SELECTIVE DETECTION IN GAS CHROMATOGRAPHY

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This technical report has been reviewed and is approved for publication.

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The most popular method of separating a complex hydrocarbon mixture, such as jet propulsion fuel, into its individual components is that of high resolution gas chromatography (HRGC) using a flame ionization detector (FID). However, neither the efficiency of HRGC or the sensitivity of the FID lend any information as to the identity of the individual components.

This report looks at 24 different gas chromatographic detectors that respond selectively to heteroatom compounds such as those containing S, N, O, P, Halogens, and some metals. Recommendations for the use of these detectors for analyses of jet propulsion fuels are made.
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INTRODUCTION

High-resolution gas chromatography (HRGC) is capable of separating a very complex hydrocarbon liquid such as jet propulsion fuel into nearly all of its individual components. However, there is no guarantee that what appears to be a single chromatographic peak is in fact a single fuel component. The method of detecting these HRGC eluents is very often the flame ionization detector (FID), a very sensitive and universally responding hydrocarbon detector.

Neither the separating power of HRGC, or the sensitivity of the FID, however, lend information as to the identity of the eluting component. The need to specifically identify HRGC eluents, especially those containing heteroatoms such as N, S, P, O, halogens and some metals has been well-documented (5, 11).

The purpose of this report is to review the progress in the area of selective gas chromatographic detectors and to recommend selective detectors for use in the HRGC analysis of jet propulsion fuels.

This report will present 24 different selective gas chromatographic detectors. These detectors, while not all directly applicable to the HRGC analysis of jet propulsion fuel, all show the concept of selective detection in gas chromatography.

The detectors presented herein range in complexity from simple modifications made to existing universal detectors or specific analyzers to detectors that are scaled down versions of high-technology, high cost spectroscopic techniques.

The applicability of these detectors to jet propulsion fuel analyses will be determined from a comparison of their complexity and cost versus their effectiveness as high-resolution gas chromatographic detectors.
The most common method of chromatographing complex hydrocarbon mixtures such as jet propulsion fuel is with a high resolution gas chromatograph using glass capillary or fused silica columns. The sample is injected into a heated injector zone where it is vaporized. A large portion of this vaporized sample is vented, and the rest is routed into the column. Once inside the column, the sample partitions into a stationary liquid phase that coats the inside of the column. Different fuel components migrate down the column at different rates due to their individual partition coefficients with this stationary phase.

Once eluted from the column, compounds are then detected with any one of a number of available gas chromatographic detectors. The 24 detectors examined in this report are separated into four distinct categories: ionization detectors, absorption detectors, emission detectors, and conductivity detectors.

A. IONIZATION DETECTORS

Virtually any type of energy (electronic, radioactive, heat, etc.) can be used to ionize GC eluents. Selectivity is realized by certain compounds being susceptible to the type of ionization method used, or by ionization products being characteristic of the parent molecules.

1. Mass Spectral (MS)

Compact mass selective GC detectors are now commonly available for use. The Ion Trap Detector (Finnigan Mat) is an excellent example of one such detector.
The eluents of a GC are transferred to the detector via a heated fused silica transfer line where they are ionized by electron impact. The ions are then ejected from the ion storage region sequentially from low to high mass and then detected by an electron multiplier. The resultant mass spectrum for any given mass-to-charge ratio can then be matched to existing mass spectrum in a library by computer searching techniques. This detector is capable of universal mass spectrum scanning and specific mass spectrum searching. (See Fig. 1.)

2. Fourier Transform Mass Spectral (FTMS)

The use of Fourier Transformation with mass spectral data greatly enhances the mass resolution of GC/MS data. For example, the GC-FTMS system referenced here is capable of collecting 8,192 data points every 100 ms. This fact combined with the background separation capability of Fourier Transformation makes GC-FTMS an extremely powerful GC peak identification technique (2).

3. Oxygen-Flame Ionization (O-FID)

The Oxygen Flame Ionization Detector obtains selectivity for O2 containing compounds by utilizing two microreactors in line between the column outlet and the flame. The first, a cracking reactor operated at 1500-K, forms CO from the O2 compounds. The second reactor operated at 620-K, hydrates the CO to form CH4. The CH4 is then detected by normal flame ionization (20). (See Fig. 2.)

4. Hydrogen Atmosphere Flame Ionization (HAFID)

The Hydrogen Atmosphere Flame Ionization Detector realizes selectivity for Si, P, Cl and Fe by merely altering the type and flow rates of gases to the detector. The normal 300 ml/min air and 30 ml/min H2 is changed to 130 ml/min O2 and 1600 ml/min H2. This results in suppressed responses to normal hydrocarbons and selective responses to the above-mentioned compounds (19).
Figure 3 Procedure for compound identification using the ITD.

Table 3

<table>
<thead>
<tr>
<th>Compound class</th>
<th>Characteristic mass</th>
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<tbody>
<tr>
<td>Paraffins</td>
<td>43, 57, 71, 85</td>
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<tr>
<td>Aliphatic carboxylic acid methyl esters</td>
<td>74, 87</td>
</tr>
<tr>
<td>Alkenes</td>
<td>41, 55, 69</td>
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<td>Trittersanes</td>
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<td>Steranes</td>
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<td>Silioxanes</td>
<td>73</td>
</tr>
<tr>
<td>PCBs</td>
<td>256, 290, 324, 358, 392</td>
</tr>
</tbody>
</table>

*For a more detailed list of masses common for compound classes, see Ref. 3.

FIG 1: Identification of chlorocresol by GC/MS using a reference spectrum. (Reproduced by permission from ref. 1, Copyright International Scientific Communications.)
FIG 2: Selective detection of 02 compounds in starting fluid by GC/O-FID. (Reproduced by permission from ref. 20, Copyright Elsevier Science Publishers B.V.)
5. Thermionic Specific (TSD)

The thermionic emission of charged particles from the surface of a hot bead consisting of alkali metal embedded in a ceramic matrix provides highly specific responses to compounds containing N and P as well as halogenated compounds. To a small extent the detector is selective to aromatic compounds in hydrocarbon fuel samples (17) (22).

6. Ion Mobility (IMD)

Ion Mobility Spectrometry is a method closely related to the Electron Capture Detector where organic compounds are ionized by charge transfer reactions in the presence of a constant electric field gradient. Product ions drift down this field at different rates, dependent on mass, charge and ion size. By doping the detector with oxygen, increased selectivity and sensitivity are gained for halogenated compounds. (13).

7. Electron Capture Detector (ECD)

The constant signal of a radioactive electron source, commonly Np-63, in an inert gas flow is interrupted by the introduction of molecules with "capture free" electrons, such as halogenated compounds, which generate a negative response. The eluents are ionized but the response is read as the reduction of the ion current (5).

8. Photoionization (PID)

The Photoionization Detector selectively detects gases such as H2S, NH3, AsH3, and I2. A U.V. lamp irradiates the GC eluents. Any compound with an ionization potential less than the U.V. lamp will be ionized. The resultant ion current is measured by a sensitive electrometer system (11).
9. Catalytic Ionization

When organic compounds are oxidized on the surface of a heated platinum filament only certain compound types yield an ion current comparable to that of the flame ionization detector. Specifically, these compounds are highly branched hydrocarbons such as alkanes and alcohols (17).

B. ABSORPTION DETECTORS

Comparing the absorption characteristics of an eluting GC compound to known standards in some region of the electromagnetic spectrum truly defines the nature of that compound. This technique is used either by scanning a specific region of the spectrum or at a set wavelength specific to the compound of interest.

1. Infrared (IR)

Eluting GC components are trapped or isolated in a GC/IR cell through which a dispersive I.R. spectrophotometer provides an I.R. spectrum of the component. This is not a continuous analysis, and is limited by the I.R. scan time (3). (See Fig.3.)

2. Fourier Transform Infrared (FTIR)

By use of a interferometric system allowing the infrared to scan eluting chromatographic compounds at a rate of up to 10 scans per second, the Fourier Transform Infrared system can produce a spectrum of a single chromatographic peak while it is eluting. The spectrum is then compared to a library of infrared spectrum for peak identification. Absorbance versus time can also be plotted to produce a more traditional chromatogram (4).
Figure 12 Chromatogram of a fragrance mixture, recorded as in Figure 1. Chromatographic conditions are the same as for Figure 1, except the oven was programmed from 90°C to 220°C at 10°C/min.

Figure 13 Infrared chromatograms of the fragrance mixture in Figure 11, obtained on an IBM IR/85 GC/IR system.

Figure 14a) Eucalyptol and b) eugenol are infrared spectra in transmittance of peaks 1 and 2 (Figures 12 and 13), respectively.

**FIG 3:** Selective detection of eucalyptol and eugenol by GC/IR.
(Reproduced by permission from ref. 4, Copyright International Scientific Communications.)
3. Atomic Absorption Spectrometry (AAS)

Through the simple interfacing of a high resolution gas chromatograph to an atomic absorption spectrophotometer organomercury (II) compounds can be selectively analyzed at trace levels in organic mixtures. The background interference from the organic molecules is minimal for two reasons. The AAS is not particularly sensitive to organic molecules and these molecules are mostly cracked in the 600 Deg.-C decomposition furnace used to convert the organomercurials to elemental mercury (9).

4. Ultraviolet (UV)

GC column eluents are directed to a sample cell with a quartz window. Through this window, the sample is irradiated by U.V. light at a fixed wavelength. The choice of this wavelength determines the compound-type sensitivity. Typically, a setting of 250 nm renders the detector very sensitive to aromatic hydrocarbons and insensitive to aliphatic or naphthenic species (5). (See Fig.4.)

C. EMISSION DETECTORS

Eluting GC compounds are converted to an excited state and allowed to relax yielding characteristic emissions. These detectors combine the advantages of ionization and absorption detectors in that the type and amount of energy applied is controllable and the emissions are characteristic in the electromagnetic spectrum.

1. Rotationally Cooled Laser-Induced Fluorescence (RC-LIF)

GC eluents are excited by a high-intensity laser beam and caused to fluoresce. By taking advantage of the fact that any compound will fluoresce at least one wavelength unique to that compound, RC-LIF is virtually unsurpassed in selectivity. However, at present, the system is restricted to scanning for a particular compound at its unique wavelength (8). (See Fig.5.)
FIG 4: Selective detection of aromatics in a jet propulsion fuel by GC/UV. (25)
Figure 2. Chromatograms of an equimolar mixture of naphthalene, α-methylnaphthalene, and β-methylnaphthalene. Excitation wavelengths used for each injection are indicated.

Figure 1. Diagram of the RC-LIF/GC apparatus. The portion enclosed by the dotted line forms a simple GC. Temperature variation along the column is shown below.

Figure 3. Chromatograms of a Wilmington Crude Oil sample: (a-d) chromatograms of the crude oil excited at the wavelengths indicated; (e) a laboratory prepared mixture of naphthalene, α-methylnaphthalene, and β-methylnaphthalene at the concentrations determined for the crude oil sample.

FIG 5: Selective detection of naphthalene isomers by GC/RC-LIF. (Reproduced by permission from ref. 8, Copyright American Chemical Society.)
Fig. 11. Separation of a monochlorobiphenyl isomer mixture and subsequent detection by a helium micro-mass plasma. Peak concentrations are (1) 184 μg g⁻¹ ortho; (2) 325 μg g⁻¹ para; and (3) 448 μg g⁻¹ meta. Chlorine response is at 481.0 nm with 2 μl injections being made of the mixture. The column is the same as that used in Fig. 12. Note absence of matrix response compared to Fig. 12.

FIG 6: Selective detection of monochlorobiphenyl isomers by GC/MIP. (Reproduced by permission from ref. 10, Copyright Pergamon Press, Inc.)
2. Microwave-Induced Plasma (MIP)

The use of emission spectroscopy by an atmospheric pressure microwave induced helium plasma (MIP) has been proven to be a specific detection technique for N, P, S, Cl, Br, C and Fe. A small portion of the plasma image, slightly off-center of the microwave cavity, was found to exhibit intense elemental emissions for the compounds mentioned while the rest of the plasma showed no emissions for those compounds. By varying the helium flow and power to the microwave cavity, optimum responses for each of the above mentioned compounds were obtained (10). (See Fig.6.)

3. Oxygen-Inductively Coupled Argon Plasma (O-ICAP)

Use of inductively coupled argon plasma to excite non-resonance atomic emissions from oxygen in the red and near infrared regions is used to quantitatively and selectively determine oxygen in permanent gas mixtures and volatile organics (6).

4. Chemiluminescent (CLD)

The Chemiluminescent Detector provides very sensitive, very selective responses to bound nitrogen in hydrocarbon matrices. The sample is combusted to form nitric oxide. The nitric oxide is detected photometrically by its reaction with ozone to form the chemiluminescent species NO2*. Similarly, the detector can be used to specifically detect volatile polyhalogenated hydrocarbons by utilizing the halogenated compounds' reaction with sodium vapors and the resultant chemiluminescent sodium species (14) (23) (11).

5. Glow Discharge Detector (GDD)

A quartz tube, using axially mounted platinum wire electrodes, provides the spectrum for this optical emission detector. Use of a very stable power supply is necessary to maintain the tuning at the emission wavelength of choice. The detector is used primarily to indicate the presence of halogens, S, P, C and metals in GC effluents (7) (15).
6. Atmospheric Pressure Active Nitrogen (APAN)

Atmospheric Pressure Active Nitrogen sources have been used to selectively detect organo- Hg, Pb, and Sn as trace GC eluents. These compounds react with metastable atoms and molecules of nitrogen from the active nitrogen source. In these reactions the fragmentation and excitation of species lead to characteristic atomic and/or molecular emissions (16).

7. Flame Photometric Detector (FPD)

Emissions from a flame burning chromatographic eluents are viewed through appropriate bandpass filters by a photomultiplier. The maxima of the characteristic filters are 394 nm for sulfur and 526 nm for phosphorus. Filters of other wavelengths are available for other compound types (5) (17).

8. Thermal Energy Analyzer (TEA)

The Thermal Energy Analyzer can be used as a GC detector to selectively detect N-nitroso compounds in organic matrices. The sample is pyrolyzed and converted to nitrosyl and organic radicals. The organic fragments either decompose or rearrange to stable forms. The nitrosyl radical is converted to NO$_2^*$ through ozonolysis. The NO$_2^*$ decays rapidly with an emission in the near infrared region of the spectrum (18) (21).

D. CONDUCTIVITY DETECTORS

By converting GC eluents to the appropriate inorganic forms their specific electrical nature can be detected. This detection can be accomplished either by direct measurement of electrolytic conductivity or by titration with electrolytically generated reactive ions.
Figure 2 Block diagram of bipolar pulse, differential conductivity circuit.

Figure 3 Cross section of microsector and differentials.

Figure 18 Arochlor 1254 in transformer oil, extracted with methanol.

FIG 7: Selective detection of chlorinated compounds in transformer oil by GC/HECD. (Reproduced by permission from ref. 12, Copyright Preston Publications, Inc.)
1. Hall Electrolytic Conductivity (HECD)

The Hall Electrolytic Conductivity Detector can be operated with different
modes of selectivity for halogen-, sulfur-, carbon-, nitrogen-, or nitrosamine-
containing compounds. The species of interest are converted to stable
inorganic forms by combustion or hydrogenolysis. Upon dissolving these
converted species in water or alcohol they can be detected by measuring the
resultant electrolytic conductivity (12). (See Fig. 7.)

2. Microcoulometry (MC)

Microcoulometry has been successfully used as a specific GC detector for S,
N and Cl. The elements in question are converted to the inorganic forms SO2,
NH3, and HCL respectively. The species are then titrated by electrolytic
generation of reactive ions. For example, SO2 is titrated with
electrogenerated I2 via a redox reaction. The quantity (i.e. microcoulombs) of
electricity required for this generation is the area under the GC-MC peak.
GC-MC peaks are broad with delayed response times and some tailing (11).

3. Molecular Weight Detector

By directing GC eluents to one of two carrier gas flow paths which are legs
of a Wheatstone Bridge the density changes caused by the eluents can be used as
a detector signal. While this method is not specific to eluent types, the use
of two of these systems in parallel, each using a different carrier gas, can be
used to determine molecular weights of eluting compounds. The molecular weight
is calculated by the simultaneous solution of the gas density balance equations
for the two systems (17). (See Fig. 8.)

Table 1 is a listing of the detectors covered in this report by their types
of selectivity. Table 2 lists performance data for some of the detectors
covered in this report as available in the references listed herein.
The basis for the method can be seen by examining the basic equation for the gas density balance:

\[ \rho_x = k \left( \frac{N_{x}}{N_{x} - N_{CG}} \right) \]  

(1)

where \( \rho_x = \frac{W}{V} = \frac{W}{A} \) and \( A \) is the peak area, \( N_W \) is molecular weight, and \( CG \) denotes carrier gas. For a system with two gas density balances with different carrier gases,

\[ \rho_x = \frac{W_x}{A_x} = k_x \left( \frac{N_{x}}{N_{x} - N_{CG}} \right) \]  

(2)

The two weights are related to the total weight by the split flow ratio:

\[ W_1 = C_1 W_{\text{total}} \]

(3)

\[ W_2 = C_2 W_{\text{total}} \]

where \( C_1 \) is the fraction of sample through detector 1 and \( C_1 + C_2 = 1 \).

and \( C_2 \) is the fraction of sample through detector 2

and \( C_1 + C_2 = 1 \).

\[ W_{\text{total}} = \frac{W_1}{C_1} = \frac{W_2}{C_2} \]  

(4)

Substituting Equation 2 in Equation 4

\[ W_{\text{total}} = \frac{k A_x}{C_1} \left( \frac{N_{x}}{N_{x} - N_{CG}} \right) \]  

(5)

Simplifying, we obtain

\[ \frac{k_1 A_1}{C_1} = \frac{N_{x}}{N_{x} - N_{CG}} \]

(6)

where \( \frac{k_1 A_1}{C_1} \) can be determined after which \( Y \) is determined from Equation 7, and is then used in Equation 9. Typical results are shown in Table 1.

\[ k_1 A_1 \]

(7)

\[ \frac{N_{x}}{N_{x} - N_{CG}} \]

(8)

\[ \frac{N_{x} - N_{CG}}{N_{x}} \]

(9)

Equation 6 is first used to determine \( X \) using a standard material whose molecular weight is known. Substituting for \( N_W \), and the measured areas \( A_1 \) and \( A_2 \), \( X \) can be determined after which \( Y \) is determined from Equation 7, and is then used in Equation 9. Typical results are shown in Table 1.

Figure 3 (Ref. 3). Chemalytics molecular weight chromatograph splits sample between two columns, uses two carrier gases.

FIG 8: Molecular Weight Detector system and density balance equations.

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<thead>
<tr>
<th>DETECTOR</th>
<th>TYPE</th>
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<td>Mass Spectral Ionization</td>
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<tr>
<td>Infrared Absorption</td>
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<td></td>
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<tr>
<td>Rotationally Cooled-Emission Laser Induced Fluorescence</td>
<td>Various</td>
<td></td>
</tr>
<tr>
<td>Molecular Weight Conductivity</td>
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<td></td>
</tr>
<tr>
<td>Ion Mobility Ionization</td>
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<td>Chemiluminescent Emission</td>
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<td>Thermionic Specific Ionization</td>
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<td>Microcoulometry Conductivity</td>
<td>N,S,X</td>
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<td>Hall Electrolytic Conductivity</td>
<td>N,S,X</td>
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<tr>
<td>Microwave Induced Plasma Emission</td>
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<td>Glow Discharge Emission</td>
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<td>0-Flame Ionization Ionization</td>
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<td>0-Inductively Coupled Argon Plasma Emission</td>
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SECTION III
CONCLUSIONS

A chromatographer with a specific G.C. detection problem clearly has two choices, either to attack the problem with a high-technology detector such as the GC-FTMS, or to approach the problem with a novel device such as the O-FID.

High technology, high cost detectors such as the GC-FTMS and GC-FTIR have advantages over even the most novel element specific detectors in that they are "universally specific." However, one cannot afford to overlook the phenomenal achievements made over the last two decades in the development of element specific detectors.

For the purpose of analyzing heteroatom constituents of jet propulsion fuels, the following detectors are recommended: the O-FID for the detection of oxygen-containing compounds, the FPD for the detection of phosphorous and sulfur-containing compounds, either the CLD or TSD for the detection of nitrogen-containing compounds. A system containing either MSD, FTIR, or both capabilities is also highly recommended for these analyses.
SECTION IV
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