/z 783) is 1%. A decrease in intensity of M-I peak with increasing in molecular weight of M is the result of increasing in mass ratio (M -I:I) 783:127 = 6.2. M-I ion with m/z 783 either fragments with emissions of OC₂F₄, OC₃F₆ and CF₂, forming a series of perfluoroalkyl ions (series marked in red), or (after emission of CF(CF₃)CF(CF₃)CF₂OC₂F₄ group and detachment of two fluorine atoms) forms a rearrangement ion (m/z 479) with a terminal fluorocarbonyl group (series marked in blue). During fragmentation, the fluorocarbonyl group is lost (ion with m/z 266), and then, after emission of C₂F₄ and fluorine atom, the fluorocarbonyl group is re-formed (ion with m/z 147).

Chain breaks of +M-517 (with formation of ion IC₂F₄OCF₂+CF(CF₃) with m/z 393 (series marked in green) and +M-393 with additional emission of two fluorine atoms and formation of ion +C₃F₆OCF(CF₃)CF₂OCF(CF₃)FC=O with m/z 479 (series marked in blue) occur in the same chain link (see Fig. 20). The first fragmented iodine-containing ion IC₂F₄OCF₂+CF(CF₃) with m/z 393 fragments in two ways: both with detachment of OC₃F₆ and formation of base peak of ICF₂+CF₂ ion, and with emission of C₃F₆ and two fluorine atoms with formation of “parallel” series of iodine-containing ions.

**Conclusion**

Mass spectrum of compound, presented in the form of its ion series (with corresponding peak intensities, ion structures and fragmentation sequences) allows us to solve the same problems as
analysis of peaks of metastable ions. When analyzing metastable ions, a summary fragmentation scheme is usually considered, without isolating ion series and discussing their structures [5]. The series of ion fragmentation chains, presented in this report, correspond to mass spectra of compounds (with real intensities of their peaks). Consideration of primary ions and finite fragment ions compositions, as a rule, allows us to determine the number of ion series of mass spectrum. The ion series will branch if the resulting ion cannot be formed as a result of detachment of regular fragment group. When the ion is fragmented, along with regular detachment, a parallel detachment may also occur, which differs from regular one by a regular fragment group, unless this of course contradicts the composition and structure of this ion. As a result, this ion can move from one ion series to another.

Molecular ions do not appear in mass spectra of polyoxaperfluoroalkanes, regardless of their molecular weight. During fragmentation of symmetric molecules with MW 570-702 Da, the detachments of terminal radicals (with or without an oxygen atom) occurs in two central groups: O(CFCF₃)₂O. Fragmentation of polyoxaperfluoroalkane with two different terminal groups CF₃CF₂CF₂ and C(CF₃)₃ begins with CF₃CF₂CF₂ group and terminates with C(CF₃)₃ group.

In mass spectrum of symmetric polyoxaperfluoroalkane C₃₄F₇₀O₁₀ (MW 1898) the first fixed peak is the peak of ion with a mass of 667. Since C₃₄F₇₀O₁₀ molecule is symmetric, there is reason to believe that decay of M⁺. occurs with emission of neutral molecule [OCF(CF₃)CF₂OCF(CF₃)-] with a mass of 564 Da from central part of molecular radical cation. The resulting symmetric radical cation [C₂₄F₅₀O₆]⁺ with m/z 1334 decays into ion and radical with the same masses equal to 667Da.

In mass spectra of polyoxaperfluoroalkanes with number of oxygen atoms of 2 or more, as a rule, there are two ion series. One of them leads to fluoroalkyl ions, and the other - to ions with fluorocarbonyl group.

In mass spectrum of oxaperfluoroalkyl chloride with one oxygen atom ClC₂F₄O(CF₂)₆CF₃ (see Fig. 9), the additional (third) ionic series of perfluoroallyl ions appears. After primary detachment of CICF₂CF₂ radical, as well as two fluorine atoms from opposite terminal group, the ion +OC₅F₁₀CF=CF₂ with m/z 347 is formed. It fragments in two ways, both with formation of series of perfluoroallyl ions (with m/z 281,231,181 and 131), and ions with fluorocarbonyl group (with 197, 147, 97 and 47). The occurrence of two series of fragmentation of ion +CF₂CF₂FC=O with m/z 347 is a result of its rearrangement into O=CFCF₂C₄F₈⁺CF₂ ion.

In mass spectra of polyoxaperfluoroalkanes and polyoxaperfluoroalkils with terminal halide atoms, due to presence of oxygen atoms in the chain, the formation of series of perfluoroallyl ions is impossible.
However, CF$_2$=CF$^+$CF$_2$ ion with m/z 131 is formed. It occurs as a result of oxygen atom detachment from ion $^+\text{CF}_2\text{CF}_2\text{FC}=\text{O}$ with m/z 147, or as a result of oxygen atom detachment from ion $^+\text{C}_3\text{F}_6$.

In spectrum of $\text{C}_{14}\text{F}_{29}\text{IO}_4$ the rearrangement of iodine-containing ion with m/z 393 leads to formation of second iodine-containing series of ions.

In spectra of polyoxaperfluoroalkyl bromides the primary detachments of two radicals with different masses lead to formation of two series of bromine-containing ions.

Two fragmentation pathways for $^+\text{M}$ (C$_{14}$Br$_2$F$_{28}$O$_4$), leading to formation of two series of bromine-containing ions, are either the result of difference in excitation energies $^+\text{M}$, or the result of removal of two electrons from molecule that are topologically different with respect to bromine atom.

Presented examples of ionic series of polyoxaperfluoroalkanes and polyoxaperfluoroalkyl with terminal halide atoms allow us to conclude about a certain «foreignness» of oxygen atoms in perfluoroalkyl chain. During fragmentation, the oxygen atom either moves to terminal fluorocarbonyl group, or detaches as part of terminal group, or is emitted from the middle of chain (in the form of atom O, molecule O=CF$_2$, OCF$_2$CF$_2$ with formation of perfluoroalkyl and haloperfluoroalkyl ions. As a result of such detachments, the fragmented ions $^+\text{C}_n\text{F}_{2n+1}$ are formed, with a number of fluorine atoms greater than in original molecule.

When fragmenting [(CF$_3$)$_2$CFOC$_2$F$_4$I]$^+$ (see Fig. 18) the extraordinary emission of OC$_2$F$_4$ with formation of rearrangement ion (CF$_3$)$_2^+$CFI deserves attention.

In mass spectra of polyoxaperfluoroalkyl chlorides and bromides, the primary peak is M-F peak, and the peaks of molecular cation radicals do not appear. Compared with other halides, the terminal iodine atom in mass spectra of polyoxaalkyl iodides contributes to maximum stability of molecular ion, especially if the linear chain of compound is not very large. As the molecular weight increases with increasing in chain mass by more than 3 times (compared to mass of terminal iodine atom), the intensity of peaks of molecular iodide ions decreases.

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