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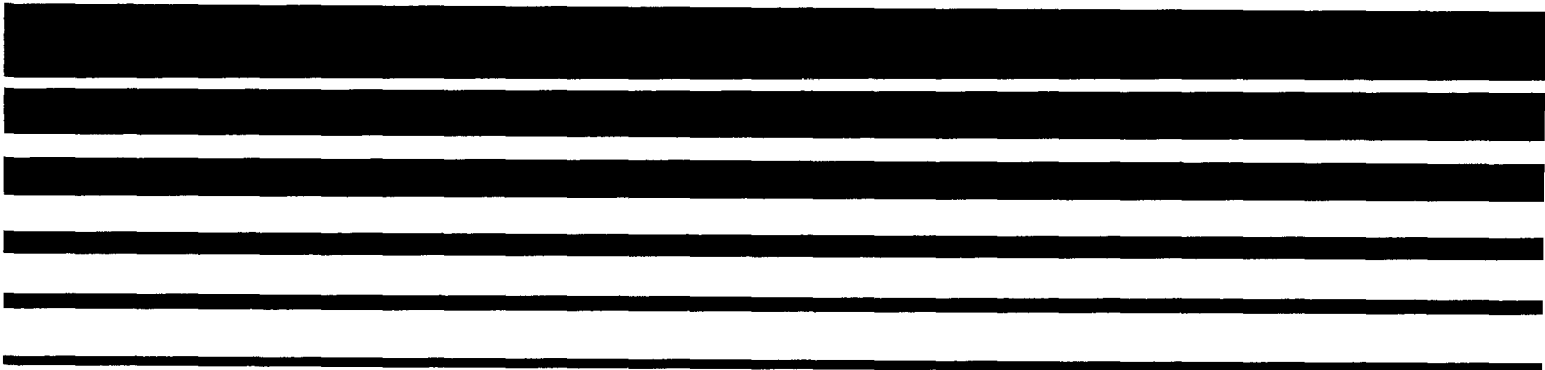
Office of Mobile Source Air Pollution Control
Emission Control Technology Division
2565 Plymouth Road
Ann Arbor, Michigan 48105

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Air



Calculation of Emissions and Fuel Economy When Using Alternate Fuels



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Calculation of Emissions and Fuel Economy When Using Alternate Fuels

by

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FOREWORD

This Work Assignment was initiated by the Control Technology Assessment and Characterization Branch, Environmental Protection Agency, 2565 Plymouth Road, Ann Arbor, Michigan 48105. The effort on which this report is based was accomplished by the Department of Emissions Research of Southwest Research Institute, 6220 Culebra Road, San Antonio, Texas 78284. This program, authorized by Work Assignment 8 under Contract 68-03-3073, was initiated May 26, 1982 and was completed September 28, 1982. The program was identified within Southwest Research Institute as Project 05-6619-008.

This Work Assignment was conducted by Mr. Charles Urban, Project Leader. Mr. Charles Hare was Project Manager and was involved in the initial technical and fiscal negotiations and subsequent major program decisions. The EPA Project Officer was Mr. Robert J. Garbe of the Technical Support Staff, Environmental Protection Agency.

ABSTRACT

This report provides methods for the calculation of vehicle emissions and fuel consumption when nonstandard fuels are used. Methods of analysis, required for evaluation of alternate fuels, are included by reference or as Appendices to this report.

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I. INTRODUCTION

The objective of this Work Assignment was to compile and document procedures to follow when calculating and reporting exhaust or evaporative emissions data and carbon balance fuel economy data from vehicles operating on hydrogen and carbon containing fuels other than commercial gasoline and Diesel fuel.

This report describes calculational procedures for use in light-duty applications when using alternate fuels. In addition to the calculational changes that are required to the light-duty certification procedure, some equipment and instrumentation changes may also be required. Discussion of these equipment and instrumentation requirements is also included in this report.

The composition of the evaporative and the exhaust organic matter with alternate fuels generally differs from the composition of the hydrocarbons used to establish the HC emissions standard. The relationship between the organic matter value for an alternate fuel and the HC emissions value for a standard gasoline or diesel fuel have not been established. Therefore, no procedure is included for the calculation of a single composite value for organic matter when alternate fuels are used.

The format used for mathematical equations in this report is in accordance with that used in the Code of Federal Regulation.^{(1)*} Computations are to be performed in accordance with the IBM hierarchy of operations.

Although specific application to the heavy-duty transient procedure is not included, the procedures given are generally applicable. The methods given can be fairly readily adapted for use with the heavy-duty transient procedure.

*Numbers in parentheses designate references at the end of this report.

II. EQUIPMENT, INSTRUMENTATION AND ANALYTICAL METHODS

Fuels having higher hydrogen-to-carbon ratios produce more water and less CO₂ for equal energy output. Therefore, a higher than normal CVS dilution flow rate could be required to keep water condensation from occurring. A higher CVS flow rate (CO₂ emissions would also be directly affected by lower CO₂ production) could result in emissions concentrations that are lower than the prescribed accuracies of the normally used emissions instruments. Therefore, the CVS flow rate and the exhaust emissions instrumentation are essential considerations when alternate fuels are to be evaluated.

A. CVS Dilution Requirements

The CVS flow capacity of 300 to 350 scfm that is designated in the light-duty certification procedure as being adequate, is based on currently formulated gasoline and Diesel fuels.⁽¹⁾ Fuels containing a higher hydrogen-to-carbon ratio produce more water and can require a higher CVS dilution flow rate to keep condensation of water from occurring in the sample bags. Since the currently specified CVS flow rate generally prevents water condensation from occurring, when using gasoline or Diesel fuel, the CVS flow rate for an alternate fuel that is sufficient for most light-duty vehicles can be estimated as follows:

$$\bullet \text{ ACVSFR} = (\text{AFFH}/\text{ALHV}) \times 5 \times 10^7$$

Where:

ACVSFR = CVS flow rate for alternate fuel that is
sufficient for most vehicles, cfm

AFFH = Fuel fraction hydrogen of the alternate fuel

ALHV = Lower heating value of the alternate fuel, Btu/lb

The resulting flow rate (ACVSFR), however, is often much higher than is actually required, especially for smaller, fuel efficient vehicles. Therefore, the determination of the actual required CVS flow rate is given as follows:

$$\bullet \text{ RCVSFR} = 16.6 \times \text{AMPH} \times \text{FDEN} \times \text{FFH} / (\text{MPG} \times (\text{SSH} - \text{TSH}))$$

Where:

RCVSFR = Required CVS flow rate, scfm

AMPH = Average speed, mph (16 minimum for the
cycles of the FTP)

FDEN = Density of the fuel, g/ml

FFH = Fuel fraction hydrogen, mass/mass

MPG = Fuel economy, mpg

SSH = Specific humidity of saturated air at
minimum sample bag temperature, lb/lb

TSH = Actual specific humidity of the dilution air
at test conditions, lb/lb

Note: Derivation of RCVSRF is given in Appendix A-1.

Combining the provisions in the ACVSFR and the RCVSFR equations, and making several assumptions to enable simplification, results in the following equation for estimation of the required CVS flow rate for the FTP:

$$ECVSFR = 45,000 \times FDEN \times FFH / MPG$$

When the fuel economy for the vehicle is not known, it can be estimated using the fuel economy for a similar vehicle operating on certification fuel multiplied by the ratio of the lower heating values per unit volume:

$$AMPG \approx CPMG \times CLHV / ALHV$$

Where:

AMPG = Fuel economy with the alternate fuel, mpg

CPMG = Fuel economy with certification fuel, mpg

CLHV = Lower heating value of certification fuel (18,300 Btu/lb) ⁽²⁾

ALHV = Lower heating value of the alternate fuel

B. Emissions Instrumentation

When fuels with high hydrogen-to-carbon ratios are used, emissions instrumentation having lower ranges may be required. Unless the emission rates are known, however, the necessary instrument ranges can not be pre-determined. Therefore, the recommendation is that applicability of ranges be determined from actual measurement of emissions at the onset of testing.

CO₂ emission concentrations, however, can be estimated using the CVS flowrate, the carbon fraction of the fuel, and the fuel consumption on a weight basis. The equation, derived in Appendix A-2, is as follows:

$$CO_2\% = (451 \times AMPH \times FDEN) / (FE \times CVSFR)$$

Where:

CO₂% = CO₂ concentration in the Dilute Exhaust Sample, Percent

AMPH = Average Speed of the Cycle, mph

FDEN = Fuel Density, g/ml

FE = Cycle Fuel Economy, mpg

CVSFR = Flowrate of the CVS, scfm

C. Analytical Methods

In several of the calculations required for alternate fuels, additional analyses of the fuel or the emissions are specified or recommended. Some analytical methods are listed on the following page.

| <u>Method</u> | <u>Source</u> |
|---|-----------------------------|
| Determination of FID Response Factor | Appendix A-3 |
| Measurement of Aldehydes and Ketones | Appendix A-4 ⁽³⁾ |
| Measurement of Individual Hydrocarbons | Reference 3 |
| Methane Measurement Using Gas Chromatograph | SAE J1151a ⁽⁴⁾ |

Additional methods for special applications are listed as follows:

| <u>Method</u> | <u>Source</u> |
|---------------------------------------|-----------------------------|
| Measurement of Methanol | Appendix A-5 ⁽⁵⁾ |
| Measurement of Ethanol | Appendix A-6 ⁽⁶⁾ |
| Measurement of Tertiary Butyl Alcohol | Reference 6 |

III. FUEL COMPOSITION

In the procedure for certification of light-duty vehicles, a specified hydrogen-to-carbon ratio and density are assumed to be applicable for all petroleum based fuels meeting the prescribed specifications. When using alternate fuels, however, the differences in fuel composition and density are sufficiently large to make it essential that the actual values be determined and used.

The calculations of emissions and fuel economy, for alternate fuels, require the use of the weight ratios of carbon, hydrogen, oxygen, and the sum of the other elements in the fuel. For the calculation of fuel economy, the density of the fuel is also required.

For blends of known composition, the weight ratios of the constituents can either be measured or calculated. For fuels of unknown composition, however, determination of the elemental composition is essential. Due to the uncertainties involved when various liquids are blended together, it is recommended that fuel density always be determined by actual measurement (unless the effects of blending on density are specifically known for the specific blended fuel).

For gaseous fuels, such as propane, a significant quantity of impurities (e.g., CO_2) may be present in the fuel which would have an impact on the emissions and fuel economy calculations. Therefore, it is also important to determine the weight ratios of carbon, hydrogen, oxygen, and the sum of the other elements when gaseous fuels are used.

A. Blends of Known Composition

When blending a fuel of known composition, the elemental weight ratios can be determined by calculation. The method will be demonstrated using a blend of a gasoline, meeting the certification fuel specifications, and methanol (assuming such a blend is achieved).

For gasoline, a hydrogen-to-carbon ratio (HCR) of 1.85 to 1 can be assumed. This results in fuel carbon and hydrogen fractions of weight (GFC and GFH) of 0.866 and 0.134, respectively. Methanol has an MFC and MFH of 0.375 and 0.126, respectively. The remaining component in methanol is oxygen, with an MFO of 0.499. The resulting carbon, hydrogen, and oxygen fractions in a blend of gasoline and methanol are on the following page.

$$\text{BFX} = ((\text{Wt}\% \text{ Gas}) \times \text{GFX} + (\text{Wt}\% \text{ Meth}) \times \text{MFX}) / 100\%$$

$$\text{Where: Wt}\% \text{ Gas} = \frac{(\text{Vol.}\% \text{ Gas}) \times \text{GDEN} \times 100}{(\text{Vol.}\% \text{ Gas}) \times \text{GDEN} + (\text{Vol.}\% \text{ Meth}) \times \text{MDEN}}$$

B, G, & M = Blend, Gasoline and Methanol

X = Carbon, hydrogen or oxygen

DEN = Density, g/ml

Assume 90% Gasoline and 10% Methanol by Weight:

$$\text{BFC} = (90 \times 0.866 + 10 \times 0.375) / 100 = 0.817$$

$$\text{BFH} = (90 \times 0.134 + 10 \times 0.126) / 100 = 0.133$$

$$\text{BFO} = (90 \times 0.000 + 10 \times 0.499) / 100 = 0.050$$

The elemental fractions in a blended fuel can also be determined by elemental analyses of the fuel blend. Methods for such analyses are discussed in the following Section III.B. Due to the uncertainties involved in blending fuels, it is recommended that generally the density of the resultant fuel blend be determined by actual measurement.

B. Fuels of Unknown Composition

For fuels of unknown composition, the elemental fractions can be determined only by actual analysis. The carbon, hydrogen and oxygen values, and occasionally the nitrogen value, are required. Analysis of liquid fuels for these elements is extremely difficult, and standard ASTM methods are not currently available. It is recommended that such analysis be conducted only by a laboratory known to have the necessary expertise.

There does not appear to be general agreement on the best methods of analysis for these elements. The commercially available C, H, N, O and S analyzers were not specifically designed for, and do not generally appear to provide, satisfactory analysis of liquid fuels. Therefore, such analyzers are not a simple solution to the measurement problem and their use should be approached with caution.

Some example methods of elemental analysis for liquid fuels are as follows: Carbon and hydrogen have been satisfactorily measured using an elemental analyzer, or gravimetrically using a modified ASTM D-3178 method. Nitrogen has been satisfactorily measured using pyrochemiluminescence. Oxygen has been satisfactorily measured using neutron activation analysis. In all cases, it appears that a very high level of expertise is essential to obtaining satisfactory analysis of these elements in liquid fuels.

For general guidance purposes, the accuracies of the analyses should be within the following limits:

| Element | Analysis Accuracy as: | |
|--------------|---------------------------------|----------------------------------|
| | % of Value ^a | % of Fuel ^a |
| Carbon | ±0.5 | ±0.3 |
| Hydrogen | ±0.5 | ±0.3 |
| Oxygen | ±(0.2/Value ^b + 0.3) | ±(0.2 + 0.3×Value ^b) |
| Nitrogen | ±(0.2/Value ^b + 0.3) | ±(0.2 + 0.3×Value ^b) |
| Total of All | -- | 1.0 |

^aUse the analysis accuracy having the lowest absolute number

^bValue as a weight fraction of total fuel

The limits given represent the extremes; an effort should be made to obtain better analysis accuracies than these extreme limits given.

The total of the weight fractions of the C, H, O and N in the fuel may not add up to a total of 1.000. Computer inputs include the fuel fraction of C, H, O and other elements. If the total for C, H, O and N is less than 1.000, add the difference to the value for other elements. If the total for C, H, O and N is greater than 1.000, subtract the difference from the value for oxygen (O) or the value for other elements, as appropriate.

The acceptable level of sulfur is generally sufficiently low so that the amount of sulfur present does not significantly affect the standard emissions and fuel consumption calculations. Amount of sulfur in the fuel, however, is generally considered to be of sufficient importance to justify determination of sulfur. Depending on the type of fuel involved and its source, it may also be worthwhile to include additional analyses, such as ash content, gum, and some additional specific elements. Standard ASTM methods are available for sulfur, ash, gum, and several specific elements.

IV. EXHAUST EMISSIONS

The basic calculations used in the certification of light-duty vehicles are applicable for use with alternate fuels. A number of the parameters utilized, however, must be changed to account for the potentially large differences in fuel and exhaust organic matter composition. In addition, no generally acceptable method is currently available for determination of a meaningful composite value for the organic matter when using alternate fuels. This section describes the determination of the required parameters and the calculational methods to enable incorporation of these parameters.

A. Exhaust Organic Matter Composition

The composition of the exhaust hydrocarbon emissions affects the response factor of the FID instrument and the density of the HC emissions. The current light-duty certification procedure assumes that the overall FID response factor for the hydrocarbons in the exhaust is essentially equal to the FID response factor for propane. For alternate fuels with significantly different composition such response factor assumption is inappropriate.

The only method determined to be generally acceptable for alternate fuels is to measure the individual components of the organic matter in the exhaust, and then to calculate and report all of the major constituents. Unfortunately, this method is impractical in many situations. Therefore, it is recommended that measurement, calculation, and reporting of the individual components of the exhaust organic matter be done to the extent practical. Generally, the summation of the individual components is inappropriate, and should not be done, when alternate fuels are used.

In cases where an exact determination is not essential, a simpler determination can be made by assuming the composition of the organic matter in the exhaust is equal to that in the fuel. Determine the FID response factor for the fuel using the method given in Appendix A-3, and apply this factor to the FID measured value of the exhaust. The method used for determination of organic matter should always be identified for any such data reported.

B. Calculation of Gaseous Exhaust Emissions

The following calculations for exhaust emissions are recommended for use when non-standard carbon bearing fuels are utilized. These calculations are based on the calculations given in 40 CFR Part 86, Subpart B, §86.144-78, with latest revisions incorporated. The basic format used in §86.144 has been retained. Changes or additions to the calculational methods in §86.144-78 are identified by asterisk(*) and are derived in Appendix B-2.

*Analyses, in addition to those required in the certification procedure are as follows:

Composition of the Fuel

FFC = Fuel fraction carbon by weight
FFH = Fuel fraction hydrogen by weight
FFO = Fuel fraction oxygen by weight
FFX = Fuel fraction of other elements by weight

The following is based on §86.144-78 Calculations; exhaust emissions.

The final reported test results shall be computed by use of the following formula:

(a) For light-duty vehicles and light-duty trucks:

$$Y_{wm} = 0.43((Y_{ct} + Y_s)/(D_{ct} + D_s)) + 0.57((Y_{ht} + Y_s)/(D_h + D_s))$$

Where:

Y_{wm} = Weighted mass emissions of each pollutant, i.e.; HC, CO, NO₂, CO₂ in grams per vehicle mile.
 Y_{ct} = Mass emissions as calculated from the "transient" phase of the cold start test, in grams per test phase.
 Y_{ht} = Mass emissions as calculated from the "transient" phase of the hot start test, in grams per test phase.
 Y_s = Mass emissions as calculated from the "stabilized" phase of the cold start test, in grams per test phase.
 D_{ct} = The measured driving distance from the "transient" phase of the cold start test, in miles.
 D_{ht} = The measured distance from the "transient" phase of the hot start test, in miles.
 D_s = The measured driving distance from the "stabilized" phase of the cold start test, in miles.

(b) The mass of each pollutant for each phase of both the cold start test and the hot start test is determined from the following:

- (1) Organic matter mass (to be calculated individually for each major component of organic matter in the exhaust).

$$OM_{mass} = V_{mix} \times \text{Density}_{OM} \times (OM_{conc}/1,000,000)$$

- (2) Oxides of nitrogen mass:

$$NO_{xmass} = V_{mix} \times \text{Density}_{NO_x} \times K_H \times (NO_{xconc}/1,000,000)$$

- (3) Carbon monoxide mass:

$$CO_{mass} = V_{mix} \times \text{Density}_{CO} \times (CO_{conc}/1,000,000)$$

- (4) Carbon dioxide mass:

$$CO_{mass} = V_{mix} \times \text{Density}_{CO_2} \times (CO_{2conc}/100)$$

(c) Meaning of symbols:

- (1) OM_{mass} = Organic matter emissions, in grams per test phase.

$$*Density_{OM} = 14.135/EFC \text{ g/ft}^3 (0.4493/EFC \text{ kg/m}^3) *$$

Based on: Density of hydrocarbons is 16.33 g/ft³ (0.5768 kg/m³), assuming an average carbon-to-hydrogen ratio of 1:1.85, at 68°F (20°C) and 760 mm Hg (101.3 kPa) pressure.

Where:

*EFC = Fraction carbon in exhaust organic matter.

* OM_{conc} = Organic matter concentration of the dilute exhaust sample corrected for background in ppm carbon equivalent.

Where:

$$*OM_{conc} = OM_c - OM_d (1 - 1/DF) *$$

Where:

OM_c = Organic matter concentration of the dilute exhaust sample or, for Diesel, average organic matter concentration of dilute exhaust sample as calculated from the integrated OM traces, in ppm carbon equivalent.

OM_d = Organic matter concentration of the dilution air as measured, in ppm carbon equivalent.*

- (2) NO_{xmass} = Oxides of nitrogen emissions, in grams per test phase.

$Density_{NO_2}$ = Density of oxides of nitrogen is 54.16 g/ft³ (1.913 kg/m³) assuming they are in the form of nitrogen dioxide, at 68°F (20°C) and 760 mm Hg (101.3 kPa) pressure.

NO_{xconc} = Oxides of nitrogen concentration of the dilute exhaust sample corrected for background, in ppm.

$$NO_{xconc} = NO_{xc} - NO_{xd}(1 - 1/DF)$$

Where:

NO_x = Oxides of nitrogen concentration of the dilute exhaust sample as measured, in ppm.

NO_{xd} = Oxides of nitrogen concentration of the dilution air as measured, in ppm.

- (3) CO_{mass} = Carbon monoxide emissions, in grams per test phase.

$Density_{CO}$ = Density of carbon monoxide is 32.97 g/ft³ (1.164 kg/m³), at 68°F (20°C) and 760 mm Hg (101.3 kPa) pressure.

CO_{conc} = Carbon monoxide concentration of the dilute exhaust sample corrected for background, water vapor, and CO₂ extraction, in ppm.

$$CO_{conc} = CO_c - CO_d(1 - 1/DF)$$

Where:

* CO_c = Carbon monoxide concentration of the dilute exhaust sample volume corrected for water vapor and carbon dioxide extraction, in ppm.*

$$*CO_c = (1 - (0.01 + 0.005 \times HCR) CO_{2c} - 0.000323R) CO_{cm} *$$

Where:

CO_{cm} = Carbon monoxide concentration of the dilute exhaust sample as measured, in ppm.

CO_{2c} = Carbon dioxide concentration of the dilute exhaust sample, in percent.

R = Relative humidity of the dilution air, in percent (see §86.142(n)).

*HCR = Hydrogen-to-carbon ratio of the fuel
= (FFH/1.008)/(FFC/12.011)*

CO_d = Carbon monoxide concentration of the dilution air corrected for water vapor extraction, in ppm.

$$CO_d = (1 - 0.000323R) CO_{dm}$$

Where:

CO_{dm} = Carbon monoxide concentration of the dilution air sample as measured, in ppm.

Note: If a CO instrument which meets the criteria specified in §86.111 is used and the conditioning column has been deleted, CO_{cm} can be substituted directly for CO_c and CO_{dm} can be substituted directly for CO_d .

(4) CO_{2mass} = Carbon dioxide emissions, in grams per test phase.

$Density_{CO_2}$ = Density of carbon dioxide is 51.81 g/ft³ (1.830 kg/m³), at 68°F(20°C) and 760 mm Hg(101.3 kPa)

CO_{2conc} = Carbon dioxide concentration of the dilute exhaust sample corrected for background, in percent.

$$CO_{2conc} = CO_{2c} - CO_{2d} (1 - 1/DF)$$

Where:

CO_{2d} = Carbon dioxide concentration of the dilution air as measured, in percent

$$(5) *DF = SPCO_2 / [CO_{2c} + (\Sigma OM_c + CO_c) 10^{-4}] *$$

Where:

* $SPCO_2$ = Stoichiometric percent of CO_2 in undiluted exhaust
 $= (FFC / (FFC + 5.958 FFH + 12.011 \times FFX / MWX + 0.3278 \times SAFR)) \times 100 *$

* ΣOM_c = Summation of the individual components of the organic matter in ppm carbon equivalent. (A value for organic matter as measured by FID can generally be used for ΣOM_c when OM_c by FID divided by $(CO_{2c} + CO_c)$ is less than 0.02).

* FFX and MWX are the fuel fraction and the molecular weight, respectively, of any unknown compound in the fuel.
 If FFX is less than 0.03 and MWX can not readily be determined, assume $MWX = 28.162.$ *

Where:

* $SAFR$ = Stoichiometric air to fuel ratio
 $= 11.514 \times FFC + 34.298 \times FFH - 4.322 \times FFO *$

K_H = Humidity correction factor.

$$K_H = 1 / [1 - 0.0047(H - 75)]$$

for SI units = $1 / [1 - 0.0329(H - 10.71)]$

Where:

H = Absolute humidity in grains (grams) of water per pound (kilogram) of dry air.

$$H = [(43.478) R_c \times R_d] / [P_B - (P_d \times P_c / 100)]$$

for SI units, $H = [(6.211) R_c \times R_d] / [P_B - (P_d \times R_d / 100)]$

R_a = Relative humidity of the ambient air, in percent.

P_d = Saturated vapor pressure, in mm Hg (kPa) at the ambient dry bulb temperature.

P_B = Barometric pressure, in mm Hg (kPa).

V_{mix} = Total dilute exhaust volume in cubic feet per test phase corrected to standard conditions of 528°F (293°K) and 760 mm Hg (101.3 kPa).

For PDP - CVS, V_{mix} is:

$$V_{min} = V_o \times \frac{N(P_B - P_d)}{(760 \text{ mm Hg}) (T_p)} \times \frac{528 \text{ R}}{(T_p)}$$

for SI units,

$$V_{mix} = V_o \times \frac{N(P_B - P_d) (293.15 \text{ K})}{(101.325 \text{ kPa}) (T_p)}$$

Where:

V_o = Volume of gas pumped by the positive displacement pump, in cubic feet (m^3) per revolution. This volume is dependent on the pressure differential across the positive displacement pump.

N = Number of revolutions of the positive displacement pump during the test phase while samples are being collected.

P_B = Barometric pressure, in mm Hg (kPa).

P_d = Pressure depression below atmospheric measured at the inlet to the positive displacement pump, in mm Hg (kPa) (during an Idle mode).

T_p = Average temperature of dilute exhaust entering positive displacement pump during test, R(K).

(d) Example calculation of mass values of exhaust emissions using positive displacement pump and Indolene fuel of $CH_{1.85}$. For Indolene Fuel, the exhaust organic matter are hydrocarbons).

(1) For the "transient" phase of the cold start test assume the following:

$V_o = 0.29344 \text{ ft}^3/\text{revolution}$; $N = 10,485$; $R = 48.0$ percent;

$R_a = 48.2$ percent; $P_B = 762 \text{ mm Hg}$; $P_d = 22.225 \text{ mm Hg}$; $P_4 = 70 \text{ mm Hg}$;

$T_p = 570 \text{ R}$; $HC_c = 105.8 \text{ ppm}$, carbon equivalent; $NO_{xc} = 11.2 \text{ ppm}$;

$CO_{cm} = 306.6 \text{ ppm}$; $CL_c = 1.43$ percent; $HC_d = 12.1 \text{ ppm}$; $NO_{xd} = 0.8 \text{ ppm}$;

$CO_{dm} = 15.3 \text{ ppm}$.

$CO_{2d} = 0.032$ percent; $D_{ct} = 3.598$ miles;

*FFC = 0.8656; FFH = 0.1344;

FFO = 0; FFX = 0; FID Response Factor = 1.0.*

Then:

$$V_{mix} = (0.29344) \times (10.485) \times (762 - 70) \times (528) / (760) (570) = 2595.0 \text{ ft}^3 \text{ per test phase.}$$

$$H = (43.478) (48.2) (22.225) / [782 - (22.225 \times 48.2 / 100)] = 62 \text{ grains of water per pound of dry air.}$$

$$K_H = 1 / [1 - 0.0047(62 - 75)] = 0.9424$$

$$*HCR = (0.1344/1.008)/(0.8656/12.011) = 1.85*$$

$$*CO_c = (1 - (0.01 + 0.005 \times 1.85)1.43 - 0.000323 \times 48)306.6 = 293.4 \text{ ppm}$$

$$CO_d = (1 - 0.000323 \times 48)15.3 = 15.1 \text{ ppm}$$

$$*SAFR = 11.514 \times 0.8656 + 34.298 \times 0.1344 = 14.576*$$

$$*SPCO_2 = (0.8656/(0.8656 + 5.958 \times 0.1344 + 0.3278 \times 14.576)) \times 100 = 13.4*$$

$$*DF = 13.4[1.43 + (105.8/1.0 + 293.4) \times 10^{-4}] = 9.116*$$

$$*HC_{CC} = 105.8 - 12.1(1 - 1/9.116) + 95.03 \text{ ppm.}*$$

$$*HC_{conc} = HC_{CC}/1.0 = 95.03 \text{ ppm}*$$

$$*Density \text{ HC} = 14.135/0.8656 = 16.33 \text{ g/ft}^3*$$

$$HC_{mass} = (2595)(16.33)(95.03/1,000,000) = 4.027 \text{ grams per test phase}$$

$$NO_{xconc} = 11.2 - 0.8(1 - 1/9.116) = 10.49 \text{ ppm}$$

$$NO_{xmass} = (2595)(54.16)(10.49/1,000,000)(0.9424) = 1.389 \text{ gram per test phase.}$$

$$CO_{conc} = 293.4 - 15.1(1 - 1/9.116) = 280.0 \text{ ppm}$$

$$CO_{mass} = (2595)(32.97)(280/1,000,000) = 23.96 \text{ grams per test phase}$$

$$CO_{2conc} = 1.43 - 0.032(1 - 1/9.116) = 1.402\%$$

$$CO_{2mass} = (2595.0)(51.85)(1.402/100) = 1886 \text{ grams per test phase.}$$

(2) For the stabilized portion of the cold start test assume that similar calculations resulted in the following:

$$HC_{mass} = 0.62 \text{ grams per test phase}$$

$$NO_{xmass} = 1.27 \text{ grams per test phase}$$

$$CO_{mass} = 5.98 \text{ grams per test phase}$$

$$CO_{2mass} = 2346 \text{ grams per test phase}$$

$$D_s = 3.902 \text{ miles}$$

(3) For the "transient" portion of the hot start test assume that similar calculations resulted in the following:

$$HC_{\text{mass}} = 0.51 \text{ grams per test phase}$$

$$NO_{x\text{mass}} = 1.38 \text{ grams per test phase}$$

$$CO_{\text{mass}} = 5.01 \text{ grams per test phase}$$

$$CO_{2\text{mass}} = 1758 \text{ grams per test phase}$$

$$D_{\text{ht}} = 3.598 \text{ miles}$$

(4) Weighted mass emission results:

$$HC_{\text{wm}} = 0.43[(4.027 + 0.52)/(3.598 + 3.902)] + 0.57[(0.51 + 0.62)/(3.598 + 3.902)] = 0.352 \text{ grams per vehicle mile.}$$

$$NO_{x\text{wm}} = 0.43[(1.389 + 1.27)/(3.598 + 3.902)] + 0.57[(1.38 + 1.27)/(3.598 + 3.902)] = 0.354 \text{ grams per vehicle mile.}$$

$$CO_{\text{wm}} = 0.43[(23.96 + 5.98)/(3.598 + 3.902)] + 0.57[(5.01 + 5.98)/(3.598 + 3.902)] = 2.55 \text{ grams per vehicle mile.}$$

$$CO_{2\text{wm}} = 0.32[(1886 + 2346)/(3.598 + 3.902)] + 0.57[(1758 + 2346)/(3.598 + 3.902)] = 555 \text{ grams per vehicle mile}$$

*(e) Example calculation of mass values of exhaust emissions when using methanol fuel (CH_4O). All inputs except for fuel composition and exhaust organic matter are assumed to be the same as in previous example (d).

(1) Fuel composition is as follows:

$$FFC = 0.375; FFH = 0.126; FFO = 0.499;$$

Then:

$$V_{\text{mix}} = 2595.0 \text{ ft}^3$$

$$K_H = 0.9424$$

$$HCR = (0.126/1.008)/0.375/12.011 = 4.00$$

$$CO_C = (1 - (0.01 + 0.005 \times 4.00))1.43 - 0.000323 \times 48)306.6 = 288.7 \text{ ppm}$$

$$CO_d = 15.1 \text{ ppm}$$

$$\text{SAFR} = 11.514 \times 0.375 + 34.298 \times 0.126 - 4.322 \times 0.499 = 6.48$$

$$\text{SPCO}_2 = (0.275 / (0.375 + 5.958 \times 0.126 + 0.3278 \times 6.482)) \times 100 = 11.54$$

$$(\text{OM}_C \text{ by FID}) / (\text{CO}_{2C} + \text{CO}_C) = 105.8 / (1.43 \times 10^4 + 288.7) = 0.007 < 0.02.$$

Therefore:

$$\text{DF} = 11.54 / [1.43 + (105.8 + 288.7) \times 10^{-4}] = 7.84$$

$$\text{Density}_{\text{OM}} = 14.135 / 0.375 = 37.69 \text{ g/ft}^3$$

$$\text{OM}_{\text{conc}} = (\text{Data are not available})$$

$$\text{OM}_{\text{mass}} = (\text{can not be determined from available data})$$

$$(\text{OM}_{\text{mass}} \text{ by FID} = (2595) (37.69) (105.8 / 1,000,000) = 10.35 \text{ grams per test phase}).$$

$$\text{NO}_{x\text{conc}} = 11.2 - 0.8(1 - 1/7.84) = 10.50 \text{ ppm}$$

$$\text{NO}_{x\text{mass}} = (2595) (54.16) (10.51 / 1,000,000) (0.9424) = 1.391 \text{ grams per test phase.}$$

$$\text{CO}_{\text{conc}} = 288.7 - 15.1(1 - 1/7.84) = 275.5 \text{ ppm}$$

$$\text{CO}_{\text{mass}} = (2595) (32.97) (275.5 / 1,000,000) = 23.57 \text{ grams per test phase.}$$

$$\text{CO}_{2\text{conc}} = 1.43 - 0.032(1 - 1/7.84) = 1.402\%$$

$$\text{CO}_{2\text{mass}} = (2595.0) (51.85) (1.402 / 100) = 1886 \text{ grams per test phase.}^*$$

Assume 100.0 ppm of methanol was measured in the exhaust using the method given in Appendix A-5:

$$\text{Methanol}_{\text{mass}} = (2595) (37.69) (100.0 / 1,000,000) = 9.781 \text{ gram per test phase}$$

Note: Other components of organic matter may also be present in the exhaust; these should be determined and reported to the extent practical.

C. Calculation of Particulate Exhaust Emissions

Use of an alternate fuel does not affect the procedure or the calculations for diesel particulate emissions. Therefore, the procedure and the calculations for diesel particulate given in 40 CFR, Part 86, Subpart B can be used, without modification.

D. Non-Methane Hydrocarbons

Consideration has been given by the EPA to basing the HC emissions standard on non-methane hydrocarbons. This, however, has not been done, but remains under consideration. Therefore, for the present, it is recommended that the composition of the exhaust HC be defined to the extent practical (e.g., with methanol fuel, measure exhaust for methanol, aldehydes, and methane). Such data could provide input for future decisions as more becomes known about individual hydrocarbon photochemical reactivity.

E. Unregulated Exhaust Emissions

It has been determined that some alternate fuels may be prone to produce higher levels of some currently unregulated emissions (e.g., alcohol fuels tend to produce higher levels of aldehydes). At the present time, the only specific recommendation is to include the analyses of aldehydes and the specific alcohol, when an alcohol fuel is used.

Other emissions to measure for, if any, should be based on the composition of the fuel and on whichever emission(s) is (are) considered to be of sufficient importance at the time. No specific recommendations can be provided, but some general guidance can be given. Under certain operating conditions the following relationships might occur:

| <u>Fuel Description</u> | <u>Other Exhaust Emissions to Consider</u> | <u>Source for Analytical Method</u> |
|---------------------------|---|---|
| Methanol | aldehydes (primarily formaldehyde) unburned methanol | Appendix A-4 Appendix A-5 |
| Ethanol | aldehydes (primarily acetaldehyde) unburned ethanol | Appendix A-4 Appendix A-6 |
| High sulfur | sulfate sulfur dioxide | Reference 3 Reference 3 |
| High aromatic (diesel) | phenols nitropyrenes Ames response | Reference 7 Reference 8 Reference 9 |
| High Nitrogen (diesel) | nitropyrenes Ames Response | Reference 8 Reference 9 |

V. FUEL ECONOMY

The procedure for determination of fuel economy, given in the fuel Economy Regulations,⁽⁹⁾ can be made applicable for use with alternate fuels by incorporating the actual carbon fraction and density of the fuel. For alternate fuels, however, the resulting expression of fuel economy, in terms of distance traveled per unit volume of fuel used, is not sufficient to enable making meaningful comparisons. Therefore, an additional method for expression of fuel economy (consumption) has been selected and its use is recommended.

A. Calculation of Fuel Economy

The following calculations for fuel economy are recommended for use when non-standard carbon containing fuels are utilized. These calculations are based on the method given in 40 CFR Part 600, Subpart B, §600.113-78 and Appendix II, with latest revisions incorporated. The basic format used in §600.113 has been retained. Changes or additions to the calculation method in §600-113-78 are identified by asterisk(*). A required analysis, in addition to those required in §600.113 or in the previous calculation procedure for exhaust emissions given in Section IV of this report, is the density of the fuel.

The following is based on §600-113-78 Fuel economy calculations.

The calculations of vehicle fuel economy values require the weighted grams/mile values for OM, CO, and CO₂. (OM = Organic matter).

(a) Calculate the weighted grams/mile values for the city fuel economy test for OM, CO, and CO₂ as specified in the modifications to §86.144 as given in Section IV.B. of this report.

(b)(1) Calculate the mass values for the highway fuel economy test for OM, CO, and CO₂ as specified in modifications to paragraph (b) of §86.144 as given in Section IV.B. of this report.

(2) Calculate the grams/mile values for the highway test for OM, CO, and CO₂ by dividing the mass values obtained in (b)(1) by the actual distance traveled, measured in miles, as specified in paragraph (h) of §86.135.

(c) Calculate the city fuel economy and highway fuel economy from grams/mile values for OM, CO, and CO₂. The OM values (obtained per paragraph (a) or (b) as applicable) used in each calculation in this section are rounded to the nearest 0.01 grams/mile. The CO values (obtained per paragraph (a) or (b) as applicable) used in each calculation in this section are rounded to the nearest 0.1 grams/mile. The CO₂ values (obtained per paragraph (a) or (b) of this section as applicable) used in each calculation in this section are rounded to the nearest gram/mile.

(d) and (e) Calculate the fuel economy in miles per gallon of gasoline by dividing the grams of carbon per gallon of fuel (GCPG) by the sum of three terms:

*Where: $GCPG = 3785.4 \times FFC \times FDEN$

*Where: FFC = Fuel Fraction Carbon
FDEN = Fuel Density, g/ml

Note: GCPG is equal to 2421 for certification gasoline and 2778 for certification diesel fuel.

(1) OMFC multiplied by OM (in grams/mile as obtained in paragraph (c)).

(2) 0.429 multiplied by CO (in grams/mile as obtained in paragraph (c)), and

(3) 0.273 multiplied by CO₂ (in grams/mile as obtained in paragraph (c)).

*Where: OMFC is the fraction carbon in the exhaust organic matter.
[If OM_C by FID divided by (CO_{2C} + CO_C) is less than 0.02, the FID measured value can generally be used for OM_C. Also, it can be assumed that OMFC = FFC].

Round the quotient to the nearest 0.1 mile per gallon.

The following is based on Appendix II - Sample Test Value Calculation

(a) Assume that a gasoline-fueled vehicle was tested by the Federal Emission Test Procedure and the following results were calculated:

HC = 1.03 grams/mile
CO = 6.74 grams/mile
CO₂ = 785 grams/mile

According to the preceding procedure, the fuel economy or MPG_C for the vehicle may be calculated by substituting the HC, CO and CO₂ grams/mile values into the following equation.

* $MPG_C = GCPG / (HCFC \times HC + (0.429 \times CO) + (0.273 \times CO_2))$ *

*For Certification Gasoline: FFC = 0.8656; FDEN = 0.739 g/ml

$GCPG = 3785.4 \times 0.8656 \times 0.739 = 2421$

$HCFC = FFC = 0.8656$ (or 0.866) *

$MPG_C = 2421 / ((0.866 \times 1.03) + (0.429 \times 6.74) + (0.273 \times 785))$
 $= 2421 / 218.1 = 11.1$ MPG

(b) Assume that the same vehicle was tested by the Federal Highway Fuel Economy Test Procedure and a calculation similar to that shown in (a) resulted in a highway fuel economy of MPG_h of 18.6. According to the procedure in §600.113, the combined fuel economy (called $MPG_{c/h}$) for the vehicle may be calculated by substituting the city and highway fuel economy values into the following equation:

$$\begin{aligned} MPG_{c/h} &= 1/(0.55/MPG_c + 0.45/MPG_h) \\ &= 1/(0.55/11.1 + 0.45/18.6) = \\ MPG_{c/h} &= 13.6 \text{ MPG} \end{aligned}$$

*(c) Assume that a methanol-fueled vehicle was tested and the same emission results were obtained (OM = 1.03 grams/mile).

Then: $FFC = 0.375$

$FDEN = 0.7914 \text{ g/ml}$

$GCPG = 3785.4 \times 0.375 \times 0.7914 = 1123$

$OMFC = FFC = 0.375$

$MPG = 1123 / ((0.375 \times 1.03) + (0.429 \times 6.74) + (0.273 \times 785))$
 $= 1123 / 217.6 = 5.2 \text{ MPG}^*$

B. Fuel Economy Related Considerations

There are several potentially important additional fuel economy related considerations.

Energy Based Fuel Consumption - Alternate fuels can have a greatly different energy content per unit of volume and weight than is contained in current petroleum fuels. Therefore, fuel economy (or consumption) expressed only in miles per gallon (l/100 km) can present a distorted picture of the actual fuel energy consumed. Fuel consumption expressed in terms of Btu per mile (J/km) can provide potentially useful information regarding the actual fuel energy consumed. Therefore, it is recommended that fuel consumption be expressed in Btu per mile, in addition to expressing fuel economy in miles per gallon. The calculation for energy based fuel consumption, derived in Appendix C-1, is as follows:

$$EBFC = 8.34 \times LHV \times FDEN / FE$$

Where:

$EBFC$ = Energy Based Fuel Consumption, Btu/mi

LHV = Fuel Lower Heating Value, Btu/lb

$FEDN$ = Fuel Density, g/ml

FE = Fuel Economy, mpg

Effect of Particulate Emissions - The fuel economy calculation in Subpart B does not account for the carbon in the particulate emissions. With a standard hydrocarbon diesel fuel and particulate emissions below 0.6 g/mi, the effect of the particulate emissions on the calculated fuel economy is considered to be negligible.

The effect of the particulate emissions on the calculated fuel economy can be determined as follows:

$$\text{MPG}_w = \text{MPG} / (1 + 0.00036 \times \text{Part.} \times \text{PFC} \times \text{MPG})$$

Where:

- MPG_w = Calculated miles per gallon with particulate emissions included
- MPG = Calculated miles per gallon using the method in §600.113-78
- Part. = Particulate emissions rate in g/km
- PFC = Mass fraction of carbon in the particulate

Example values are given as follows:

| <u>Particulate Emissions, g/mi</u> | <u>Calculated Fuel Economy, mpg</u> | |
|--|-------------------------------------|------------------------|
| | <u>MPG</u> | <u>MPG_w</u> |
| 0.2 | 25 | 24.9 |
| 0.2 | 50 | 49.8 |
| 0.6 | 25 | 24.8 |
| 0.6 | 50 | 49.5 |

For alternate fuels, the expression of fuel economy in terms of distance traveled per unit volume of fuel used may not always be sufficient to enable making meaningful comparisons.

Other Considerations - For additional discussion of methods for expressing fuel economy, the reader is referred to Reference 2.

VI. EVAPORATIVE EMISSIONS

Calculation of a mass value for evaporative emissions requires knowing the molecular weight per carbon atom of the emissions. The calculation method given in the light-duty certification procedure uses molecular weights derived from empirical data.⁽¹⁾ Those molecular weights, however, are only applicable to gasolines meeting the prescribed fuel specifications. Therefore, for alternate fuels, molecular weights have to be determined or satisfactorily estimated. Derivations of equations are given in Appendix C-2.

A. Evaporative HC Composition for Hydrocarbon Fuels

Detailed analyses for the composition of the evaporative hydrocarbon emissions is extremely difficult, and is considered to be an impractical requirement in the routine determination of evaporative emissions for experimental blends of hydrocarbon fuels. Therefore, assumption and methods for estimation of the composition of the evaporative HC emissions are given for various categories of such fuels as follows:

Standard Cuts of Hydrocarbon Fuels - Standard fuel cuts, containing hydrocarbons composed of carbon and hydrogen (at least ninety-nine percent) and providing for acceptable vehicle performance, would be expected to have reasonably similar distribution of hydrocarbon types regardless of the basic source of the fuel. For such fuels, the value of the molecular weight can be assumed to be the same value as used for certification gasoline.

Nonstandard Cuts or Blends of Hydrocarbon Fuels - For blends of individual hydrocarbons, these components could be analyzed for in the evaporative emissions using GC. For nonstandard cuts and for unknown blends, however, it will generally be necessary to estimate the composition of the evaporative emissions.

The extremes of the hydrogen-to-carbon ratios for hydrocarbons, that could be in gasoline or otherwise used as fuel, range from 1:1 (benzene) to 4:1 (methane). This results in extremes in the K value of 2.7 and 3.3. (The K value is equal to 0.208 times the molecular weight per carbon atom of the fuel and is used in the calculation of evaporative emissions. With the K value approaching 3.3, it is likely that the constituents would be known or could be readily determined. In all likelihood, the K value for a liquid hydrocarbon fuel consisting of essentially all carbon and hydrogen would be equal to or only slightly lower than 3.0. Therefore, for all liquid hydrocarbon fuels for use in gasoline engines, it is recommended that the K values for certification gasoline be used. For gaseous hydrocarbon fuels, it is recommended that the K value be determined using GC analyses of the evaporative hydrocarbon emissions (as an alternative a K value derived from the fuel constituent having the highest H/C ratio could be used).

B. Evaporative Organic Matter Composition for Other Fuels

The only method found to be generally acceptable for determination of evaporative organic matter composition with alternate fuels is to measure the individual evaporative components. The evaporative emissions should then be computed and reported separately for each of the major components.

C. Calculation of Evaporative Emissions

The following calculations for evaporative emissions are recommended for use when nonstandard carbon containing fuels are utilized. These calculations are based on the method given in 40 CFR Part 86, Subpart B, §86.143-78, with latest revisions incorporated. The basic format used in §86.143 has been retained. Changes or additions to the calculation method in §86.143-78 are identified by asterisk(8).

The following is based on §86.143-78 Calculations; evaporative emissions.

The calculation of the net hydrocarbon mass change in the enclosure is used to determine the diurnal and hot soak mass emissions. The mass is calculated using initial and final hydrocarbon concentrations in ppm carbon, initial and final enclosure ambient temperatures, initial and final barometric pressures, and net enclosure volume using the following equation:

$$C_{OM} = kV_n \times 10^{-4} \left[\frac{C_{OMf} P_{Bf}}{T_f} - \frac{C_{OMi} P_{Bi}}{T_i} \right]$$

Where:

M_{OM} = organic matter mass, g

C_{OM} = organic matter concentration as ppm carbon

* V_n = net enclosure volume, ft³(m³) as determined by subtracting 50 ft³ (1.42 m³) or an approved measured volume, (volume of vehicle with trunk and windows open) from the enclosure volume.*

P_B = barometric pressure, in Hg (kPa)

T = enclosure ambient temperature, R(K)

i = indicates initial reading

f = indicates final reading.

K = 0.208×(Mol. Wt. per carbon atom)

for SI units, $K = 1.2 \times (\text{Mol. Wt. per C})$

For Certification Gasoline:

$K = 2.98(17.2)$ for diurnal emissions, (MW/C = 14.33)

$K = 2.95(17.0)$ for hot soak emissions, (MW/C = 14.18)

*For Other Fuels:

Molecular Weight per carbon atom is to be determined.*

The reported results shall be computed by summing the individual evaporative emission results determined in the diurnal breathing-loss test, the running-loss test, and the hot-soak test.

REFERENCES

1. Code of Federal Regulations, Title 40, Chapter 1, Part 86, Subpart B.
2. Harvey, Craig A., "Gasoline-Equivalent Fuel Economy Determined for Alternate Automotive Fuels," SAE Paper 820794, 1982.
3. Dietzmann, Harry E., et al, "Analytical Procedures for Characterizing Unregulated Pollutant Emissions from Motor Vehicles," Final Report EPA 600/2-79-017, February 1979.
4. Methane Measurement Using Gas Chromatography, SAE J1151a, Published in the 1980 SAE Handbook.
5. Smith, Lawrence R. and Urban, Charles, "Characterization of Exhaust Emissions from Methanol and Gasoline Fueled Automobiles," Final Report EPA 460/3-82-004, March 1982.
6. Bykowski, Bruce B., "Gasohol, TBA, MTBE Effects on Light-Duty Emissions," Final Report of Task No. 6 of EPA Contract 68-03-2377, October 1979.
7. Dietzmann, Harry E., et al, "Analytical Procedures for Characterizing Unregulated Pollutant Emissions from Vehicles Using Middle-Distillate Fuels," Interim Report EPA-600/2-80-068, April 1980.
8. Tejada, Silvestre B., et al, "Analysis of Nitroaromatics in Diesel and Gasoline Emissions," SAE Paper 820775.
9. Ames, B., et al, "Methods for Detecting Carcinogens and Mutagens with the Salmonella/Mammalian-Microsome Mutagenicity Test." Mutation Research, 31, pp, 347-364, 1975.
10. Code of Federal Regulations, Title 40, Chapter 1, Part 600, Subpart B.
11. Weast, Robert C., Ph.D., "Handbook of Chemistry and Physics," 54th Edition, CRC Press, 1974.

SELECTED BIBLIOGRAPHY

A number of other sources of information, although not directly utilized in this report, had an influence on the selection of the specific methods recommended and on the comments provided in this report.

Furey, Robert L., and King, Jack B., "Evaporative and Exhaust Emissions from Cars Fueled with Gasoline Containing Ethanol or Methyl tert-Butyl Ether," SAE Paper 800261.

Espinola, Stephen A., and Pefley, Richard K., "Alternate Fuel Influences on Emissions Test Procedures," Paper to be presented at the American Chemical Society Symposium on "Chemistry of Oxygenates in Fuel," September 12-17, 1982.

Wagner, T. O., et al. "Practicality of Alcohols as Motor Fuels," SAE Paper 790429.

Lawrence, D.C., and Niemczak, D.J., "Evaporative and Exhaust Emissions of Two Automobiles Fueled with Volatility Adjusted Gasohol," Paper EPA-AA-TED-81-12, December 1980.

Lang, J. M., and Black, F. M., "Impact of Gasohol on Automobile Evaporative and Tailpipe Emissions," SAE Paper 810438.

Dimitriades, B., and Joshi, S. B., "Application of Reactivity Criteria in Oxidant-Related Emission Control in the USA," International Conference Report EPA 600/3-77-001b, January 1977.

Memorandum from Frank Black, Chief, ETCS to Karl Hellman, Chief, CTAB, "Emissions and Fuel Consumption Measurements with Alternate Fuels," September 24, 1981.

Lawrence, Richard, "Fuel Economy Measurement Carbon Balance Method," Draft Report, October 1981.

California Air Resources Board, "Alcohol Fueled Fleet Test Program," Project 3T8001, Second Interim Report, July 1981.

Alson, Jeff, "A Brief Summary of the Technical Feasibility, Emissions and Fuel Economy of Pure Methanol Engines," Technical Report EPA-AA-SDSB-82-1, December 1981.

California Air Resources Board, "Fuel Economy Calculations for LPG Based on the Cold Start CVS-1975 Federal Test Procedure," Proposed amendments transmitted by letter dated August 3, 1982.

California Air Resources Board, "California Exhaust Standards and Test Procedures for Systems Designed to Convert Motor Vehicles to Use Alcohol or Alcohol/Gasoline Fuels," Proposed procedure received in September 1982.

APPENDICES

- A. APPENDICES FOR SECTIONS I THROUGH III
- B. APPENDICES FOR SECTION IV
- C. APPENDICES FOR SECTIONS V AND VI

NOTE: The format used for mathematical equations is in accordance with that used in the Code of Federal Regulations.⁽¹⁾ Computations are to be performed in accordance with the IBM hierarchy of operations.

APPENDIX A

- A-1. CVS FLOWRATE DETERMINATION
- A-2. DERIVATION OF CO₂ CONCENTRATION ESTIMATION
- A-3. DETERMINATION OF FID RESPONSE FACTOR
- A-4. MEASUREMENT OF ALDEHYDES AND KETONES
- A-5. MEASUREMENT OF METHANOL
- A-6. MEASUREMENT OF ETHANOL

APPENDIX A-1

CVS FLOWRATE DETERMINATION

$$\text{System H}_2\text{O Capacity} = \text{CVSFR} \times \text{DEN}_{\text{AIR}} \times (\text{SatSH} - \text{TestSH}), \text{ wt/time}$$

Where:

CVSFR = CVS flowrate, scfm

DEN_{AIR} = Density of air = 0.075 lb/scf

SatSH = Saturation specific humidity, lb/lb

TestSH = Specific humidity at test conditions, lb/lb

$$\text{Fuel H}_2\text{O Addition} = \text{FC} \times \text{FFH} \times 18.016 / 2.016, \text{ wt/time}$$

Where:

FC = Fuel Consumption in lb/min

= $1 / (\text{MPG} \times \text{Avg.MPH} / 60 \times 8.34 \times \text{FDEN})$

FFH = Fuel fraction hydrogen

MPG&MPH = Mile per gallon and per hour

FDEN = Density of the fuel, g/ml

Then:

$$\text{CVSFR} \times 0.075 (\text{SSH} - \text{TSH}) = \text{AMPH} / \text{MPG} \times \text{FDEN} \times \text{FFH} \times 1.24$$

$$\text{CVSFR} = 16.56 \times \text{AMPH} \times \text{FDEN} \times \text{FFH} / (\text{MPG} (\text{SSH} - \text{TSH}))$$

APPENDIX A-2

DERIVATION OF CO₂ CONCENTRATION ESTIMATION

$$\text{CO}_2\% = (\text{MRCO}_2/\text{MRAIR}) \times (\text{MWAIR}/\text{MWCO}_2) \times 100$$

Where:

$$\begin{aligned}\text{MRCO}_2 &= \text{Mass flowrate of CO}_2 \\ &= 1/\text{FE} \times \text{AMPH} \times 8.34 \times \text{FDEN} \times (44.011/12.011)/60 \\ &= 0.14 \times \text{AMPH} \times \text{FDEN}/\text{FE}\end{aligned}$$

Where: FE = Fuel economy, mpg
AMPH = Average speed for the test cycle, mph
FDEN = fuel density, g/ml

$$\text{MRAIR} = 0.075 \times \text{CVSFR}$$

Where: CVSFR = flowrate of the CVS, scfm

$$\text{MWAIR} = 28.966$$

$$\text{MWCO}_2 = 44.011$$

$$\begin{aligned}\text{CO}_2\% &= (0.514 \times \text{AMPH} \times \text{FDEN}/\text{FE}) / (0.075 \times \text{CVSFR}) \times (28.966/44.011) \times 100 \\ &= (451 \times \text{AMPH} \times \text{FDEN}) / (\text{FE} \times \text{CVSFR})\end{aligned}$$

APPENDIX A-3

DETERMINATION OF FID RESPONSE FACTOR

For determination of the response factor of a specific FID hydrocarbon analyzer, with a liquid fuel, a standard of known concentration is prepared using the apparatus shown in Figure A-1. (Figure A-1 was adapted from SAE Paper 810438).

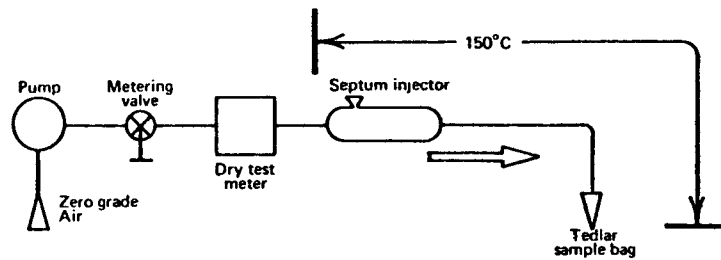


Figure 1. Apparatus for preparation of calibration mix

In preparing this standard, a known volume of the fuel of interest is injected, using a microliter syringe, into the heated mixing zone (150°C) of the apparatus. The liquids are vaporized and swept into a Tedlar bag with a known volume of air measured by a dry test meter. The concentration of the fuel (in ppm C) in the bag is determined as follows:

$$FCONC = FINJ \times FDEN / AIRV \times 1000$$

Where:

FCONC = Concentration of the fuel, $\mu\text{g}/\text{m}^3$
 FINJ = Volume of fuel injected, μl
 FDEN = Density of the fuel, g/ml
 AIRV = Volume of air, m^3 (scfm $\times 0.02832$)

$$FPPMC = FCONC \times 0.02406 / FMWC$$

Where:

FPPMC = Fuel concentration as ppmC
 0.02406 = Volume in m^3 of one mole at 29.92 in. Hg and 68°F
 FMWC = Fuel molecular weight per carbon atom

Note: It is recommended that the response factor of the FID analyzer be determined at several concentrations in the range of the concentrations in the exhaust samples.

For gaseous fuels, the standards can be prepared using a precision proportioning system or any other available method that provides accurate volume measurements of the blending gases. For gaseous fuels:

$$GFPPMC = GFVOL \times CPM / (AIRV + GFVOL) \times 10^6$$

Where:

GFPPMC = ppmC of the gaseous fuel
GFVOL = Volume of the gaseous fuel
AIRV = Volume of air
CPM = Carbon atoms per molecule

The calibration sample is evaluated using the FID analyzer and the FID response factor is determined as follows:

$$FIDRF = FIDPPM / FPPMC$$

Where:

FIDRF = Response factor for the FID
FIDPPM = ppmC value as read on the FID
FPPMC = Actual fuel concentration as ppmC

Example A. Assume a gasoline with a density of 0.739 g/ml and a fuel fraction carbon of 0.8656.

$$FMWC = 1 / 0.8656 \times 12.011 = 13.86$$

$$\begin{aligned} \text{Let: } FINJ &= 1.00 \\ AIRV &= 1 \text{ scfm} = 0.02832 \text{ m}^3 \end{aligned}$$

$$\begin{aligned} \text{Then: } FCONC &= 1.00 \times 0.739 / 0.02832 \times 1000 = 26,095 \\ FPPMC &= 26,095 \times 0.02406 / 13.86 = 45.4 \end{aligned}$$

$$\text{Let: } FIDPPM = 45.4$$

$$\text{Then: } FIDRF = 45.4 / 45.4 = 1.00$$

Example B. Assume methanol is used as the fuel.

$$\begin{aligned} FMWC &= 32.04 \\ FDEN &= 0.7914 \text{ g/ml} \end{aligned}$$

$$\begin{aligned} \text{Let: } FINJ &= 1.00 \text{ } \mu\text{l} \\ AIRV &= 1 \text{ scfm} = 0.02832 \text{ m}^3 \end{aligned}$$

$$\begin{aligned} \text{Then: } FCONC &= 1.00 \times 0.7914 / 0.02832 \times 1000 = 27,945 \\ FPPMC &= 27,945 \times 0.02406 / 32.04 = 21.0 \end{aligned}$$

$$\text{Let: } FIDPPM = 17.4$$

$$\text{Then: } FIDRF = 17.4 / 21.0 = 0.83$$

APPENDIX A-4

MEASUREMENT OF ALDEHYDES AND KETONES

The measurement of aldehydes (formaldehyde, acetaldehyde, isobutyraldehyde, crotonaldehyde, hexanaldehyde, and benzaldehyde) and ketones (acetone and methylethylketone) in exhaust is accomplished by bubbling the exhaust through glass impingers containing 2,4 dinitrophenylhydrazine (DNPH) in dilute hydrochloric acid. The exhaust sample is collected continuously during the test cycle. The aldehydes and ketones (also known as carbonyl compounds) react with the DNPH to form their respective phenylhydrazone derivatives. These derivatives are insoluble or only slightly soluble in the DNPH/HCl solution and are removed by filtration followed by pentane extractions. The filtered precipitate and the pentane extracts are combined and then the pentane is evaporated in a vacuum oven. The remaining dried extract contains the phenylhydrazone derivatives. The extract is dissolved in a quantitative volume of toluene containing a known amount of anthracene as an internal standard. A portion of this dissolved extract is injected into a gas chromatograph and analyzed using a flame ionization detector. The detection limits for this procedure under normal operating conditions are on the order of 0.005 ppm carbonyl compound in dilute exhaust.

LIST OF EQUIPMENT

The equipment required for the analysis of aldehydes and ketones is divided into three groups: sample acquisition, sample preparation, and sample analysis. Manufacturer, stock number and any pertinent descriptive information are listed.

Sample Acquisition

1. Glass impingers, Ace Glass Products, Catalog #7530-11, plain tapered tip stoppers with 18/7 arm joints and 29/42 bottle joints.
2. Flowmeter, Brooks Instrument Division, Model 1555, tube size R-2-15-C, graduated 0-15, sapphire float, 0-5 l/min range.
3. Sample pump, Thomas Model 106 CA18, capable of free flow capacity of 4 l/min.
4. Dry gas meter, American Singer Corporation, Type AL-120, 60 CFH capacity.
5. Regulating valve, Nupro 4MG, stainless steel.
6. Teflon tubing, United States Plastic Corporation, 1/4" OD \times 1/8" ID and 5/16" OD \times 1/8" ID.

7. Teflon solenoid valve, The Fluorocarbon Company, Model DV2-144NCA1.
8. Drying tube, Analabs, Inc., Catalog #HGC-146, 6" long, 1/4" brass fittings.
9. Miscellaneous Teflon nuts, ferrules, unions, tees, clamps, connectors, etc.
10. Digital readout for dry gas meter.
11. Miscellaneous electrical switches, lights, wirings, etc.
12. Six channel digital thermometer, Analog Devices, Model #2036/J/1.
13. Iron/Constantan type J single thermocouple with 1/4" OD stainless steel metal sheath, Thermo Sensors Corporation.
14. Variable autotransformer, Staco Inc., Type 3PN 1010.
15. Heating sleeve wrapped with insulation and insulation tape.
16. Class A, 20 ml volumetric pipets.
17. Class A, 1000 ml volumetric flask.
18. Teflon coated stirring bar.
19. Hot plate-stirrer, Corning, PC-351.

Sample Analysis

1. Varian 1700 gas chromatograph equipped with dual flame ionization detectors in differential operation, and a linear temperature programmer.
2. Soltec Model B-281 1 mv recorder.
3. Hewlett-Packard Model 3354 gas chromatograph computer system with remote teletype printout.
4. Syringe, 10 ml, Hamilton Company, #701.
5. Dual columns, 24 x 1/8" OD, stainless tubing packed with 6.7 percent Dexsil 300 GC on Chromosorb G 60/80 mesh, DMCS treated and acid washed.

Sample Preparation

1. Fritted glass filters, Ace Glass Company, porosity D, ASTM 10 - 20 microns pore size, 24/40 ground glass joint, vacuum takeoff.

2. Constant temperature vacuum oven, National Appliance Company.
3. Pump for oven, Thomas Industries, Model 907CA18 2.
4. Flasks, 125 ml capacity, 24/40 ground glass joints.
5. Separatory funnels, 125 ml.
6. Separatory funnels, 250 ml.
7. Separatory funnel shaker, Burrell Corporation, Wrist-Action (R) type with appropriate funnel holders, Model 75.
8. Ring stands, labels, holders, tubing, vacuum tubing, fittings and clamps needed for equipment manipulation.
9. Wash bottles, 500 ml.
10. Graduated cylinders, 50 ml.
11. Vials, Kimble, 1/2 dram.
12. Vacuum pump, Sargent-Welch.

LIST OF REAGENTS

A list of the reagents used in the determination of the aldehydes and ketones in exhaust is provided along with chemical formula, molecular weight, purity, manufacturer, and catalog number.

1. Hydrochloric acid, HCl, 36.46 g/mole, concentrated (37%), analytical reagent, Mallinckrodt, Cat. #2612.
2. Pentane, C_5H_{12} , 72.15 g/mole, Distilled in glass (bp 35-37°C), Burdick and Jackson Laboratories, Inc.
3. 2,4 Dinithrophenylhydrazine (2,4-DNPH), $(NO_2)_2C_6H_3CH=N-NH_2$, 210.149 g/mole, Aldrich analyzed, Aldrich, Cat. #D19,930-3.
4. Sodium Bicarbonate, $NaHCO_3$, 84.00 g/mole, Mallinckrodt, Cat. #7412.
5. Anthracene, $C_{14}H_{10}$, 178.24 g/mole, K and K Laboratories, Cat. #10714.
6. Toluene, $C_6H_5CH_3$, 92.14 g/mole Baker Analyzed Reagent, Baker Cat. #3-9460.
7. Methylene Chloride, CH_2Cl_2 , 84.93 g/mole, Reagent ACS, Eastman, Cat. #13022.

PREPARATION OF ABSORBING SOLUTION

To prepare the absorbing solution, 163 ml of concentrated HCl and 2.5 g of 2,4-DNPH crystals are added to a one liter volumetric flask containing about 500 ml of deionized water. The flask is diluted to mark and stirred for several hours at room temperature with an automatic stirrer/teflon coated stirring bar to dissolve the DNPH. Fresh absorbing solution is prepared daily as needed.

PREPARATION OF TOLUENE/ANTHRACENE SOLUTION

Toluene containing approximately 0.05mg anthracene per ml of toluene is used to dissolve the dried phenylhydrazone extracts. This solution is made by adding 100 mg of anthracene to a two liter volumetric flask and diluting to mark with toluene.

PREPARATION OF PHENYLHYDRAZONE DERIVATIVES

In order to obtain response factors for each of the phenylhydrazone derivatives to anthracene, pure derivatives were prepared from their respective aldehydes and ketones. These derivatives were made by adding each of the carbonyl compounds separately to a 2N HCl-DNPH solution. The resulting orange to red precipitates were filtered and dried. The derivatives were then recrystallized from hot absolute ethanol. The melting points for each of the derivatives were compared to literature values before use. A GC trace was also made on each of the derivatives to further check the purity.

PREPARATION OF STANDARD SOLUTION OF PHENYLHYDRAZONE DERIVATIVES AND ANTHRACENE

A standard containing the phenylhydrazone derivatives and anthracene in toluene is prepared to obtain a response factor of each of the derivatives to anthracene. The solution is made by dissolving weighed amounts of anthracene and each of the derivatives in a quantitative volume of toluene. These solutions contain approximately 0.05mg anthracene per ml of toluene and approximately 0.2 mg of each derivative per ml of toluene.

SAMPLING SYSTEM

Two glass impingers in series, each containing 40 ml of 2N HCl-2,4 dinitrophenylhydrazine, are used to collect exhaust samples for the analysis of the aldehydes and ketones. A flow schematic of the sample collection system is shown in Figure 1. The two impingers together trap approximately 98 percent of the carbonyl compounds. The temperature of the gas stream is monitored by a thermocouple immediately prior to the dry gas meter. The dry gas meter determines the total flow through the impinger during a given driving cycle. The sample pump is capable of pulling a flow rate of 4 l/min. A drier is included to prevent condensation in the pump, flowmeter, dry gas meter, etc. The flowmeter allows continuous monitoring of the sample flow to insure proper flow rates during the sampling. The Teflon line connecting the CVS and the solenoid valve is heated to 170°F in order to prevent water from condensing in the line. Several views of the sampling system are shown in Figure 2.

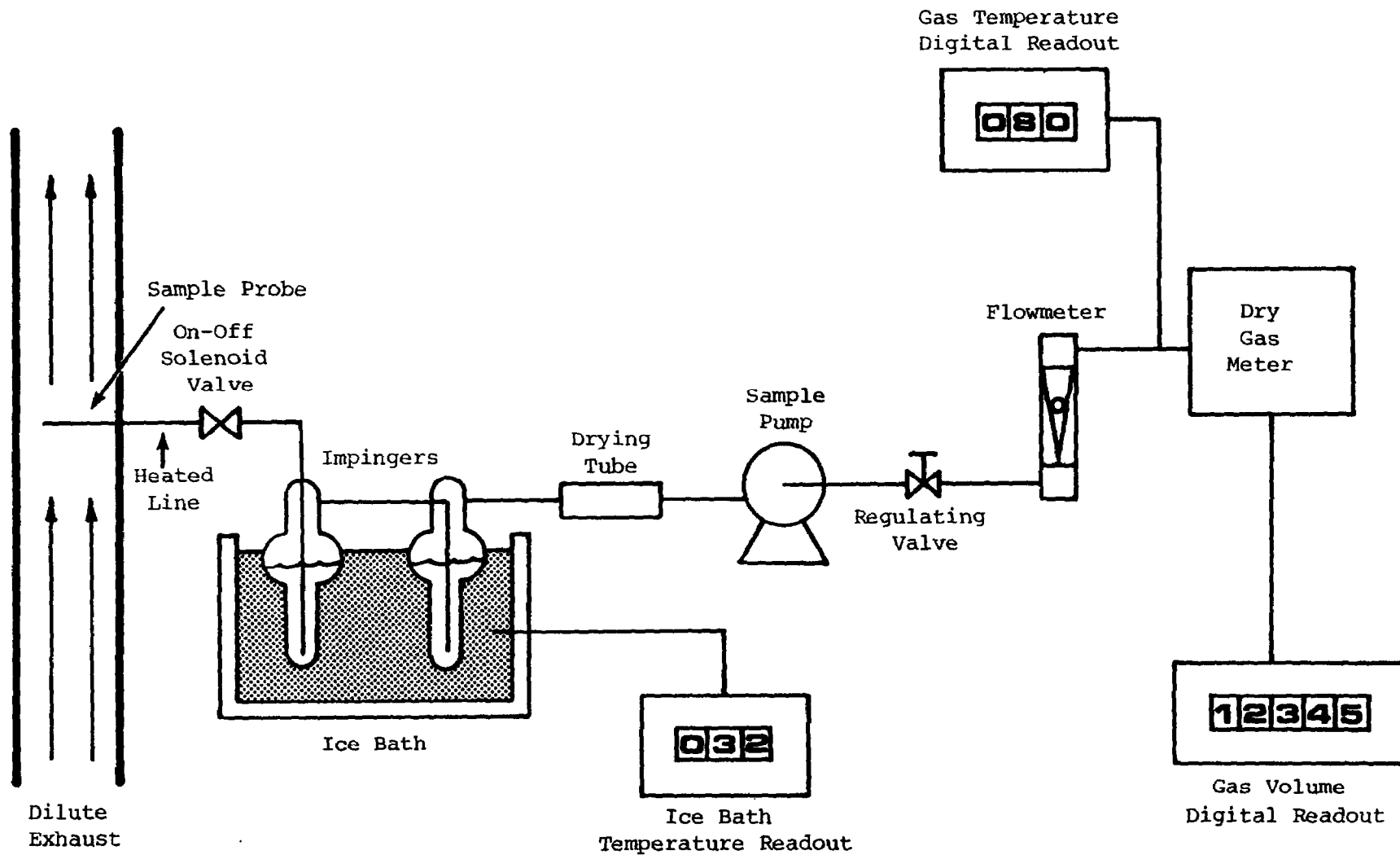
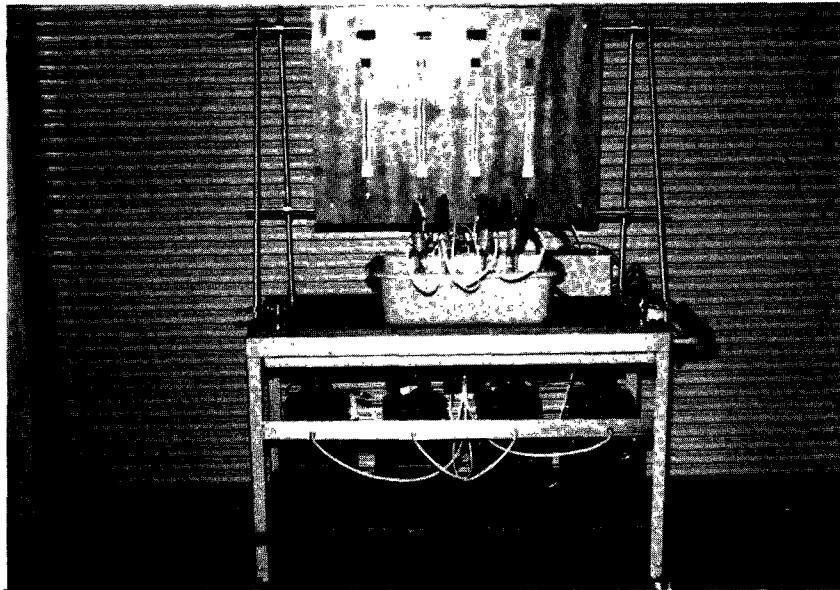
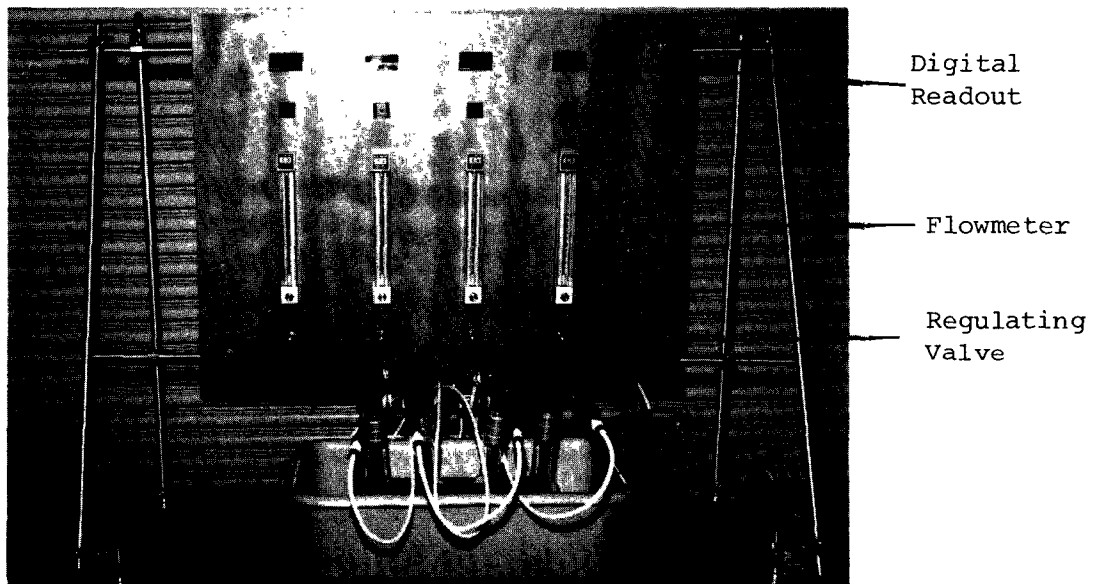


Figure 1. Aldehyde and Ketone sample collection flow schematic.



Front View

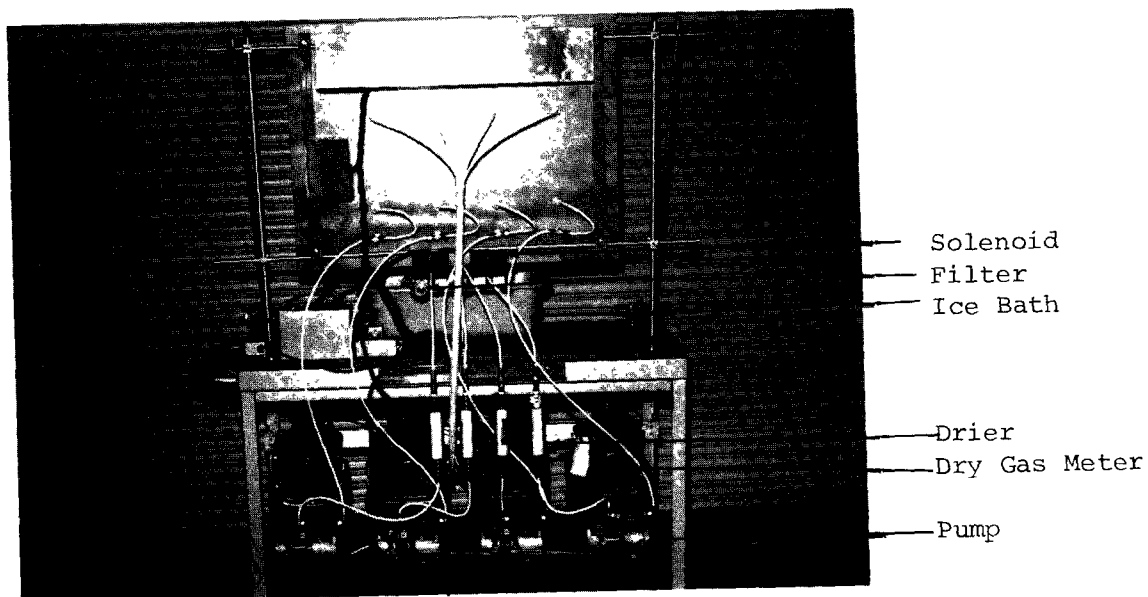


Close-Up of Upper Front

Figure 2. Aldehyde and Ketone sampling system.



Close-up of Impingers (Side View)



Rear View

Figure 2 (Cont'd). Methanol sampling system.

ANALYTICAL PROCEDURE

The analysis of the aldehydes (formaldehyde, acetaldehyde, isobutyraldehyde, crotonaldehyde, hexanaldehyde, and benzaldehyde) and of the ketones (acetone and methylethylketone) in dilute exhaust is accomplished by collecting these carbonyl compounds in a hydrochloric acid (HCl)/2,4 dinitrophenylhydrazine (DNPH) solution as their 2,4 dinitrophenylhydrazone derivatives. The derivatives are removed from the HCl/DNPH absorbing solution by filtration and/or extractions with pentane. The filtered precipitate and the pentane extracts are combined and the volatile solvents are removed. The remaining extract contains the phenylhydrazone derivatives. The derivatives are then dissolved in a quantitative volume of toluene containing a known amount of anthracene as an internal standard. This solution is analyzed by injecting a small volume of the solution into a gas chromatograph equipped with dual flame ionization detectors. From this analysis and the measured volume of exhaust sampled, the concentration of the carbonyl compounds in exhaust can be determined. The analysis flow schematic for the aldehydes and ketones is shown in Figure 3. A detailed description of the procedure follows.

The aldehydes and ketones are trapped in solution by bubbling a known volume of dilute exhaust through two glass impingers connected in series, with each impinger containing 40 ml of a 2N HCl solution saturated with DNPH. The sampling temperature and barometric pressure are recorded during this bubbling period. The carbonyl compounds in the exhaust react with the DNPH to form slightly soluble or insoluble 2,4 dinitrophenylhydrazone derivatives. The two impingers together collect 98 percent + of the carbonyls that are present in the exhaust. The impingers are removed from the sampling cart and are allowed to stand at room temperature for at least one hour before proceeding to the filtration and extraction steps. Figure 4 shows two impingers containing the HCl/DNPH absorbing solution after being removed from the sampling cart.

Under normal operating conditions the contents of the two impingers are combined and analyzed as one sample. If either of the two impingers contain a precipitate they are first subjected to a filtration step. If no precipitate is present this filtration step is omitted and the extraction step, described later in the procedure, is the first step.

For the filtration step the contents of the two impingers are poured through a fritted glass filter into a flask under vacuum (Figure 5). The two impingers are rinsed with small portions of deionized water. This wash water is also poured through the fritted glass filter. The precipitate in the filter is then washed with a few ml of deionized water. The fritted filter is then removed from the flask containing the 80 ml of absorbing reagent and the water washings. The flask is then set aside for the extraction step. The fritted glass filter containing the precipitate is connected to a dry flask. The two impingers that had previously contained the filtered precipitate are then each washed with small portions of methylene chloride. The methylene chloride dissolves any solid residue which was not removed by the water wash. These methylene chloride washings are poured into the fritted glass filter containing the precipitate. After

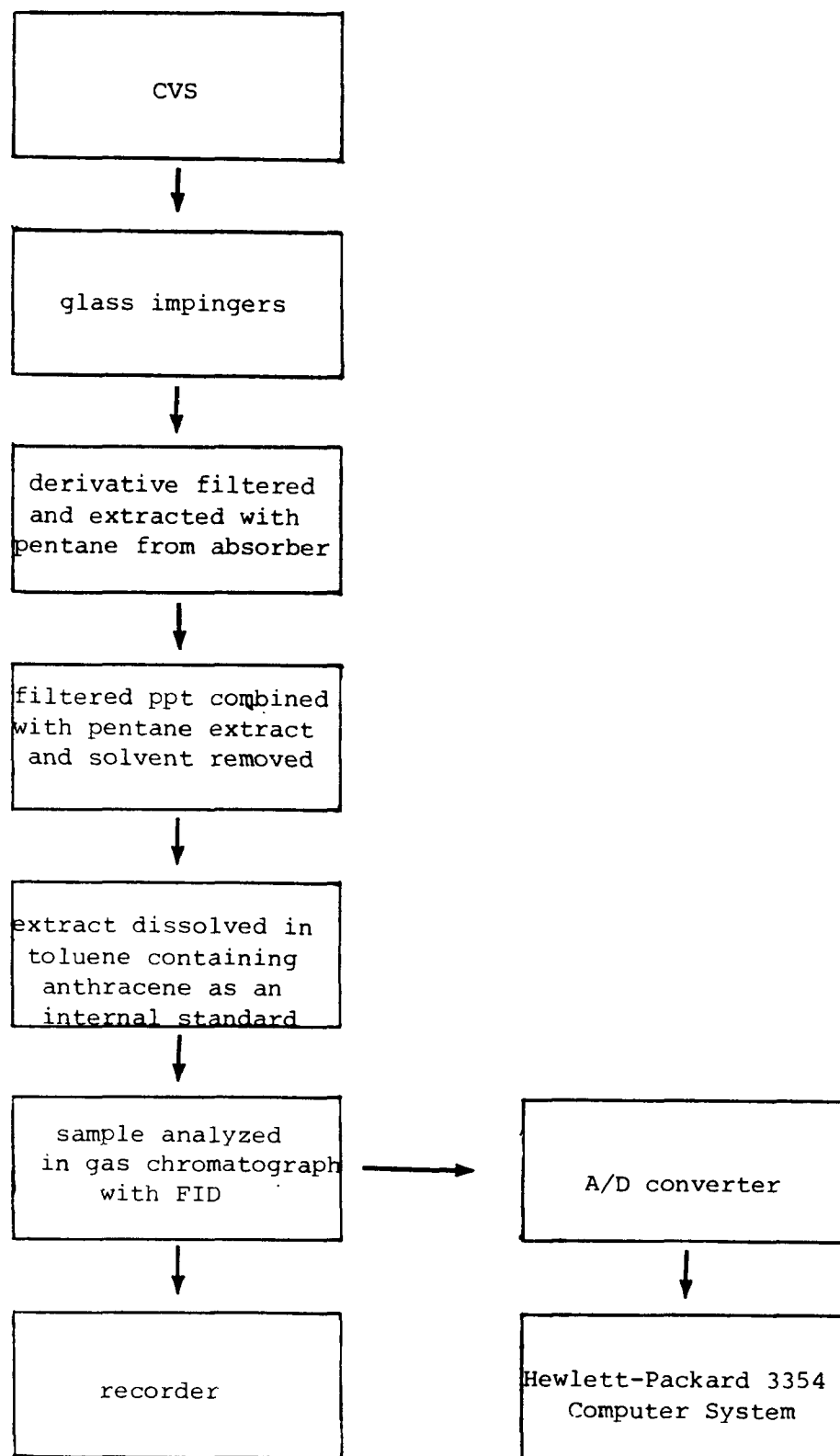


Figure 3. Aldehyde and Ketone analysis flow schematic.

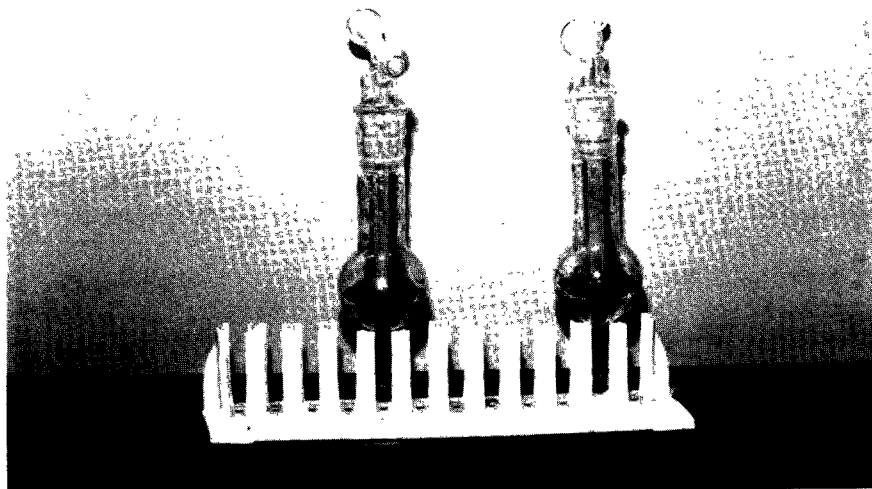


Figure 4. Impingers containing HCl/DNPH absorbing solution.



Figure 5. Filtration of absorbing solution.

the precipitate has been dissolved by the methylene chloride, a vacuum is applied to the flask and the methylene chloride solution is pulled through the filter into the flask. Another small amount of methylene chloride is poured through the filter into the flask to wash the filter. The methylene chloride solution is now saved until the extraction step is complete.

The extraction step is carried out as follows. The contents of the two impingers (if no precipitate is present) are transferred to a 250 ml separatory funnel. The impingers are each washed with small portions of deionized water which is also added to the separatory funnel. If a precipitate was found in the impingers the contents of the flask containing the filtered absorbing reagent and the water washings from the filtration step are transferred quantitatively to a 250 ml separatory funnel. The flask is washed with a small portion of water, and this water is added to the separatory funnel. Forty ml of pentane is now added to the separatory funnel containing the 80 ml of absorbing reagent and water washings. The funnel is stoppered and shaken for five minutes in an automatic shaker, Figure 6. The shaker is stopped and the funnel is vented. After the two phases are allowed to separate, the lower phase is collected in a second separatory funnel. The remaining phase is transferred to a third 250 ml separatory funnel. A second 40 ml portion of pentane is added to the already once extracted absorbing solution. The funnel is again stoppered, shaken for 5 minutes and vented. After the phases have separated the lower phase is again collected in another separatory funnel. The upper or pentane layer is combined with the pentane layer from the first extraction. A third 40 ml portion of pentane is added to the twice extracted absorbing solution and the extraction process repeated. After the third extraction the lower layer is discarded and the pentane layer is combined with the pentane layers from the first two extractions. Any absorbing solution which might have been accidentally transferred with the pentane layers is drained off. Deionized water (25-50 ml) and sodium bicarbonate (1/4-1/2 gram) is added to the 250 ml separatory funnel containing the 120 ml of pentane extract. The funnel is stoppered and manually shaken for 30 seconds. The phases are allowed to separate and the lower water phase is drained off. Another 25 ml of deionized water is added and the shaking is repeated. After the phases have separated, the water is drained off insuring that all traces of water are removed. The contents of the funnel are then combined with the methylene chloride solution which was saved from the filtration step.

The flask containing the methylene chloride solution and the pentane extracts is then placed in a vacuum oven, Figure 7, operating at 50-60°C and 65" water vacuum until the pentane and methylene chloride have been removed. At this time only the dried phenylhydrazones remain.

Each time a series of samples are collected a blank containing 80 ml of HCl/DNPH solution is extracted and dried in the same manner as the samples. This accounts for any aldehydes or interfering compounds which might be found in the reagents used for extraction.

Two ml of toluene which contain a quantitative amount of anthracene (0.05 mg/ml toluene) as an internal standard is pipetted into the flask

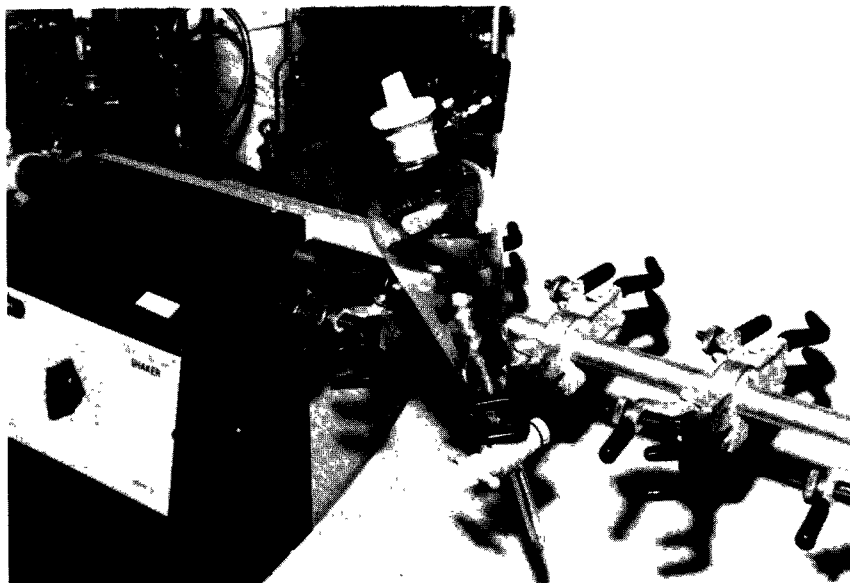


Figure 6. Automatic shaker.

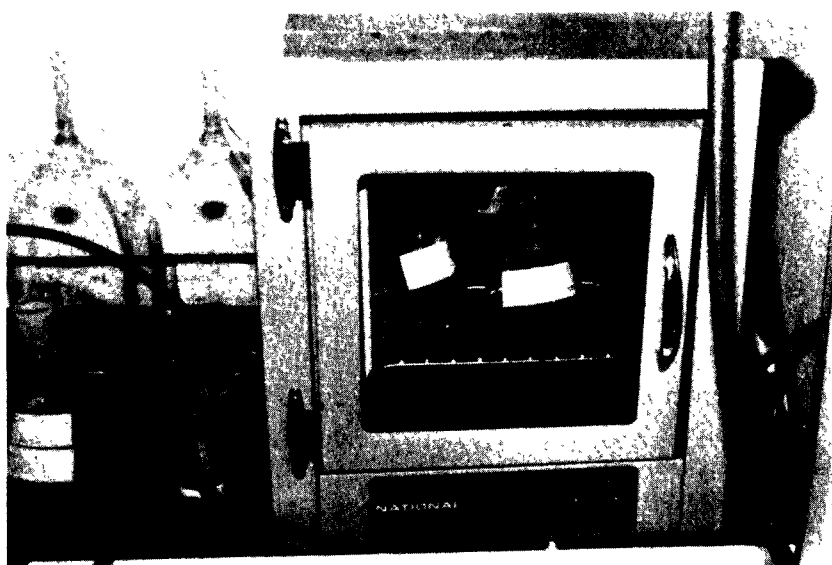


Figure 7. Vacuum oven.

containing the dried phenylhydrazone derivatives. The flask is then placed in a sonic bath until all of the residue is dissolved. After the precipitate has dissolved, the solution is transferred to a 1/2 dram vial. (See Figure 8.) At this point the derivative is ready for injection into the gas chromatograph system.

The gas chromatograph system used to analyze the toluene solution containing the phenylhydrazone derivatives is shown in Figure 9. The system consists of a Varian 1700 GC, an A/D converter, and a recorder. The GC is equipped with dual columns and dual flame ionization detectors with a single differential amplifier. The columns consist of 24 x 1/8 inch O.D. stainless steel tubing packed with 6.7 percent Dexsil (polycarboranesiloxane) 300 GC on DMCS treated and acid washed, 60/80 mesh Chromosorb G. The carrier gas is helium which flows through the columns at a rate of 40 ml/min. The optimum hydrogen and air flow rates are 500 ml/min and 35 ml/min respectively. The column temperature, after injection of the sample, is programmed from 120°C to 300°C at 8° a minute. In a chromatogram of a standard sample (Figure 10) containing anthracene and the phenylhydrazone derivatives of formaldehyde, acetaldehyde, acetone, iso-butyraldehyde, methylethylketone, crotonaldehyde, hexanaldehyde, and benzaldehyde, the first peak eluted is toluene followed by anthracene, and then the derivatives of formaldehyde, acetaldehyde, acetone, iso-butyraldehyde, methylethylketone, crotonaldehyde, hexanaldehyde and benyaldehyde. The methylethylketone derivative is added to the list of derivatives in order to name an unknown peak found in some of the exhaust samples. Data obtained from the five repetitive injections of the standard derivatives in toluene showed a maximum standard deviation of 4.56 percent for benzaldehyde and a minimum standard deviation of 0.87 percent for formaldehyde. The computer printout of the standard, Figure 10, is shown in Figure 11. This printout gives the retention time, area, and the name of each peak. The printout also gives the concentration of each of the derivatives in mg/ml. This concentration is calculated by the computer from response factors which are determined daily. Each day a standard containing known amounts of the derivatives and anthracene is injected into the GC. From the anthracene and derivative areas the computer calculates a response factor F. These F factors are used in all subsequent runs during the day to determine the concentration of the derivatives. This response is calculated from the following equation:

$$\text{Response Factor (F)} = \frac{\text{Anthracene Area}}{\text{Derivative Area}} \times \frac{\text{mg/ml Derivative}}{\text{mg/ml Anthracene}}$$

Typical response factors for each of the derivatives are listed below:

| FACTOR | NAME |
|--------|-------------------|
| 1.0000 | ANTHRACENE |
| 3.1043 | FORMALDEHYDE |
| 2.7736 | ACETALDEHYDE |
| 2.2366 | ACETONE |
| 2.4160 | ISO-BUTYRALDEHYDE |
| 2.3332 | METHYLETHYLKETONE |
| 3.4174 | CROTONALDEHYDE |



Figure 8. 1/2 dram vials.

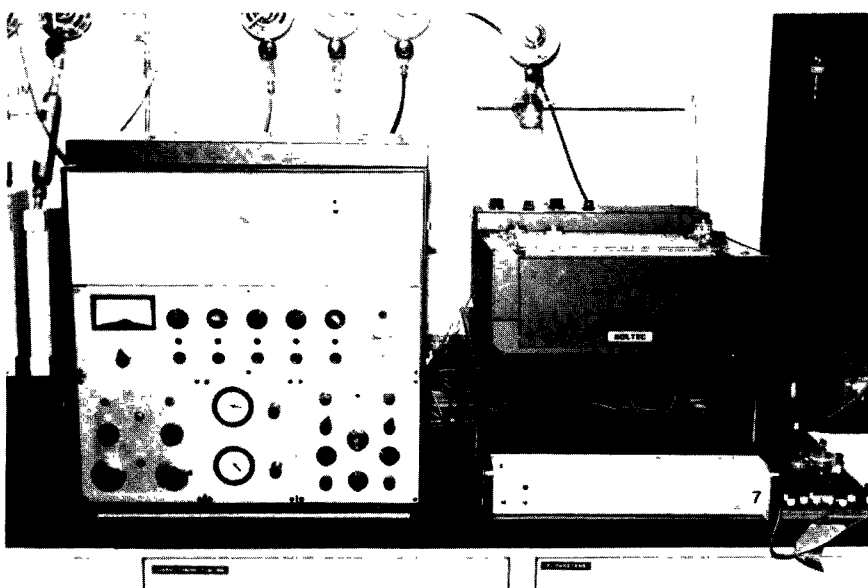


Figure 9. Aldehyde and ketone analytical system.

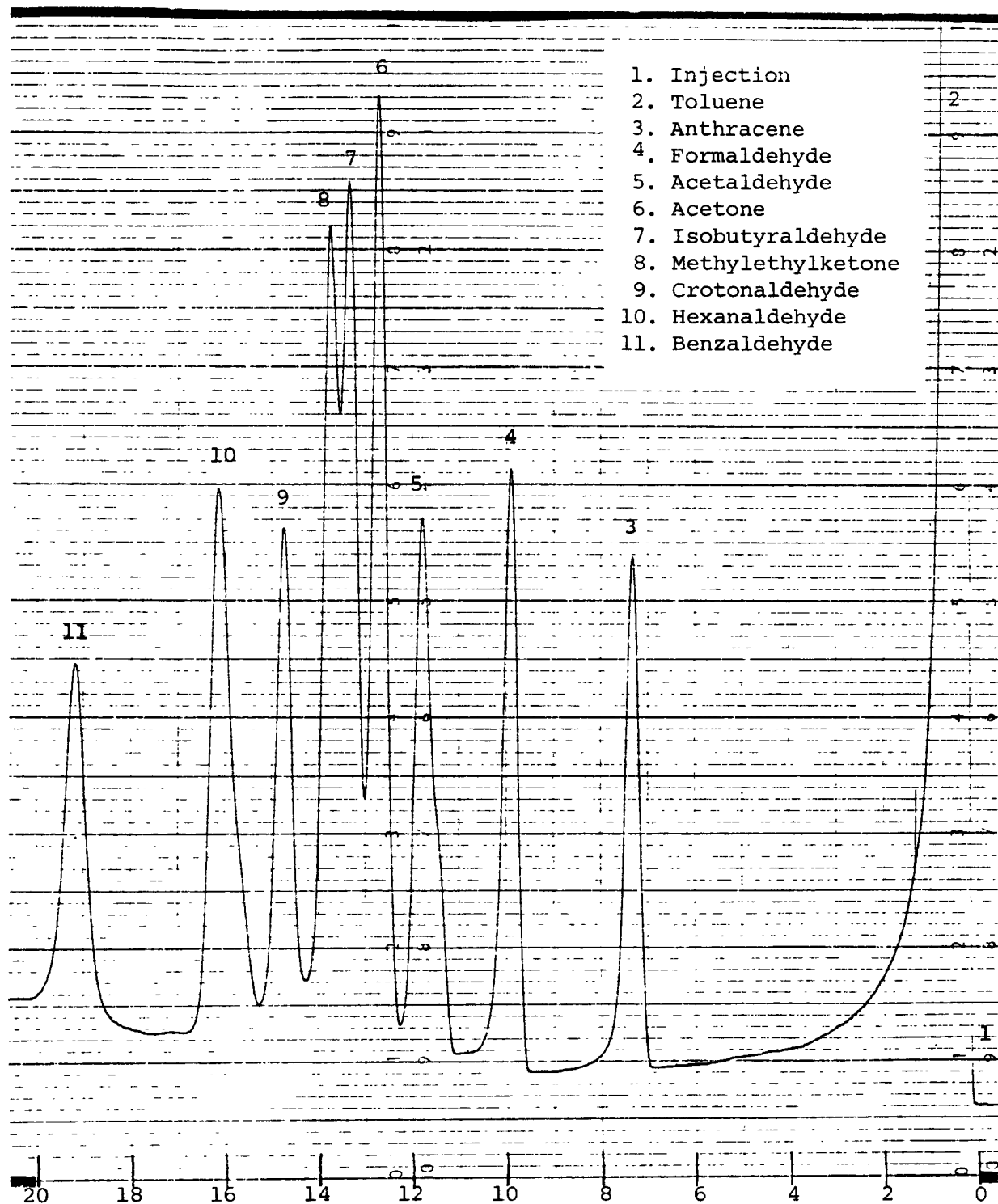


Figure 10. Chromatogram of standard.

REPORT: 14.11 CHANNEL: 11

SAMPLE: RCI INJECTED AT 11:18:27 ON MAR 1, 1978

ISTD METHOD: DNPH11

ACTUAL RUN TIME: 30.008 MINUTES

ISTD-RATIO: .0500 R MG/ML STD-AMT: .0500 SAMP-AMT: 1.0000

ENDED NOT ON BL

| RT | AREA | MG/ML | NAME |
|--------------|----------|---------|---------------------|
| 7.26 | 9638 BB | | &ANTHRACENE |
| 9.81 | 11159 BB | .190 | #FORMALDEHYDE |
| 11.71 | 13355 BV | .202 | #ACETALDEHYDE |
| 12.64 | 17898 VV | .211 | #ACETONE |
| 13.28 | 16448 VV | .210 | #ISOBUTYRALDEHYDE |
| 13.70 | 16469 VV | .207 | #MEK |
| 14.69 | 11167 VV | .208 | #CROTONALDEHYDE |
| 16.08 | 15988 VV | .206 | #HEXANALDEHYDE |
| 17.08 | 74 VB | 3.8E- 4 | |
| 19.08 | 10525 BB | .215 | #BENZALDEHYDE |
| 25.04 | 187 BB | 9.7E- 4 | |
| TOTAL AREA = | | 122909 | TOTAL MG/ML = 1.650 |

Figure 11. Computer printout of standard.

| | |
|--------|---------------|
| 2.3428 | HEXANALDEHYDE |
| 2.9329 | BENZALDEHYDE |

When the response factor is known a concentration in mg/ml for each of the derivatives can be found. This concentration, along with the volume of sampled exhaust is then used to calculate the concentration of the carbonyl compounds in exhaust. Figures 12 and 13 show a typical sample chromatogram and accompanying printout respectively.

CALCULATIONS

This procedure has been developed to provide the user with the concentrations of the aldehydes (formaldehyde, acetaldehyde, isobutyraldehyde, crotonaldehyde, hexanaldehyde, and benzaldehyde) and ketones (acetone and methylethylketone) in exhaust. The results will be expressed in $\mu\text{g}/\text{m}^3$ of exhaust and ppm for each carbonyl compound. The equations for determining the concentrations in $\mu\text{g}/\text{m}^3$ and ppm are derived in the following manner.

The first step is to correct the volume of exhaust sampled to a standard temperature, 68°F and pressure, 29.92" Hg, by use of the equation

$$\frac{P_{\text{exp}} \times V_{\text{exp}}}{T_{\text{exp}}} = \frac{P_{\text{corr}} \times V_{\text{corr}}}{T_{\text{corr}}}$$

V_{exp} = experimental volume of gas sampled in ft^3
 V_{corr} = volume of gas sampled in ft^3 corrected to 68°F and 29.92" Hg
 P_{exp} = experimental barometric pressure
 P_{corr} = 29.92" Hg
 T_{exp} = experimental temperature in °F + 460
 T_{corr} = 68°F + 460 = 528°R

Solving for V_{corr} gives:

$$V_{\text{corr}} = \frac{P_{\text{exp}} ("Hg) \times V_{\text{exp}} (\text{ft}^3) \times 528^\circ\text{R}}{T_{\text{exp}} (^\circ\text{R}) \times 29.92" \text{ Hg}}$$

The next step converts the volume from cubic feet to cubic meters by use of the conversion factor 1 cubic meter is equal to 35.31 cubic feet.

$$V_{\text{corr}}(\text{m}^3) = \frac{P_{\text{exp}} ("Hg) \times V_{\text{exp}} (\text{ft}^3) \times 528^\circ}{T_{\text{exp}} \times 29.92" \text{ Hg} \times 35.31 \text{ ft}^3/\text{m}^3}$$

(Equation 1)

The next step converts the mg/ml of derivative determined by the computer to mg of carbonyl collected in the two impingers. To obtain mg of derivative, the concentration (from the computer printout) in mg/ml is multiplied by the volume of toluene used to dissolve the solid extract.

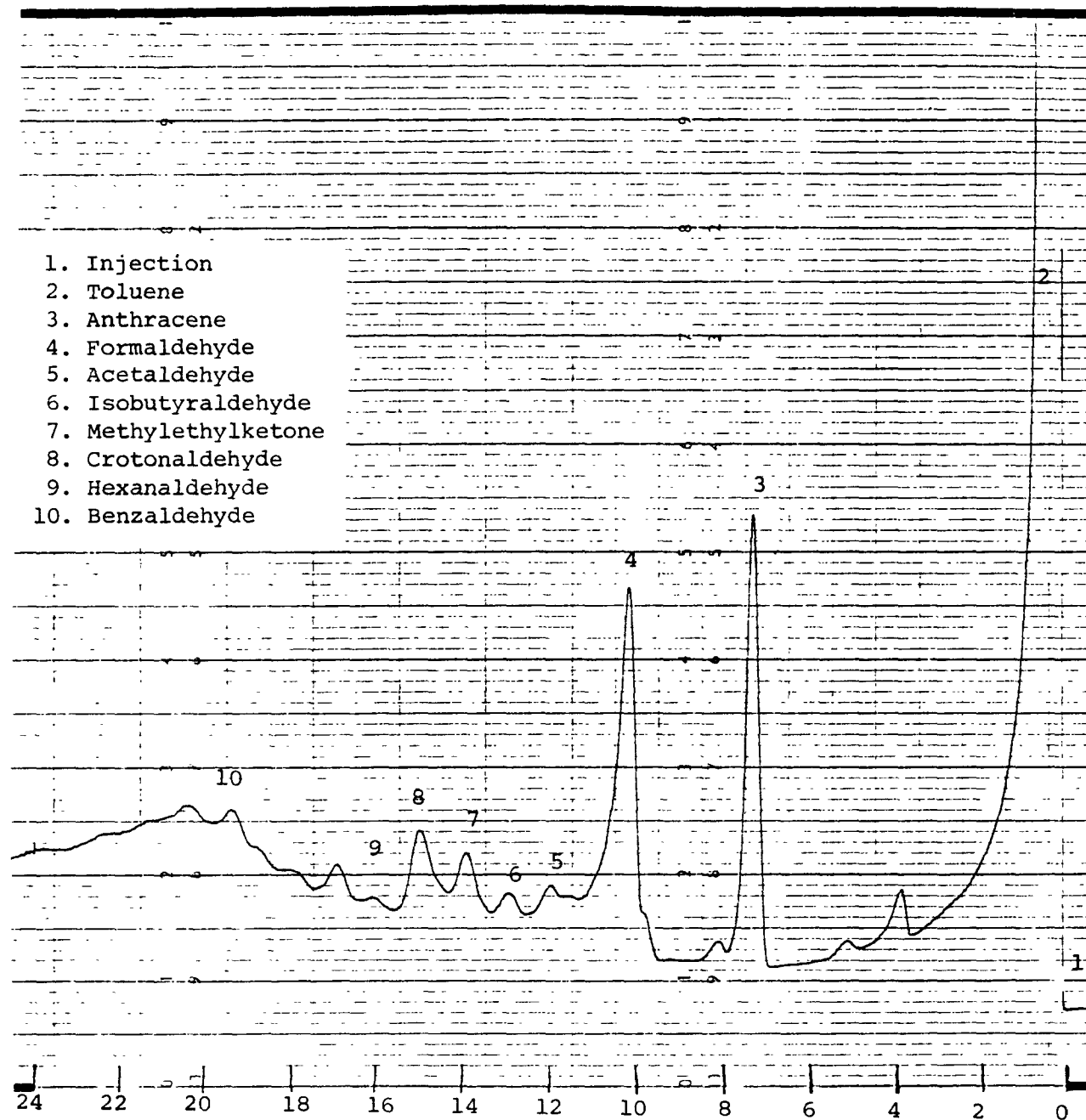


Figure 12. Sample chromatogram.

REPORT: 20 CHANNEL: 11

SAMPLE: RCI INJECTED AT 15:41:05 ON MAR 1, 1978

ISTD METHOD: DNP11

ACTUAL RUN TIME: 39.317 MINUTES

ISTD-RATIO: .050, R MG/ML STD-AMT: .0500 SAMP-AMT: 1.0000

| RT | AREA | MG/ML | NAME |
|--------------|---------|---------|--------------------|
| 7.15 | 8604 BV | | ANTHRACENE |
| 7.95 | 435 VB | .003 | |
| 10.03 | 8877 BB | .186 | FORMALDEHYDE |
| 11.88 | 575 BV | .010 | ACETALDEHYDE |
| 12.83 | 463 VV | .007 | ISO-BUTYRALDEHYDE |
| 13.78 | 1594 VV | .022 | MEK |
| 14.85 | 2630 VV | .053 | CROTONALDEHYDE |
| 15.95 | 146 VV | .002 | HEXANALDEHYDE |
| 16.77 | 675 VV | .004 | |
| 19.20 | 648 VB | .012 | BENZALDEHYDE |
| 20.23 | 1912 BV | .011 | |
| 23.52 | 217 VV | .001 | |
| 25.03 | 13 VB | 7.6E- 5 | |
| 25.48 | 84 BB | 4.9E- 4 | |
| TOTAL AREA = | | 26874 | TOTAL MG/ML = .310 |

PROCESSED DATA FILE: *PRC11 RAW DATA FILE: *RAW11

A-24

Figure 13. Computer printout of sample.

$$\text{mg derivative} = \text{Conc}_{\text{Der}} (\text{mg/ml}) \times \text{Vol}_{\text{Tol}} (\text{ml})$$

To find mg of carbonyl compound per sample the mg of derivative are multiplied by the ratio of the molecular weight of the carbonyl derivative over the molecular weight of its phenylhydrazone derivative.

$$\begin{aligned} \text{mg carbonyl} &= \text{mg derivative} \times \frac{\text{mol. wt. carbonyl}}{\text{mol. wt. derivative}} \\ &= \text{Conc}_{\text{Der}} (\text{mg/ml}) \times \text{Vol}_{\text{Tol}} (\text{ml}) \times \frac{\text{mol. wt. carbonyl}}{\text{mol. wt. derivative}} \end{aligned}$$

To obtain the number of μg of carbonyl compound the mg of carbonyl are multiplied by the conversion factor, 1000 $\mu\text{g}/\text{mg}$

$$\begin{aligned} \mu\text{g carbonyl} &= \text{Conc}_{\text{Der}} (\text{mg/ml}) \times \text{Vol}_{\text{Tol}} (\text{ml}) \times \frac{\text{Mol. Wt. carbonyl}}{\text{Mol. Wt. derivative}} \\ &\quad \times 1000 \mu\text{g}/\text{mg} \end{aligned}$$

(Equation 2)

The concentration of the carbonyl compound in exhaust can now be found in $\mu\text{g}/\text{m}^3$ by dividing equation 2 by equation 1.

$$\begin{aligned} \mu\text{g carbonyl}/\text{m}^3 &= \frac{\text{Conc}_{\text{Der}} (\text{mg/ml}) \times \text{Vol}_{\text{Tol}} (\text{ml}) \times \text{mol. wt. carbonyl}}{P_{\text{exp}} (\text{"Hg}) \times V_{\text{exp}} (\text{ft}^3) \times 528^\circ} \\ &\quad \times \frac{1000 \mu\text{g}/\text{mg} \times T_{\text{exp}} (^\circ\text{R}) \times 29.29 \text{ "Hg} \times 35.31 \text{ ft}^3/\text{m}^3}{\text{mol. wt. derivative}} \end{aligned}$$

(Equation 3)

To find the concentration of each carbonyl compound in ppm, the densities of carbonyls are needed. At 29.92" Hg and 32°F, one mole of gas occupies 22.4 liters. This volume is corrected to 68°F from the equation.

$$\frac{V}{T} = \frac{V_1}{T_1}$$

$$V_1 = 22.4$$

$$T_1 = 32^\circ\text{F} + 460 = 492^\circ\text{R}$$

$$V = \text{volume at } 68^\circ\text{F}$$

$$T = 68^\circ\text{F} + 460 = 528^\circ\text{R}$$

Solving for V gives:

$$V = \frac{V_1 \times T}{T_1} = \frac{22.4 \times 528}{492} = 24.04\text{l}$$

Since one mole of gas occupies 22.04ℓ at 68°F, the density can be found in g/ℓ by dividing the molecular weight in g/mole by 24.04 ℓ/mole

$$\text{den (g/ℓ)} = \frac{\text{mol. wt. (g/mole)}}{24.04\text{ℓ/mole}}$$

The density in μg/ml can be found by converting g to μg and μg and ℓ to ml as follows:

$$\text{den } \mu\text{g/ml} = \frac{\text{mol. wt. g/mole}}{24.04\text{ℓ/mole}} \times \frac{1 \times 10^6 \mu\text{g/g}}{1 \times 10^3 \text{ml/ℓ}} \times \frac{\text{mol. wt.} \times 1000}{24.04}$$

(Equation 4)

To obtain the concentration of each carbonyl in ppm, the concentration in μg/m³ is divided by the density in μg/ml

$$\text{ppm} = \mu\text{g/m}^3 \div \mu\text{g/ml} = \frac{\text{ml}}{\text{m}^3}$$

Using Equations 3 and 4 gives the ppm concentration in the form of the raw data.

$$\begin{aligned} \text{ppm} &= \frac{\text{Conc}_{\text{Der}} (\text{mg/ml}) \times \text{Vol}_{\text{Tol}} (\text{ml}) \times \text{mol. wt. carbonyl} \times 1000 \mu\text{g/ml}}{\text{P}_{\text{exp}} (" \text{Hg}) \times \text{V}_{\text{exp}} (\text{ft}^3) \times 528^\circ \times \text{mol. wt. derivative} \\ &\quad \times \frac{\text{T}_{\text{exp}} (^\circ \text{R}) \times 29.92" \text{ Hg} \times 35.31 \text{ ft}^3/\text{m}^3 \times 24.04 \text{ ℓ/mole}}{\text{mol. wt. carbonyl} \times 1000} \\ &= \frac{\text{Con}_{\text{Der}} (\text{mg/ml}) \times \text{Vol}_{\text{Tol}} (\text{ml}) \times \text{T}_{\text{exp}} (^\circ \text{R}) \times 29.92" \text{ Hg}}{\text{P}_{\text{exp}} (" \text{Hg}) \times \text{V}_{\text{exp}} (\text{ft}^3) \times 528^\circ} \\ &\quad \times \frac{35.35 \text{ ft}^3/\text{m}^3 \times 24.04 \text{ ℓ/mole}}{\text{mol. wt. derivative}} \end{aligned}$$

(Equation 5)

At this point, the concentration can be express in μg/m³ (Equation 3) and ppm (Equation 5) at 68°F and 29.92" Hg from the raw data.

Hewlett-Packard Calculations

In order to insure maximum turnaround in a minimum time period, two Hewlett-Packard 65 programs were developed. One calculates the aldehyde and ketone concentrations in μg/m³ from the raw data and phenylhydrazone derivative concentrations (from computer printout). The other program calculates the concentrations in ppm from the concentrations in μg/m³. These programs are presented in Figures 14 and 15.

HP-65 Program Form

Title _____ Page _____ of _____

SWITCH TO W-PRGM PRESS **F** PRGM TO CLEAR MEMORY

| KEY ENTRY | CODE SHOWN | COMMENTS | KEY ENTRY | CODE SHOWN | COMMENTS | REGISTERS |
|-----------|------------|---|-----------|------------|---|-----------|
| LBL | 23 | | X | 71 | | R1 _____ |
| A | 11 | Input sample vol., ft ³ | R/S | 84 | Out $\mu\text{g}/\text{m}^3$; In mg/ml | R2 _____ |
| 0 | | | RCL2 | 3402 | | R3 _____ |
| . | 83 | | X | 71 | | R4 _____ |
| 0 | | | 0 | | | R5 _____ |
| 0 | | | . | 83 | | R6 _____ |
| 0 | | | 2 | | | R7 _____ |
| 5 | | | 8 | | | R8 _____ |
| X | 71 | | 6 | | | R9 _____ |
| R/S | 84 | Input Barometer, "Hg | X | 71 | | LABELS |
| X | 71 | | R/S | 84 | Out $\mu\text{g}/\text{m}^3$; In mg/ml | A _____ |
| STO1 | 3301 | | RCL2 | 3402 | | B _____ |
| R/S | 84 | Input sample Temp °F | X | 71 | | C _____ |
| 4 | | | 0 | | | D _____ |
| 6 | | | . | 83 | | E _____ |
| 0 | | | 2 | | | 0 _____ |
| + | 61 | | 8 | | | 1 _____ |
| RCL1 | 3401 | | 6 | | | 2 _____ |
| g x y | 3507 | | X | 71 | | 3 _____ |
| 20 ÷ | 81 | | R/S | 84 | Out $\mu\text{g}/\text{m}^3$; In mg/ml | 4 _____ |
| g 1/x | 3504 | | RCL2 | 3402 | | 5 _____ |
| R/S | 84 | Input Vol. Toluene, ml | X | 71 | | 6 _____ |
| X | 71 | | 0 | | | 7 _____ |
| STO2 | 3302 | | . | 83 | | 8 _____ |
| R/S | 84 | In mg/ml | 2 | | | 9 _____ |
| RCL2 | 3402 | | 8 | | | FLAGS |
| X | 71 | | 0 | | | 1 _____ |
| 0 | | | X | 71 | | 2 _____ |
| . | 83 | | R/S | 84 | Out $\mu\text{g}/\text{m}^3$; In mg/ml | |
| 0.1 | | | RCL2 | 3402 | | |
| 4 | | | X | 71 | | |
| 3 | | | 0 | | | |
| X | 71 | | . | 83 | | |
| R/S | 84 | Out $\mu\text{g}/\text{m}^3$; In mg/ml | 3 | | | |
| RCL 2 | 3402 | | 5 | | | |
| X | 71 | | 9 | | | |
| 0 | | | X | 71 | | |
| . | 83 | | R/S | 84 | Out $\mu\text{g}/\text{m}^3$; In mg/ml | |
| 1 | | | RCL2 | 3402 | | |
| 0.9 | | | X | 71 | | |
| 6 | | | 0 | | | |
| X | 71 | | . | 83 | | |
| R/S | 84 | Out $\mu\text{g}/\text{m}^3$; In mg/ml | 3 | | | |
| RCL 2 | 3402 | | 7 | | | |
| X | 71 | | 1 | | | |
| 0 | | | X | 71 | | |
| . | 83 | | R/S | 84 | Out $\mu\text{g}/\text{m}^3$ | |
| 2 | | | RTN | 24 | | |
| 4 | | | | | | |
| 4 | | | | | | |

HP-65C-6

TO RECORD PROGRAM INSERT MAGNETIC CARD WITH SWITCH SET AT W PRGM

Figure 14. Aldehyde and Ketone concentrations in $\mu\text{g}/\text{m}^3$.

HP-65 User Instructions

Title _____ Page _____ of _____
 Programmer _____ Date _____



| STEP | INSTRUCTIONS | INPUT DATA/UNITS | KEYS | OUTPUT DATA/UNITS |
|----------------|---|------------------|--|-------------------|
| 0 ₁ | Switch to on; switch to run | | <input type="button" value="ON"/> <input type="button" value="RUN"/> | |
| 0 ₂ | Feed card in from right to left | | <input type="button" value="CARD"/> <input type="button" value="PUNCH"/> | |
| 0 ₃ | Initialize | | <input type="button" value="f"/> <input type="button" value="REG"/> | |
| 0 ₄ | Set decimal place | | <input type="button" value="DSP"/> <input type="button" value="2"/> | |
| 1 | Input - sample volume, ft ³ | | <input type="button" value="A"/> <input type="button" value="0"/> | |
| 2 | Input - barometric pressure, "Hg | | <input type="button" value="R/S"/> <input type="button" value="0"/> | |
| 3 | Input - Sample Temperature, °F | | <input type="button" value="R/S"/> <input type="button" value="0"/> | |
| 4 | Input - Volume Toluene, ml | | <input type="button" value="R/S"/> <input type="button" value="0"/> | |
| 5 | Input - Conc. Formaldehyde Der, mg/ml | | <input type="button" value="R/S"/> <input type="button" value="0"/> | |
| | Output - Conc. Formaldehyde μg/m ³ | | <input type="button" value="0"/> <input type="button" value="0"/> | |
| 6 | Input - Conc. Acetaldehyde Der, mg/ml | | <input type="button" value="R/S"/> <input type="button" value="0"/> | |
| | Output - Conc. Acetaldehyde μg/m ³ | | <input type="button" value="0"/> <input type="button" value="0"/> | |
| 7 | Input - Conc. Acetone Der, mg/ml | | <input type="button" value="R/S"/> <input type="button" value="0"/> | |
| | Output - Conc. Acetone, μg/m ³ | | <input type="button" value="0"/> <input type="button" value="0"/> | |
| 8 | Input - Conc. Isobutyraldehyde Der, mg/ml | | <input type="button" value="R/S"/> <input type="button" value="0"/> | |
| | Output - Conc. Isobutyraldehyde, μg/m ³ | | <input type="button" value="0"/> <input type="button" value="0"/> | |
| 9 | Input - Conc. methylethylketone Der, mg/ml | | <input type="button" value="R/S"/> <input type="button" value="0"/> | |
| | Output - Conc. methylethylketone, μg/m ³ | | <input type="button" value="0"/> <input type="button" value="0"/> | |
| 10 | Input - Conc. Crotonaldehyde Der, mg/ml | | <input type="button" value="R/S"/> <input type="button" value="0"/> | |
| | Output - Conc. Crotonaldehyde, μg/m ³ | | <input type="button" value="0"/> <input type="button" value="0"/> | |
| 11 | Input - Conc. Hexanaldehyde Der, mg/ml | | <input type="button" value="R/S"/> <input type="button" value="0"/> | |
| | Output - Conc. Hexanaldehyde, μg/m ³ | | <input type="button" value="0"/> <input type="button" value="0"/> | |
| 12 | Input - Conc. Benzaldehyde Der, mg/ml | | <input type="button" value="R/S"/> <input type="button" value="0"/> | |
| | Output - Conc. Benzaldehyde μg/m ³ | | <input type="button" value="0"/> <input type="button" value="0"/> | |
| | | | <input type="button" value="RTN"/> <input type="button" value="0"/> | |
| | | | <input type="button" value="f"/> <input type="button" value="0"/> | (cont'd.) |

Figure 14 (Cont'd). Aldehyde and Ketone concentrations in μg/ml.

HP-65 User Instructions

Title _____ Page _____ of _____

Programmer _____ Date _____



| STEP | INSTRUCTIONS | INPUT DATA/UNITS | KEYS | OUTPUT DATA/UNITS |
|----------------|--|------------------|---|-------------------|
| 0 ₁ | Switch to on; switch to run | | <input type="text"/> <input type="text"/> | |
| 0 ₂ | Feed card in from right to left | | <input type="text"/> <input type="text"/> | |
| 0 ₃ | Initialize | | <input type="text"/> f <input type="text"/> REG | |
| 0 ₄ | Set decimal place | | <input type="text"/> DSP <input type="text"/> 2 | |
| 1 | Input-conc Formaldehyde, $\mu\text{g}/\text{m}^3$ | | <input type="text"/> a <input type="text"/> | |
| | Output-conc Formaldehyde, ppm | | <input type="text"/> <input type="text"/> | |
| 2 | Input-Conc Acetaldehyde, $\mu\text{g}/\text{m}^3$ | | <input type="text"/> R/S <input type="text"/> | |
| | Output-Conc Acetaldehyde, ppm | | <input type="text"/> <input type="text"/> | |
| 3 | Input-Conc Acetone, $\mu\text{g}/\text{m}^3$ | | <input type="text"/> R/S <input type="text"/> | |
| | Output-Conc Acetone, ppm | | <input type="text"/> <input type="text"/> | |
| 4 | Input-Conc Isobutyraldehyde, $\mu\text{g}/\text{m}^3$ | | <input type="text"/> R/S <input type="text"/> | |
| | Output-Conc Isobutyraldehyde, ppm | | <input type="text"/> <input type="text"/> | |
| 5 | Input-Conc Methyl ethyl ketone, $\mu\text{g}/\text{m}^3$ | | <input type="text"/> R/S <input type="text"/> | |
| | Output-Conc Methyl ethyl ketone, ppm | | <input type="text"/> <input type="text"/> | |
| 6 | Input-Conc Crotonaldehyde, $\mu\text{g}/\text{m}^3$ | | <input type="text"/> R/S <input type="text"/> | |
| | Output-Conc Crotonaldehyde, ppm | | <input type="text"/> <input type="text"/> | |
| 7 | Input-Conc Hexanaldehyde, $\mu\text{g}/\text{m}^3$ | | <input type="text"/> R/S <input type="text"/> | |
| | Output-Conc Hexanaldehyde, ppm | | <input type="text"/> <input type="text"/> | |
| 8 | Input-Conc Benzaldehyde, $\mu\text{g}/\text{m}^3$ | | <input type="text"/> R/S <input type="text"/> | |
| | Output-Conc Benzaldehyde, ppm | | <input type="text"/> <input type="text"/> | |
| | | | <input type="text"/> RTN <input type="text"/> | |
| | | | <input type="text"/> f <input type="text"/> | |
| | | | <input type="text"/> REG <input type="text"/> | |
| | | | <input type="text"/> CLX <input type="text"/> | |
| | | | <input type="text"/> <input type="text"/> | |
| | | | <input type="text"/> <input type="text"/> | |

Figure 15. Aldehyde and Ketone concentrations in ppm.

HP-65 Program Form

Title _____ Page _____

SWITCH TO W PRGM PRESS ☐ PPGM TO CLEAR MEMORY

| KEY ENTRY | CODE SHOWN | COMMENTS | KEY ENTRY | CODE SHOWN | COMMENTS | REGISTERS |
|-----------|------------|---------------------------|-----------|------------|---------------------------|-----------|
| LBL | 23 | | | | In conc ug/m ³ | R1 |
| A | 11 | In conc ug/m ³ | | | | |
| 1 | | | | | | |
| 2 | | | | | | R2 |
| 4 | | | | | | |
| 9 | | | | | | |
| ÷ | 31 | | | 31 | | |
| R/S | 34 | Out conc ppm | R/S | 34 | Out conc ppm | R3 |
| | | In conc ug/m ³ | RTN | 24 | | |
| 1 | | | | | | R4 |
| 8 | | | | | | |
| 3 | | | | | | |
| 2 | | | | | | R5 |
| ÷ | 31 | | | | | |
| R/S | 34 | Out conc ppm | | | | R6 |
| | | In conc ug/m ³ | | | | |
| 2 | | | | | | |
| 1 | | | | | | R7 |
| 1 | | | | | | |
| ÷ | 31 | | | | | |
| R/S | 34 | Out conc ppm | | | | R8 |
| | | In conc ug/m ³ | | | | |
| 3 | | | | | | |
| 0 | | | | | | R9 |
| 0 | | | | | | |
| 0 | | | | | | |
| ÷ | 31 | | | | | |
| R/S | 34 | Out conc ppm | | | | |
| | | In conc ug/m ³ | | | | |
| 1 | | | | | | |
| 0 | | | | | | |
| 2 | | | | | | |
| 0 | | | | | | |
| ÷ | 31 | | | | | |
| R/S | 34 | Out conc ppm | | | | |
| | | In conc ug/m ³ | | | | |
| 2 | | | | | | |
| 9 | | | | | | |
| 1 | | | | | | |
| 6 | | | | | | |
| ÷ | 31 | | | | | |
| R/S | 34 | Out conc ppm | | | | |
| | | In conc ug/m ³ | | | | |
| 4 | | | | | | |
| 1 | | | | | | |
| 6 | | | | | | |
| 6 | | | | | | |
| ÷ | 31 | | | | | |
| R/S | 34 | Out conc ppm | | | | |
| | | In conc ug/m ³ | | | | |
| 4 | | | | | | |
| 1 | | | | | | |
| 6 | | | | | | |
| 6 | | | | | | |
| ÷ | 31 | | | | | |
| R/S | 34 | Out conc ppm | | | | |

TO RECORD PROGRAM INSERT MAGNETIC CARD WITH SWITCH SET AT W PRGM

Figure 15 Cont'd; Aldenylde and ketone concentrations in ppm.

Sample Calculation

Assume exhaust samples were collected in glass impingers for each portion of a three bag 1975 FTP. Raw data for these tests is presented in Figure 16. Calculations were performed using the HP-65 programs and manual calculations.

Manual calculation for driving cycle FTP-1:

$$\begin{aligned}\mu\text{g}/\text{m}^3 \text{ formaldehyde} &= \frac{\text{Conc}_{\text{Der}} (\text{mg}/\text{ml}) \times \text{Vol}_{\text{Tol}} (\text{ml}) \times \text{mol. wt. carbonyl}}{P_{\text{exp}} (\text{"Hg}) \times V_{\text{exp}} (\text{ft}^3)} \\ &\quad \times \frac{1000 \mu\text{g}/\text{mg} \times T_{\text{exp}} (^{\circ}\text{R}) \times 29.92 \text{" Hg}}{528^{\circ}\text{R}} \\ &\quad \times \frac{35.31 \text{ ft}^3/\text{m}^3}{\text{mol. wt. derivative}} \\ &= \frac{0.186 \text{ mg}/\text{ml} \times 2 \text{ ml} \times 30.03 \text{ g}/\text{mole} \times 1000 \mu\text{g}/\text{mg}}{29.80 \text{" Hg} \times 3.196 \text{ ft}^3 \times 528^{\circ}\text{R}} \\ &\quad \times \frac{535^{\circ}\text{R} \times 29.92 \text{" Hg} \times 35.31 \text{ ft}^3/\text{m}^3}{201.15 \text{ g}/\text{mole}} \\ &= 597.5 \mu\text{g}/\text{m}^3 \\ \text{ppm formaldehyde} &= \mu\text{g}/\text{m}^3 \div \text{density } \mu\text{g}/\text{ml} \\ \text{density } \mu\text{g}/\text{ml} &= \frac{\text{mol. wt. (formaldehyde)} \times 1000}{24.04\ell} \\ \text{mol. wt. formaldehyde} &= 30.03 \text{ g}/\text{mole} \\ \text{density} &= \frac{30.03 \text{ g}/\text{mole} \times 1000}{24.04\ell} = 1249 \mu\text{g}/\text{ml} \\ \text{ppm} &= 597.5 \mu\text{g}/\text{m}^3 \div 1249 \mu\text{g}/\text{ml} = 0.478 \text{ ml}/\text{m}^3 = 0.478 \text{ ppm}\end{aligned}$$

The calculations for acetaldehyde, acetone, isobutyraldehyde, methylethylketone, crotonaldehyde, hexanaldehyde, and benzaldehyde are carried out in the same manner by substituting the appropriate derivative concentrations and molecular weights into the above formulas. These calculations give the following concentrations:

acetaldehyde, 561 $\mu\text{g}/\text{m}^3$ and 0.306 ppm
acetone, 663 $\mu\text{g}/\text{m}^3$ and 0.274 ppm
isobutyraldehyde, 141 $\mu\text{g}/\text{m}^3$ and 0.047 ppm
methylethylketone, 630 $\mu\text{g}/\text{m}^3$ and 0.210 ppm
crotonaldehyde, 541 $\mu\text{g}/\text{m}^3$ and 0.186 ppm
hexanaldehyde, 250 $\mu\text{g}/\text{m}^3$ and 0.060 ppm

SWRI PROJECT NO. _____ TEST NO. _____ TEST DATE: _____ VEHICLE: _____
 FUEL: _____ CVS NO. _____ TUNNEL SIZE: _____ DRIVER: _____ MILES: _____
 SAMPLE COLLECTION BY: _____ CHEMICAL ANALYSIS BY: _____ CALCULATIONS BY: _____
 GENERAL COMMENTS: _____

| Test No. | 1 | 2 | 3 | 4 | 5 | 6 |
|--------------------------------------|-------|-------|-------|-------|-------|-------|
| Driving Cycle | FTP-1 | FTP-2 | FTP-3 | SET-7 | HFET | NYCC |
| Volume, Ft ³ | 3.196 | 1.625 | 2.010 | 3.730 | 8.241 | 1.070 |
| B.P., "Hg | 29.80 | 30.02 | 29.02 | 29.25 | 29.95 | 29.50 |
| Temp. °F | 75 | 80 | 96 | 85 | 83 | 89 |
| Vol. Toluene ml | 2 | 2 | 2 | 2 | 2 | 2 |
| Formaldehyde Der Conc mg/ml | 0.186 | 0.105 | 0.201 | 0.312 | 0.732 | 0.142 |
| Formaldehyde Conc µg/m ³ | 598 | 665 | 1100 | 891 | 921 | 1410 |
| Formaldehyde Conc ppm | 0.479 | 0.532 | 0.881 | 0.713 | 0.737 | 1.130 |
| Acetaldehyde Der Conc mg/ml | 0.127 | 0.092 | 0.157 | 0.282 | 0.612 | 0.102 |
| Acetaldehyde Conc µg/m ³ | 559 | 798 | 1170 | 1100 | 1060 | 1390 |
| Acetaldehyde Conc ppm | 0.305 | 0.436 | 0.639 | 0.600 | 0.579 | 0.759 |
| Acetone Der Conc mg/ml | 0.121 | 0.098 | 0.161 | 0.285 | 0.595 | 0.105 |
| Acetone Conc µg/m ³ | 663 | 1060 | 1500 | 1390 | 1280 | 1780 |
| Acetone Conc ppm | 0.274 | 0.439 | 0.621 | 0.575 | 0.530 | 0.737 |
| I-Bu Aldehyde Der Conc mg/ml | 0.022 | 0.011 | 0.028 | 0.023 | 0.051 | 0.009 |
| I-Bu Aldehyde Conc µg/m ³ | 141 | 139 | 305 | 131 | 128 | 179 |
| I-Bu Aldehyde Conc ppm | 0.047 | 0.046 | 0.102 | 0.044 | 0.043 | 0.060 |
| MeEt Ketone Der Conc mg/ml | 0.098 | 0.084 | 0.097 | 0.198 | 0.252 | 0.075 |
| MeEt Ketone Conc µg/m ³ | 630 | 1060 | 1060 | 1130 | 634 | 1490 |
| MeEt Ketone Conc ppm | 0.210 | 0.353 | 0.353 | 0.377 | 0.211 | 0.497 |
| Cro-Aldehyde Der Conc mg/ml | 0.086 | 0.074 | 0.076 | 0.105 | 0.286 | 0.072 |
| Cro-Aldehyde Conc µg/m ³ | 541 | 917 | 811 | 587 | 705 | 1400 |
| Cro-Aldehyde Conc ppm | 0.186 | 0.314 | 0.278 | 0.201 | 0.242 | 0.480 |
| Hex-Aldehyde Der Conc mg/ml | 0.031 | 0.018 | 0.030 | 0.027 | 0.078 | 0.011 |
| Hex-Aldehyde Conc µg/m ³ | 250 | 286 | 411 | 194 | 246 | 275 |
| Hex-Aldehyde Conc ppm | 0.060 | 0.069 | 0.099 | 0.047 | 0.059 | 0.066 |
| Benzaldehyde Der Conc mg/ml | 0.093 | 0.081 | 0.097 | 0.121 | 0.232 | 0.081 |
| Benzaldehyde Conc µg/m ³ | 775 | 1330 | 1370 | 897 | 757 | 2090 |
| Benzaldehyde Conc ppm | 0.176 | 0.301 | 0.310 | 0.203 | 0.171 | 0.473 |

Figure 16. Aldehyde Collection Sheet.

Benzaldehyde, 775 g/m³ and 0.176 ppm.

Note: The values used in these calculations are picked from a range of temperatures, derivative concentrations, etc. to validate the calculations and may not be representative of expected raw data. The calculations are presented to confirm the manual and HP-65 calculations give the same results. This was confirmed for six sets of calculations.

REFERENCES

This procedure is taken from the procedure: "Oxygenated Compounds in Automobile Exhaust-Gas Chromatograph Procedure" by Fred Stump, ESRL, Environmental Protection Agency, Research Triangle Park, North Carolina.

THIS PROCEDURE IS REPRINTED FROM EPA REPORT EPA 600/2-79-017,
"ANALYTICAL PROCEDURES FOR CHARACTERIZING UNREGULATED POLLUTANT
EMISSIONS FROM MOTOR VEHICLES." (Reference 3)

APPENDIX A-5

MEASUREMENT OF METHANOL

The measurement of methanol in exhaust is accomplished by bubbling the exhaust through glass impingers containing deionized water. The exhaust sample is collected continuously during the test cycle. For analysis, a portion of the aqueous solution is injected into a gas chromatograph equipped with a flame ionization detector (FID). External methanol standards in deionized water are used to quantify the results. Detection limits for this procedure are on the order of 0.06 ppm in dilute exhaust.

SAMPLING SYSTEM

Two glass impingers in series, with each containing 25 mL of deionized water are used to collect exhaust samples for the analysis of methanol. A flow schematic of the sample collection system is shown in Figure 1. The two glass impingers collect 99 percent of the methanol in exhaust. The temperature of the impinger is maintained at 0-5°C by an ice water bath, and the flow rate through the impinger is maintained at 4L/minute by the sample pump. A dry gas meter is used to determine the total flow through the impinger during a given sampling period. The temperature of the gas stream is monitored by a thermocouple immediately prior to the dry gas meter. A drier is included in the system to prevent condensation in the pump, flowmeter, dry gas meter, etc. The flowmeter in the system allows continuous monitoring of the sample flow to insure proper flow rates during the sampling. Several views of the sampling system are shown in Figure 2.

ANALYTICAL PROCEDURE

The analysis of methanol is accomplished by collecting methanol in deionized water and analyzing the sample with a gas chromatograph equipped with an FID. The analysis flow schematic for methanol is shown in Figure 3. A detailed description of the procedure follows.

For the analysis of methanol, dilute exhaust is bubbled through two glass impingers each containing 25 mL of deionized water. Upon completion of each driving cycle, the impinger is removed and the contents are transferred to a 30 mL polypropylene bottle, and capped.

A Perkin-Elmer 3920B gas chromatograph equipped with a flame ionization detector is used to analyze the sample. A 5 µL portion of the sample is injected into the gas chromatograph (GC). The column is a 3' X 1/8" Teflon column containing 120/150 mesh Porapak Q. The carrier gas is helium which flows through the column at a rate of 20 mL/minute. The column temperature is maintained at 100°C. A chromatogram of a standard sample containing 63 ppm methanol is shown in Figure 4. To

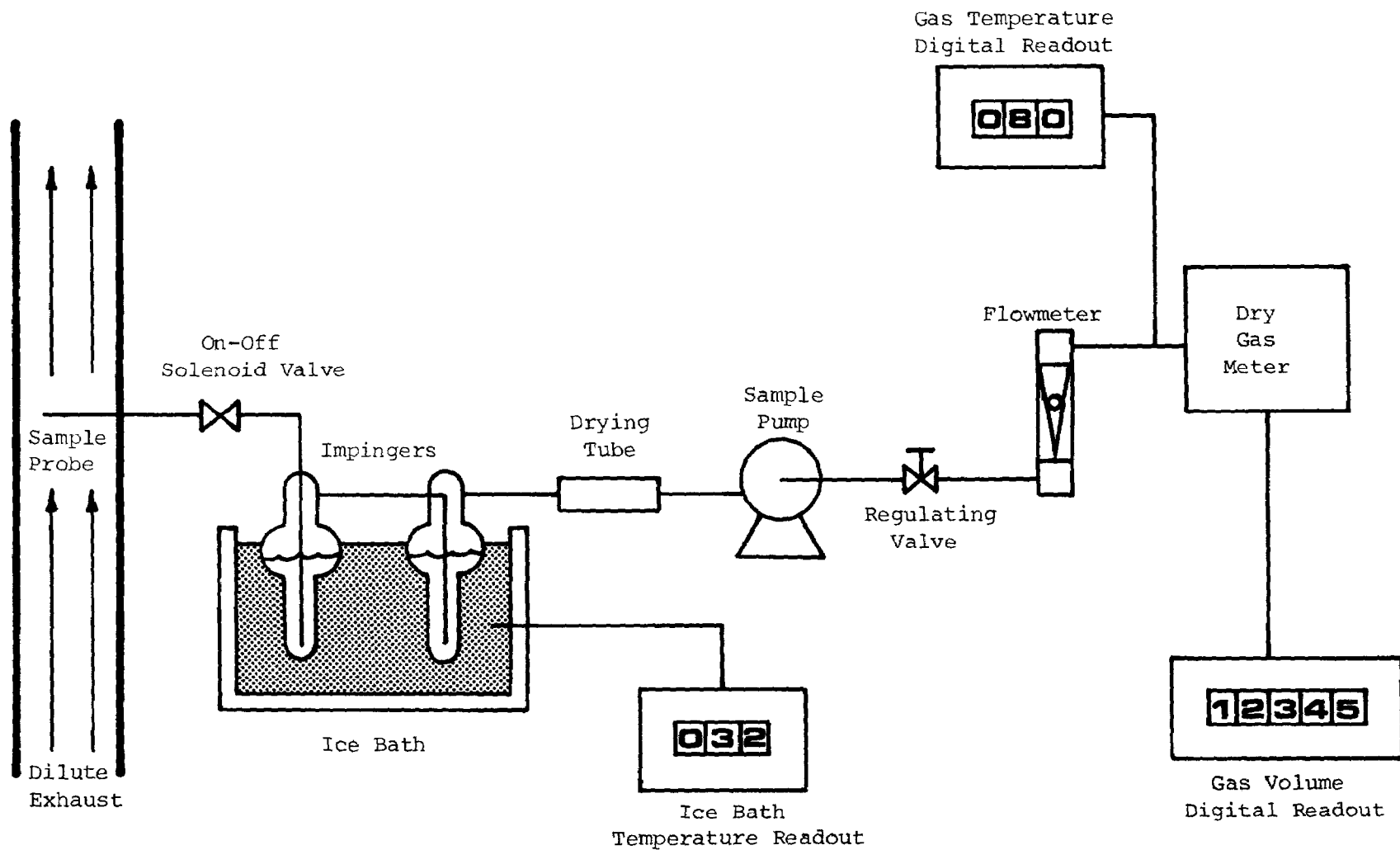
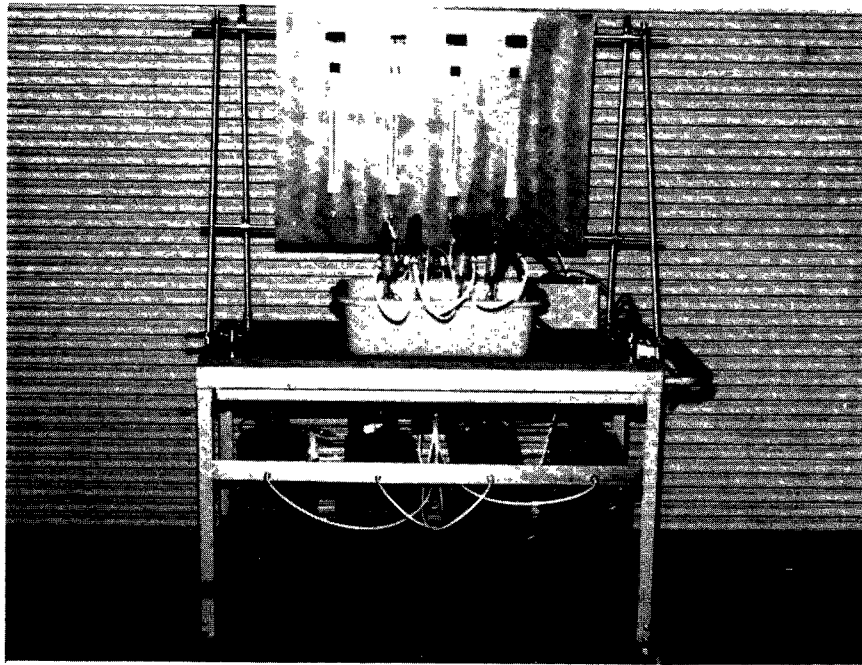
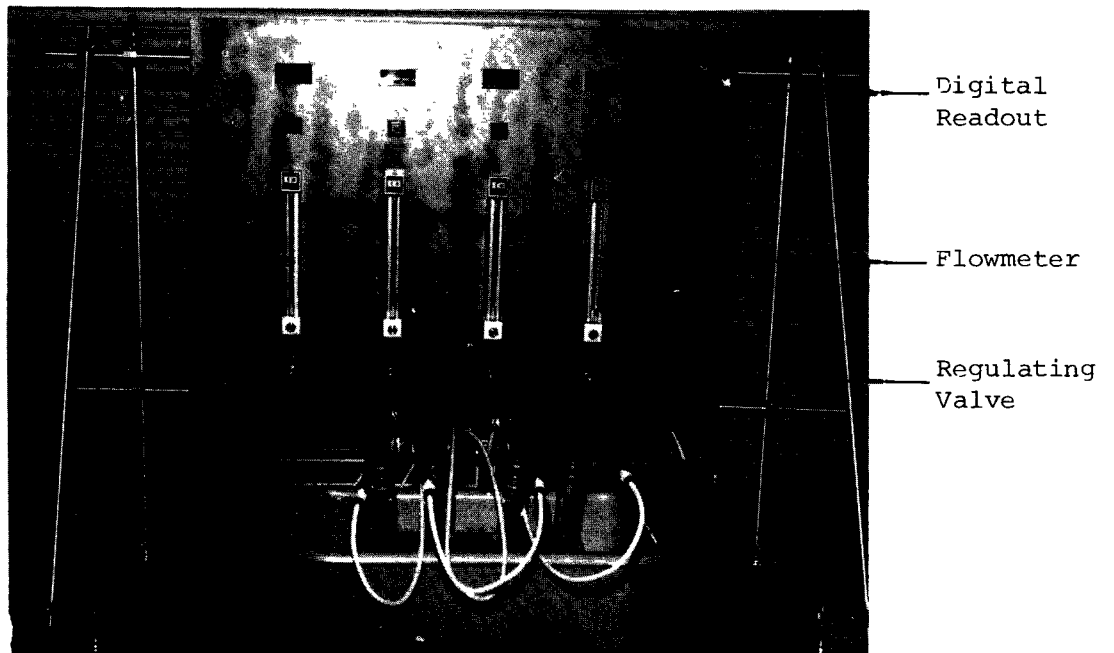


Figure 1. Methanol sample collection flow schematic.

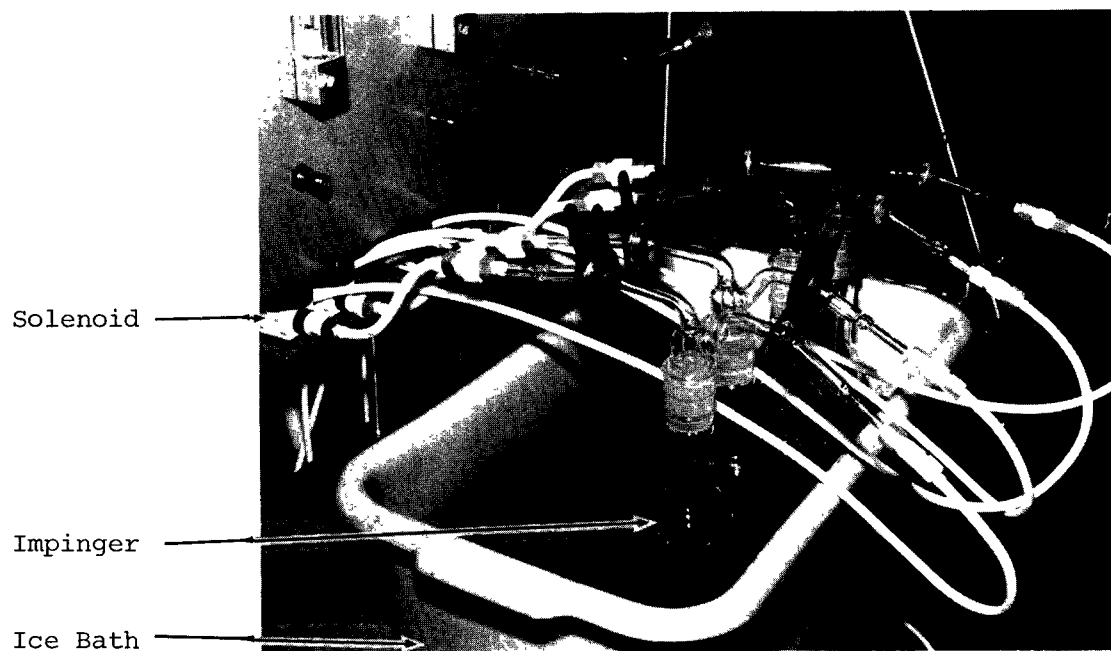


Front View

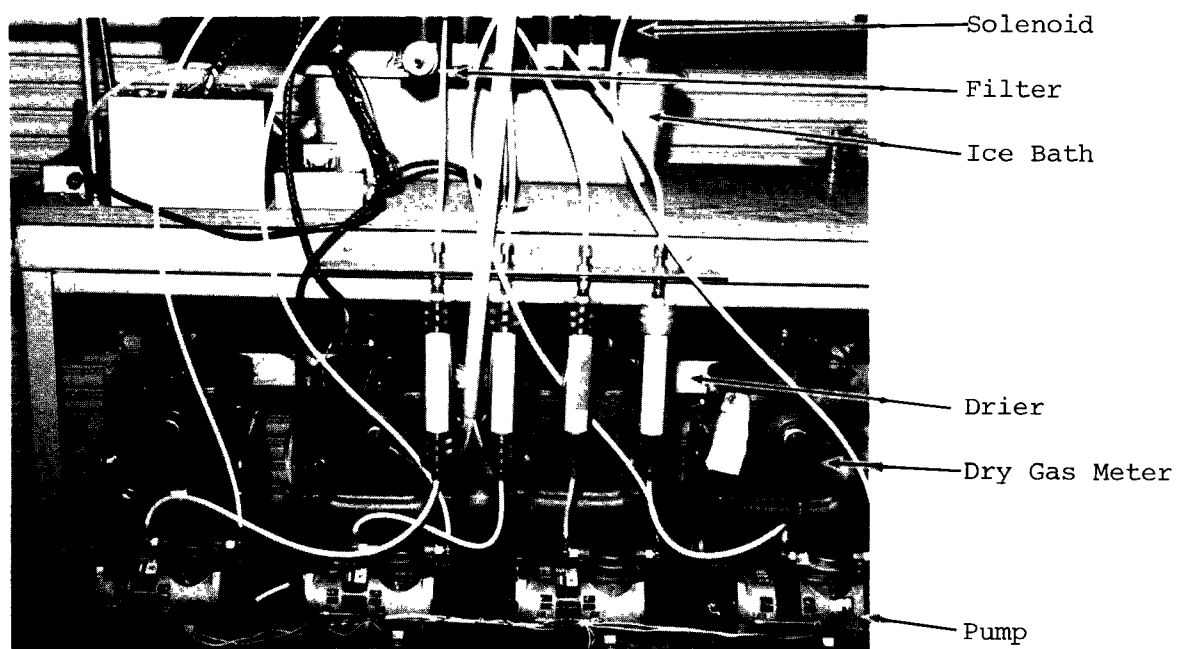


Close-up of Upper Front

Figure 2. Methanol sampling system.



Close-up of Impingers (Side View)



Rear View

Figure 2 (Cont'd). Methanol sampling system.

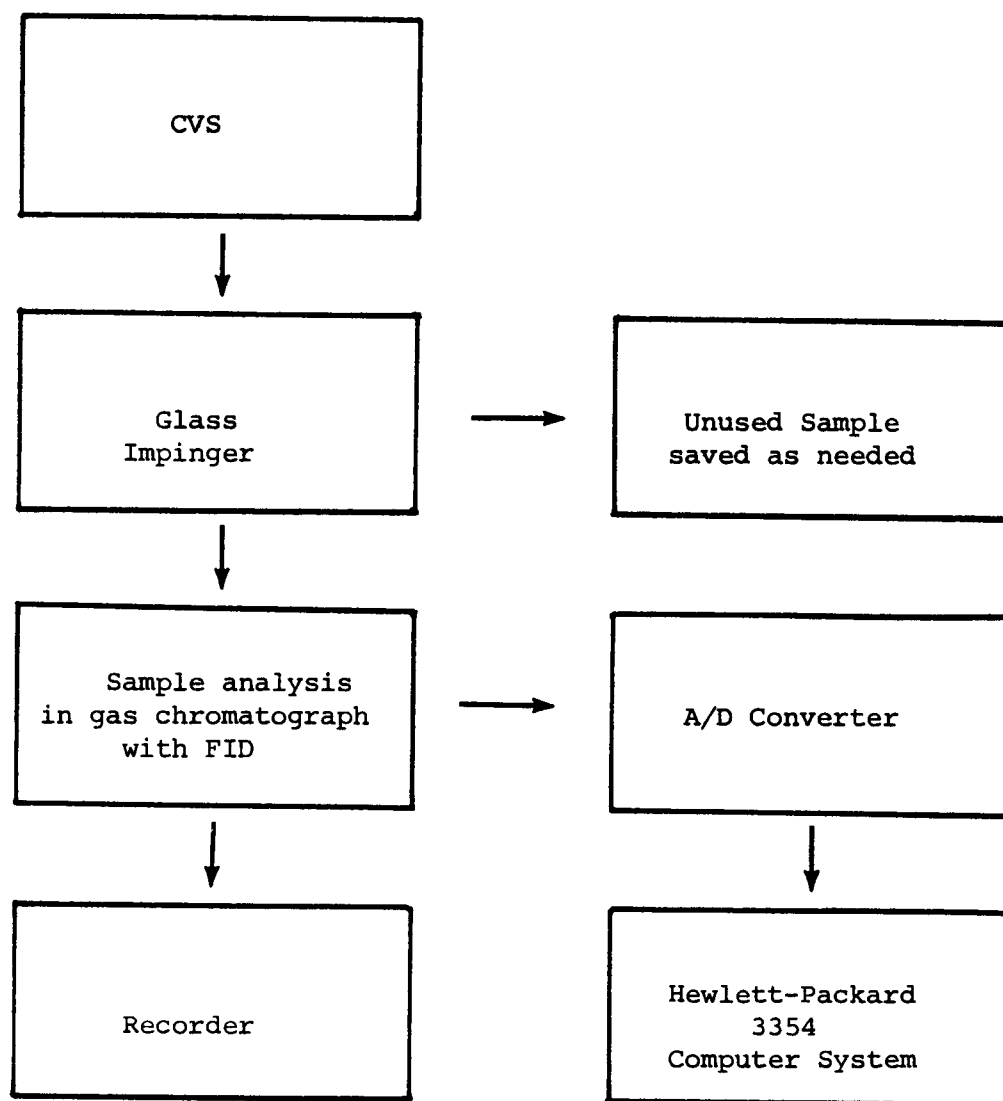


Figure 3. Methanol analysis flow schematic.

Sample Methanol Standard Date 2/11/82
 Instrument PE 3920B Operator K. Miltenberger
 Column 3 ft. 1/8" O.D. I.D. Teflon Type
 Packed with % wt. Liq. Phase
 on 120/150 mesh Porapak Q Support
 Run ISO @ 100 °C using 20 cc/min. Helium Carrier
 @ psig Rotameter Reading
 held @ °C ISO for min., prog to °C
 @ °/min. Held for min., Prog to °C at °/min
 held for min. (other)
 Inlet 150 °C Heated-Glass Lined Type
 Detector 200 °C FID Type (other)
 Hyd 33 psig Rotameter Rdg. cc/min
 Air 57 psig Rotameter Rdg. cc/min
 () psig Rotameter Rdg. cc/min
 Recorder 1 in/min speed 1 mV.F.S. Soltec Type
 Injection 5 ul Indicated 5 ul net 5 ul Actual
 Sampling Device 5 µl syringe

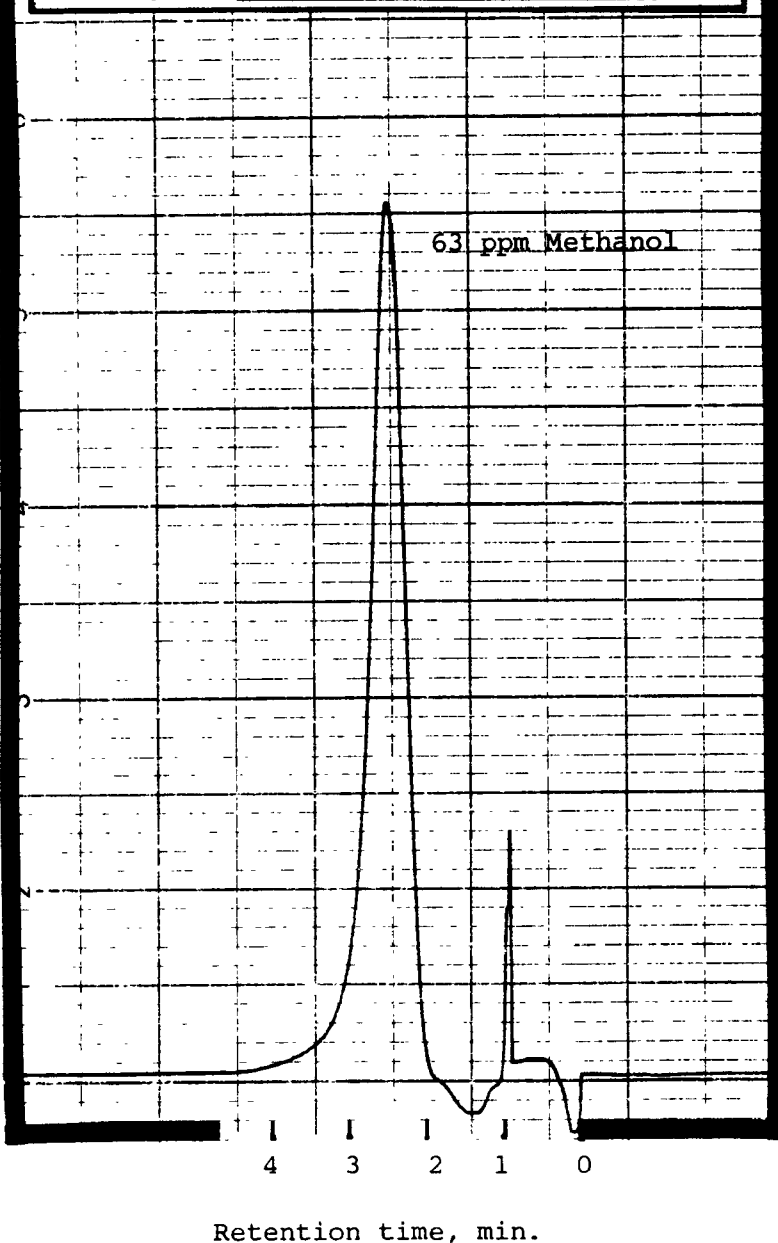


Figure 4. Chromatogram of methanol standard

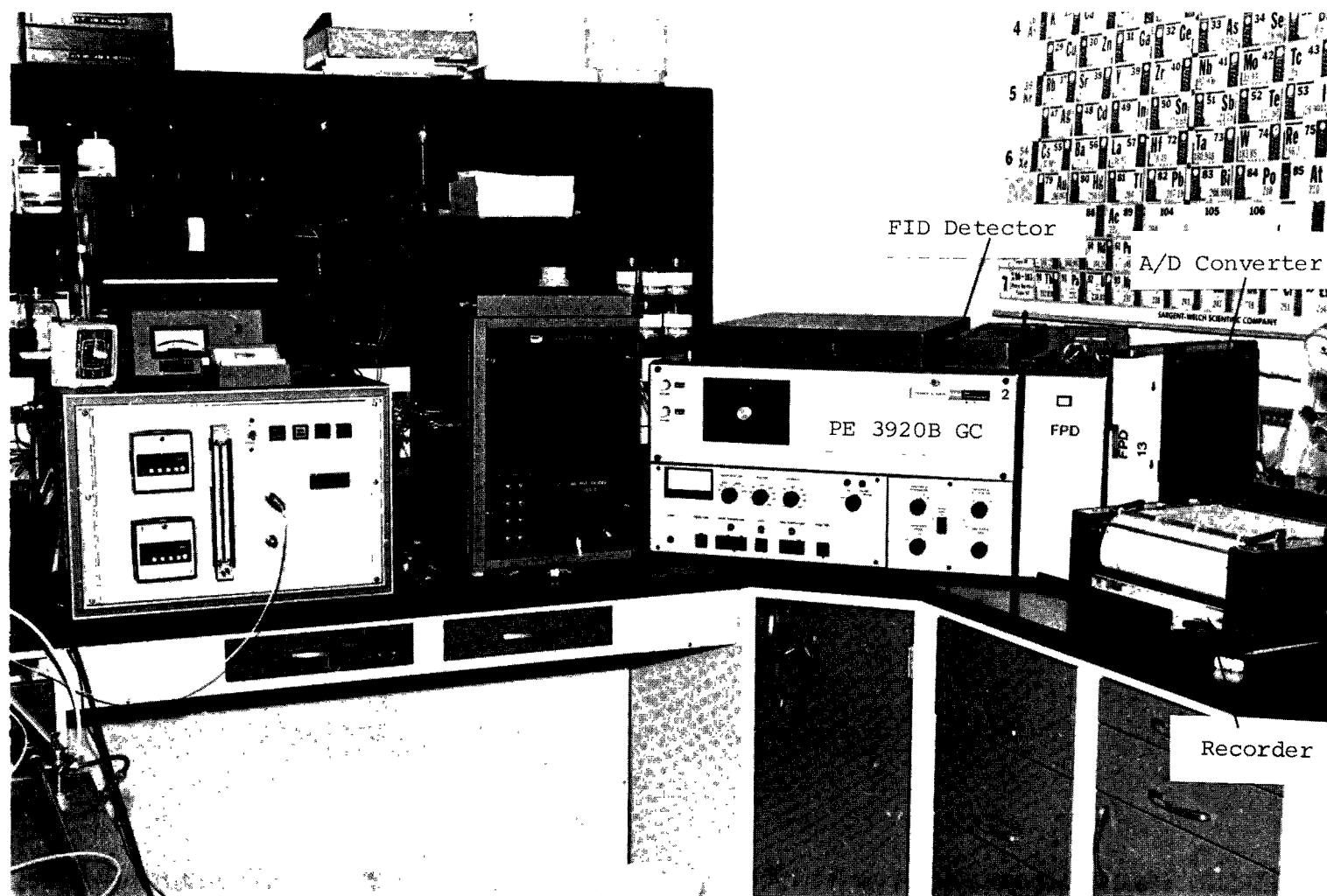


Figure 5. Methanol Analytical system.

quantify the results, the sample peak area is compared to the peak area of a standard solution. Figure 5 shows the analytical system with gas chromatograph, detector, A/D converter, and recorder.

CALCULATIONS

The procedure has been developed to provide the user with the concentration of methanol in exhaust. The results will be expressed in $\mu\text{g}/\text{m}^3$ of exhaust and ppm. The equations for determining the concentrations in $\mu\text{g}/\text{m}^3$ and ppm are derived in the following manner.

The first step is to correct the volume of exhaust sampled to a standard temperature, 68°F , and pressure, 29.92"Hg , by use of the equation

$$\frac{P_{\text{exp}} \times V_{\text{exp}}}{T_{\text{exp}}} = \frac{P_{\text{corr}} \times V_{\text{corr}}}{T_{\text{corr}}}$$

V_{exp} = experimental volume of gas sampled in ft^3

V_{corr} = volume of gas sampled in ft^3 corrected to 68°F and 29.92"Hg

P_{exp} = experimental barometric pressure

P_{corr} = 29.92"Hg

T_{exp} = experimental temperature in $^\circ\text{F} + 460$

T_{corr} = $68^\circ\text{F} + 460 = 528^\circ\text{R}$

Solving for V_{corr} gives:

$$V_{\text{corr}} = \frac{P_{\text{exp}} (\text{"Hg}) \times V_{\text{exp}} (\text{ft}^3) \times 528^\circ\text{R}}{T_{\text{exp}} (^\circ\text{R}) \times 29.92\text{"Hg}}$$

The next step converts the volume from cubic feet to cubic meters by use of the conversion factor: 1 cubic meter is equal to 35.31 cubic feet.

$$V_{\text{corr}} (\text{m}^3) = \frac{P_{\text{exp}} (\text{"Hg}) \times V_{\text{exp}} (\text{ft}^3) \times 528^\circ\text{R}}{T_{\text{exp}} \times 29.92\text{"Hg} \times 35.31 \text{ ft}^3/\text{m}^3}$$

(Equation 1)

The next step is to find the concentration of methanol in $\mu\text{g}/\text{mL}$. Since the gas chromatograph FID has a linear response in the concentration of concern, then the following equation holds.

$$\frac{C_{\text{sam}} (\mu\text{g}/\text{mL})}{A_{\text{sam}}} = \frac{C_{\text{std}} (\mu\text{g}/\text{mL})}{A_{\text{std}}}$$

C_{sam} = concentration of the sample in $\mu\text{g/ml}$
 A_{sam} = GC peak area of sample in relative units
 C_{std} = concentration of the standard in $\mu\text{g/ml}$
 A_{std} = GC peak area of standard in relative units

Solving for C_{sam} gives:

$$C_{sam}(\mu\text{g/ml}) = \frac{C_{std}(\mu\text{g/ml}) \times A_{sam}}{A_{std}}$$

The $C_{sam}(\mu\text{g/ml})$ in solution is corrected for any necessary dilution by multiplying by the dilution factor, D.F.

$$C_{sam}(\mu\text{g/ml}) = \frac{C_{std}(\mu\text{g/ml}) \times A_{sam} \times \text{D.F.}}{A_{std}}$$

To obtain the total amount in μg of methanol in the aqueous absorbing solution, the absorbing reagent volume is multiplied by the concentration to give:

$$\mu\text{g sample} = C_{sam}(\mu\text{g/ml}) \times \text{Abs. Vol. (ml)}$$

$$= \frac{C_{std}(\mu\text{g/ml}) \times A_{sam} \times \text{D.F.} \times \text{Abs. Vol. (ml)}}{A_{std}}$$

(Equation 2)

To obtain $\mu\text{g sample/m}^3$, Equation 2 is divided by Equation 1 to give:

$$\mu\text{g samp/m}^3 = \frac{C_{std}(\mu\text{g/ml}) \times A_{sam} \times \text{D.F.} \times \text{Abs. Vol. (ml)}}{A_{std} \times P_{exp}(\text{"Hg}) \times 528^\circ}$$

$$\times \frac{T_{exp} \times 29.92\text{"Hg} \times 35.31(\text{ft}^3/\text{m}^3)}{V_{exp}(\text{ft}^3)}$$

(Equation 3)

To find the concentration of methanol in ppm, the density of the methanol is needed. At 29.92"Hg and 32°F, one mole of gas occupies 22.4 liters. This volume is corrected to 68°F from the equation

$$\frac{V}{T} = \frac{V_1}{T_1}$$

$$V_1 = 22.4\ell$$

$$T_1 = 32^\circ\text{F} + 460 = 492^\circ\text{R}$$

$$V = \text{volume at } 68^\circ\text{F}$$

$$T = 68^\circ\text{F} + 460 = 528^\circ\text{R}$$

Solving for V gives:

$$V = \frac{V_1 \times T}{T_1} = \frac{22.4 \times 528}{492} = 24.04\ell$$

Since one mole of gas occupies 22.04ℓ at 68°F, the density can be found in g/ℓ by dividing the molecular weight in g/mole by 22.04 ℓ/mole

$$\text{den (g/ℓ)} = \frac{\text{mol. wt. g/mole}}{24.04 \ell/\text{mole}}$$

The density in μg/ml can be found by converting g to μg and ℓ to ml as follows:

$$\text{den } \mu\text{g/ml} = \frac{\text{mol. wt. g/mole}}{24.04 \ell/\text{mole}} \times \frac{1 \times 10^6 \mu\text{g/g}}{1 \times 10^3 \text{ml}/\ell} = \frac{\text{mol. wt.} \times 1000}{24.04}$$

(Equation 4)

To obtain the concentration of methanol in ppm, the concentration in μg/m³ is divided by the density in μg/ml

$$\text{ppm} = \mu\text{g/m}^3 \div \mu\text{g/ml} = \frac{\text{ml}}{\text{m}^3}$$

Using Equations 3 and 4 gives the ppm concentration in the form of the raw data.

$$\text{ppm} = \frac{24.04(l) \times C_{\text{std}} (\mu\text{g/ml}) \times A_{\text{sam}} \times \text{D.F.} \times \text{Abs. Vol. (ml)}}{\text{Mol. Wt. (g/mole)} \times 1000 \times A_{\text{std}} \times P_{\text{exp}} ("Hg)} \\ \times \frac{T_{\text{exp}} (^{\circ}\text{R}) \times 29.92"Hg \times 35.31 \text{ ft}^3/\text{m}^3}{528^{\circ}\text{R} \times V_{\text{exp}} (\text{ft}^3)}$$

(Equation 5)

At this point, the concentration can be expressed in $\mu\text{g}/\text{m}^3$ (Equation 3) and ppm (Equation 5) at 68°F and $29.92"Hg$ from the raw data.

Hewlett-Packard Calculations

In order to insure maximum turnaround in a minimum time period, a Hewlett-Packard 67 program was developed to calculate the methanol concentrations in $\mu\text{g}/\text{m}^3$ and ppm from the raw data. This program is presented in Figure 6.

Sample Calculation

Assume exhaust samples were collected in glass impingers for each portion of a three bag 1975 FTP. Raw data for these tests are presented in Figure 7. Calculations were performed using the HP 67 program and manual calculations.

Manual calculations for driving cycle FTP-1

For Bubbler #1

$$\mu\text{g}/\text{m}^3\text{CH}_3\text{OH} = \frac{C_{\text{std}} (\mu\text{g/ml}) \times A_{\text{sam}} \times \text{D.F.} \times \text{Abs. Vol. (ml)}}{A_{\text{std}} \times P_{\text{exp}} ("Hg)} \\ \times \frac{T_{\text{exp}} \times 29.92"Hg \times 35.31 \text{ ft}^3/\text{m}^3}{528^{\circ}\text{R} \times V_{\text{exp}} (\text{ft}^3)} \\ = \frac{(7.9 \mu\text{g/ml}) \times 1000 \times 1 \times 25}{1500 \times 29.80"Hg} \\ \times \frac{(460 + 75) \times 29.92"Hg \times 35.31 \text{ ft}^3/\text{m}^3}{528^{\circ}\text{R} \times 3.196 \text{ ft}^3} \\ = 1480 \mu\text{g}/\text{m}^3$$

User Instructions

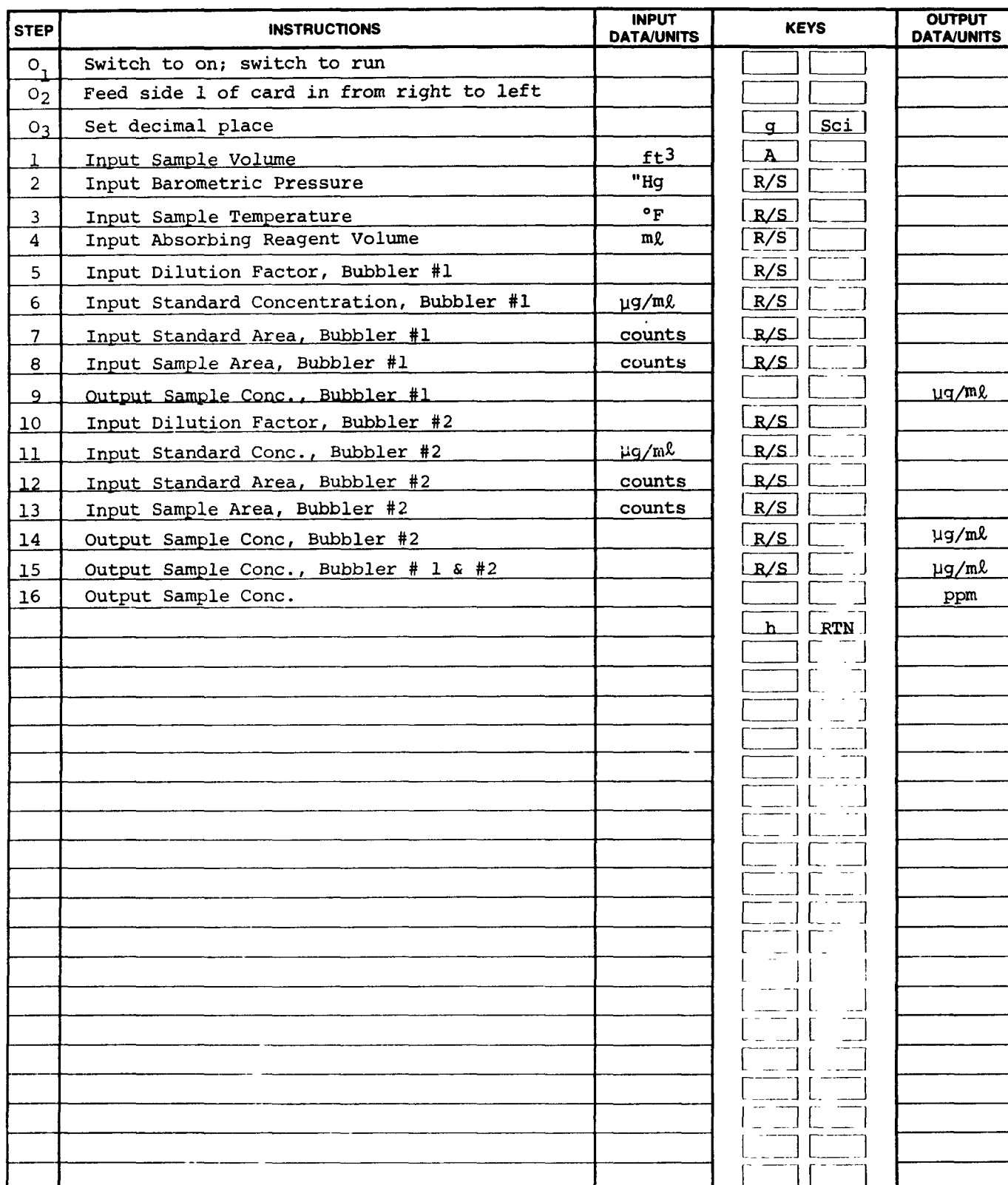


Figure 6 (Cont'd). HP-67 Program Form

[illegible]

HP-67 Program Form

[illegible]

METHANOL

SWRI PROJECT NO. _____ TEST NO. _____ TEST DATE: _____ VEHICLE: _____
 FUEL _____ CVS NO. _____ TUNNEL SIZE: _____ DRIVER: _____ MILES: _____
 SAMPLE COLLECTION BY: _____ CHEMICAL ANALYSIS BY: _____ CALCULATIONS BY: _____
 GENERAL COMMENTS _____

| Test No. | 1 | 2 | 3 | 4 | 5 | 6 |
|--|-------|-------|--------|-------|-------|-------|
| Driving Cycle | FTP-1 | FTP-1 | FTP-3 | SET-7 | HFET | NYCC |
| Volume, Ft ³ | 3.196 | 1.625 | 2.010 | 3.730 | 8.241 | 1.070 |
| B.P., "Hg | 29.80 | 30.02 | 29.02 | 29.25 | 29.95 | 29.50 |
| Temp. °F | 75 | 80 | 96 | 85 | 83 | 89 |
| Absor. Rea. Vol., ml | 25 | 25 | 50 | 50 | 75 | 75 |
| Dilution Factor, Bubbler #1 | 1 | 5 | 10 | 2 | 1 | 1 |
| Std. Conc $\mu\text{gCH}_3\text{OH/ml}$ Bub. #1 | 7.9 | 79 | 790 | 7.9 | 7.9 | 0.79 |
| Std. Area - Bubbler #1 | 1500 | 5000 | 10000 | 1000 | 5000 | 1000 |
| Sample Area - Bubbler #1 | 1000 | 1000 | 1000 | 3000 | 15000 | 3000 |
| Sample Conc $\mu\text{gCH}_3\text{OH/m}^3$, Bub#1 | 1480 | 43700 | 753000 | 23700 | 7820 | 6180 |
| Dilution Factor, Bubbler #2 | 1 | 2 | 5 | 1 | 1 | 1 |
| Std. Conc $\mu\text{gCH}_3\text{OH/ml}$ Bub. #2 | 0.79 | 7.9 | 7.9 | 0.79 | 0.79 | 0.79 |
| Std. Area - Bubbler #2 | 2000 | 1500 | 5000 | 1000 | 6000 | 5000 |
| Sample Area - Bubbler #2 | 500 | 300 | 6000 | 100 | 1000 | 15000 |
| Sample Conc $\mu\text{gCH}_3\text{OH/m}^3$, Bub#2 | 55.5 | 1750 | 45200 | 39.5 | 43.5 | 6180 |
| Total Conc. $\mu\text{gCH}_3\text{OH/m}^3$ | 1530 | 45500 | 798000 | 23700 | 7860 | 12400 |
| PPM Methanol | 1.15 | 34.1 | 599 | 17.8 | 5.90 | 9.28 |

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Figure 7. Methanol sample collection sheet

The concentration in Bubbler #2 is calculated in the same manner using the appropriate dilution factor, standard concentration, standard area, and sample area:

For Bubbler #2:

$$\begin{aligned}\mu\text{g}/\text{m}^3 &= \frac{0.79 \mu\text{g}/\text{mL} \times 500 \times 1 \times 25}{2000 \times 29.80} \\ &\times \frac{(460 + 75) \times 29.92^\circ\text{Hg} \times 35.31 \text{ ft}^3/\text{m}^3}{528^\circ\text{R} \times 3.196 \text{ ft}^3} \\ &= 55.5 \mu\text{g}/\text{m}^3\end{aligned}$$

The concentrations from the two bubblers can be added for a total concentration:

$$\begin{aligned}\text{Total } \mu\text{g methanol}/\text{m}^3 &= \text{conc. (Bubbler \#1)} + \text{conc. (Bubbler \#2)} \\ &= 1480 \mu\text{g}/\text{m}^3 + 55.5 \mu\text{g}/\text{m}^3 \\ &= 1535 \mu\text{g}/\text{m}^3\end{aligned}$$

$$\text{PPM CH}_3\text{OH} = \mu\text{g}/\text{m}^3 \div \text{density } \mu\text{g}/\text{mL}$$

$$\text{density } \mu\text{g}/\text{mL} = \frac{\text{Mol. Wt. (CH}_3\text{OH)} \times 1000}{24.04\text{l}}$$

$$\text{Mol. Wt. CH}_3\text{OH} = 32.04 \text{ g/mole}$$

$$\text{density} = \frac{32.04 \times 1000}{24.04\text{l}} = 1333 \mu\text{g}/\text{mL}$$

$$\text{ppm} = 1535 \div 1333 \mu\text{g}/\text{mL} = 1.15 \text{ mL}/\text{m}^3 = 1.15 \text{ ppm}$$

Note: The values used in these calculations are picked from a range of temperatures, standards, dilution factors, etc., to validate the calculations and may not be representative of expected raw data. These calculations are presented to confirm that the manual and HP-67 calculations give the same results. This was confirmed on six sets of calculations.

LIST OF EQUIPMENT AND REAGENTS

The equipment and reagents for the analysis of the methanol is divided into two groups. The first involves the sample acquisition and the second the instrumental analysis of the sample once it has been obtained. Manufacturer, stock number and any pertinent descriptive information are listed. The preparation of standards is also discussed.

Sampling

1. Glass impingers, Ace Glass Products, Catalog #7530-11, plain tapered tip stoppers with 18/7 arm joints and 29/42 bottle joints.
2. Flowmeter, Brooks Instrument Division, Model 1555, tube size R-2-15-C, graduated 0-15, sapphire float, 0-5 l/min range.
3. Sample pump, Thomas Model 106 CA18, capable of free flow capacity of 4 l/min.
4. Dry gas meter, American Singer Corporation, Type AL-120, 60 CFH capacity.
5. Regulating valve, Nupro 4MG, stainless steel.
6. Teflon tubing, United States Plastic Corporation, 1/4" OD x 1/8" ID and 5/16" OD x 1/8" ID.
7. Teflon solenoid valve, the Fluorocarbon Company, Model DV2-144NCA1.
8. Drying tube, Analabs Inc., Catalog #HGC-146, 6" long, 1/4" brass fittings.
9. Miscellaneous Teflon nuts, ferrules, unions, tees, clamps, connectors, etc.
10. Digital readout for dry gas meter.
11. Miscellaneous electrical switches, lights, wirings, etc.
12. Six channel digital thermometer, Analog Devices, Model #2036/J/1.
13. Iron/Constantan type J single thermocouple with 1/4" OD stainless steel metal sheath, Thermo Sensors Corporation.
14. 30 ml polypropylene sample storage bottles, Nalgene Labware, Catalog #2006-0001.
15. Deionized or distilled water.
16. Class A, 10 ml volumetric pipet.
17. Class A, 1000 ml volumetric flask.

Instrumental Analysis

1. 5 μ l syringe, Hamilton Co., Reno, Nevada.
2. Perkin-Elmer Model 3920 B gas chromatograph equipped with flame ionization detector.
3. Soltec Model B-281 1 mv recorder.
4. Hewlett-Packard Model 3354 gas chromatograph computer system with remote teletype printout.

Preparation of Primary Standards

The primary standard for the methanol analysis is prepared by diluting a known volume of methanol with deionized (or distilled) water. Standards less than 500 ppm are prepared by diluting higher concentration standards with deionized water.

THIS PROCEDURE IS REPRINTED FROM EPA REPORT EPA 460/3-82-004,
"CHARACTERIZATION OF EXHAUST EMISSIONS FROM METHANOL AND
GASOLINE FUELED AUTOMOBILES." (Reference 5)

APPENDIX A-6

MEASUREMENT OF ETHANOL

The measurement of ethyl alcohol in exhaust and evaporative emissions was accomplished by direct bag analysis using a gas chromatograph equipped with a flame ionization detector. A limited number of qualification and validation experiments were conducted to insure the accuracy and reliability of the procedure. Contacts were made with key individuals at EPA, Ann Arbor and Research Triangle Park to coordinate the approach to this analysis with other participants in the evaporative emissions program.

Analytical System

The analysis for ethyl alcohol in exhaust and evaporative emissions is conducted using a gas chromatograph equipped with a flame ionization detector. The system employs two pneumatically operated electrically controlled Seiscor valves, one in a gas sampling valve configuration and the other in the back-flush configuration. Figure A-1 illustrates the flow schematic of the analytical system.

The system was designed for bag analysis and used normal CVS bag samples for the exhaust emissions. In the case of the SHED emissions, bag samples were collected during the first and last minute of the soak. These bags were then transported to the gas chromatograph for subsequent analysis.

The analytical column selected for this separation was a 10' x 1/8" SS with 15% TCEP on 60/80 Chromosorb P. The separation of ethyl alcohol from benzene and toluene was satisfactory and no known interferences were encountered. The gas chromatograph oven was maintained at 50°C and the system was backflushed after the toluene peak. The total analysis time was 45-60 minutes. A typical separation of ethyl alcohol and the two closest eluting compounds is presented in Figure A-2. The specific GC operating parameters for this separation is presented in Table A-1.

Control System

The control of the two Seiscor valves is accomplished by ATC timers and ASCO electric solenoid valves. The electrical schematic for the control of the Seiscor valves using these timers and electric solenoid valves is shown in Figure A-3. The flow schematic for the vacuum and pressure lines to the Seiscor valves are presented in Figure A-4.

Equipment

This analysis is performed using a gas chromatograph equipped with a flame ionization detector. The gas chromatograph, recorder and data acquisition system are major components in the detection system. A control console was fabricated to house the mechanical hardware items that are necessary for the proper operation of the ethyl alcohol analysis system. The major items that are included in each of these systems is listed as follows:

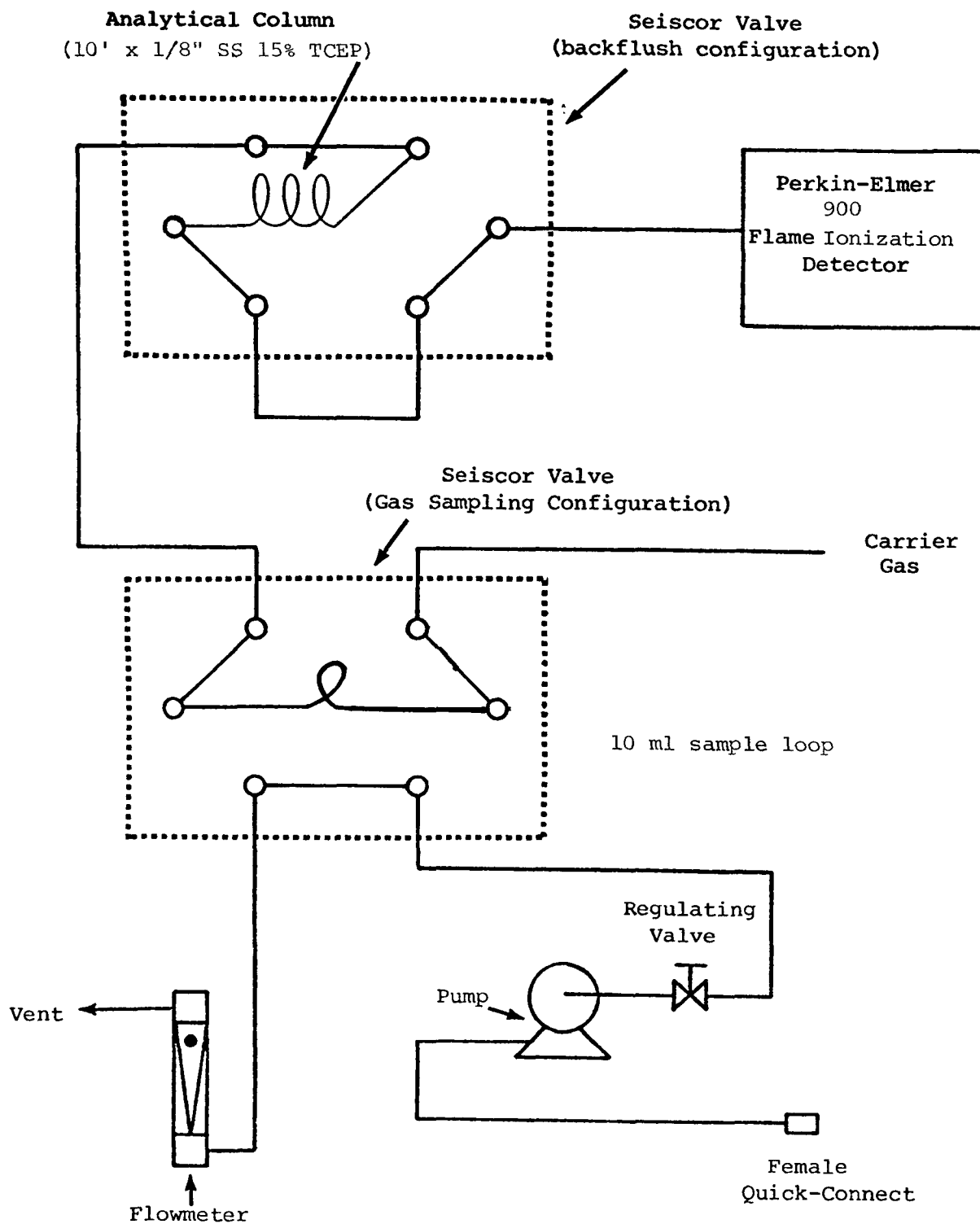


Figure 1. Ethyl Alcohol Analytical Flow Schematic

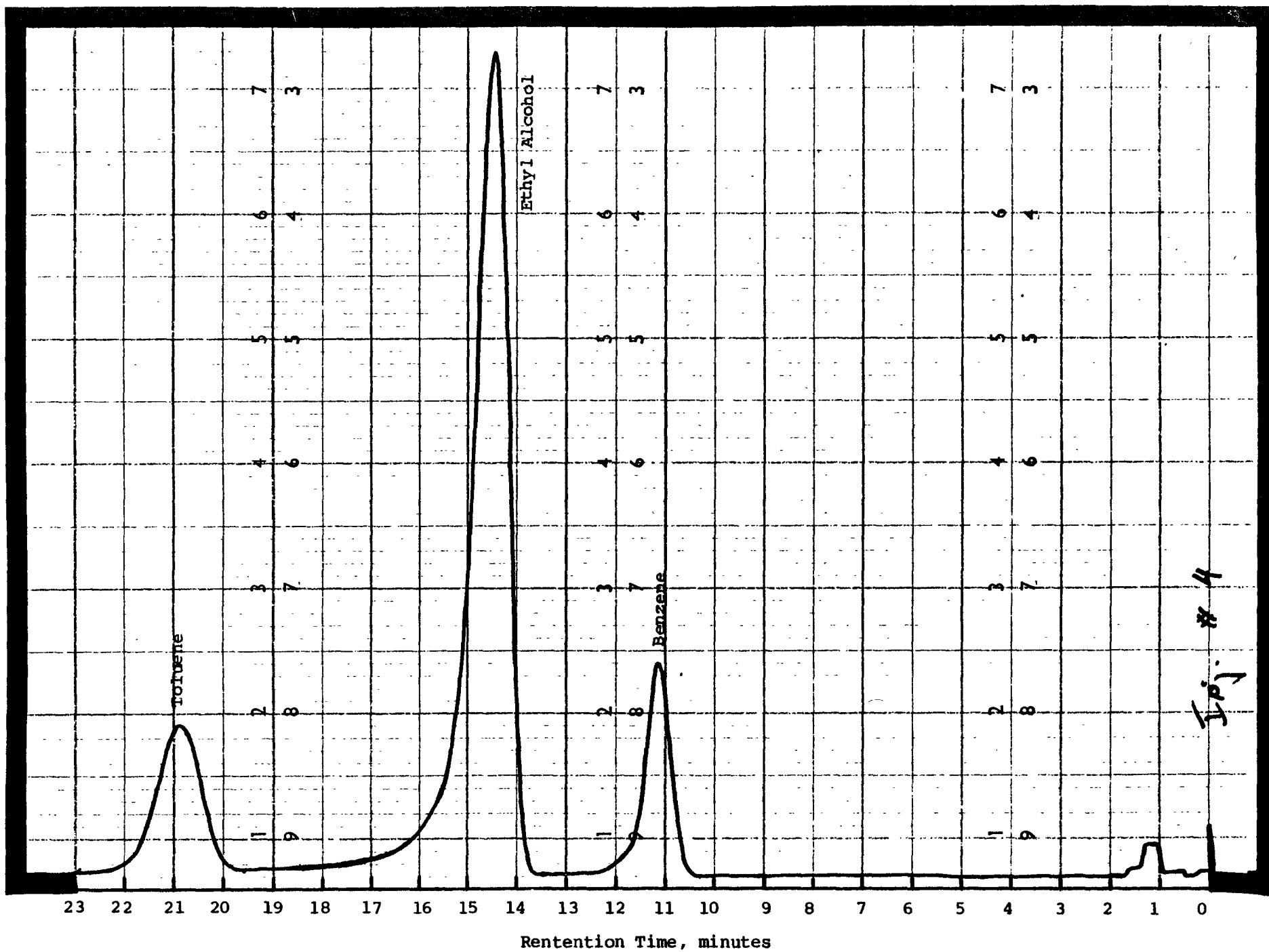


Figure 2. Typical Separation of Ethyl Alcohol from Benzene and Toluene

TABLE 1. ETHYL ALCOHOL ANALYSIS GAS CHROMATOGRAPH OPERATING PROCEDURES

| | |
|------------------------|--|
| Gas Chromatpgraph: | Perkin-Elmer Model 900 |
| Detector Type: | Flame Ionization Detector |
| Gas Sampling Valve: | Seiscor Model VIII (GSV Configuration) |
| Backflush Valve | Seiscor Model VIII (BF Configuration) |
| Sample Loop Size: | 10 ml |
| Column Temperature: | 50°C |
| Detector Temperature: | 50°C |
| Injection Temperature: | 27°C |
| Carrier Gas: | Helium |
| Carrier Gas Flow: | 30 ml/min |
| Hydrogen Gas Flow: | 40 ml/min |
| Air Flow Rate: | 400 ml/min |
| Column Dimensions: | 10' x 1/8" SS |
| Column Material: | 15% TCEP on 60/80 Chromosorb P-AW |

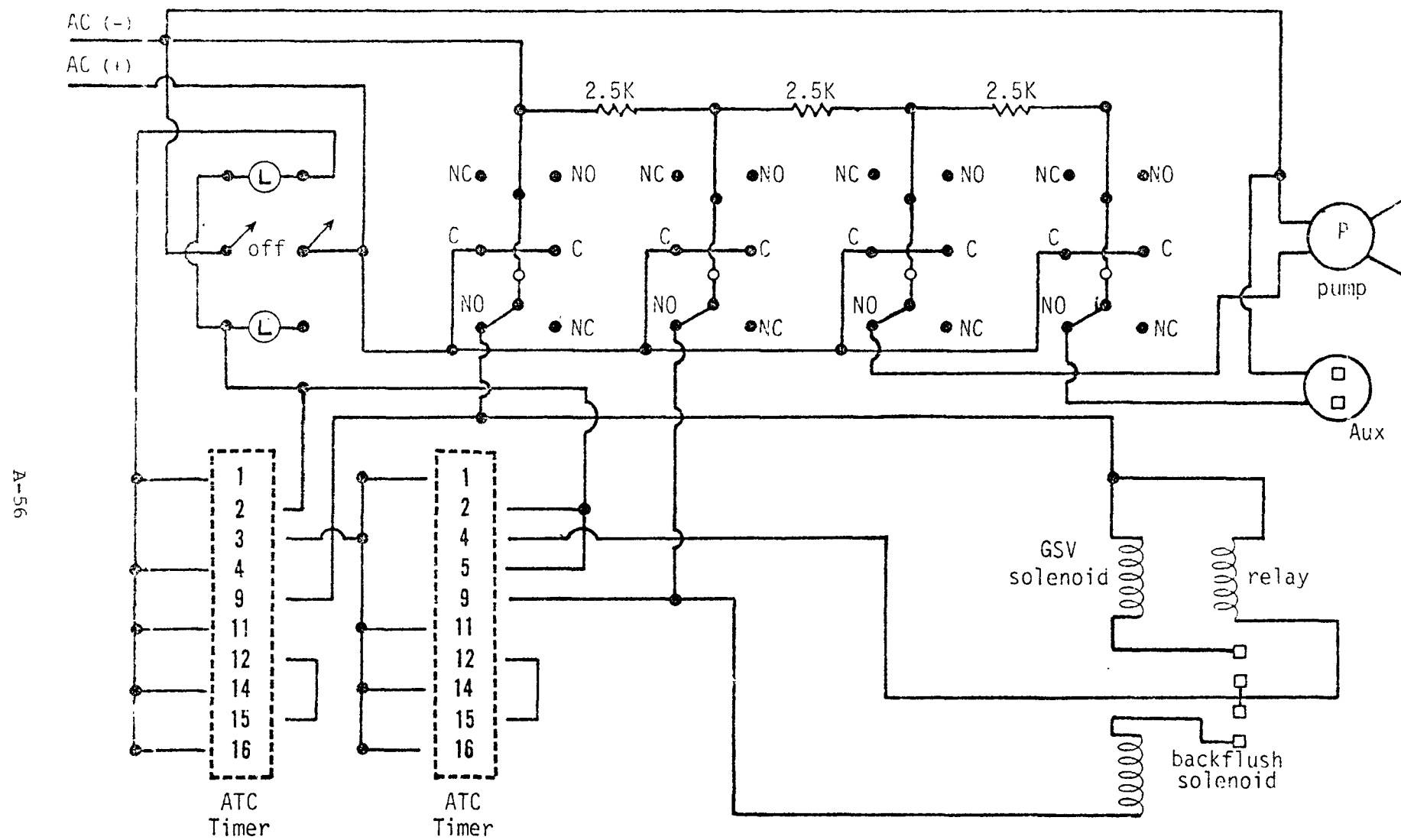


Figure 3. Electrical Schematic for Nitrous Oxide Analysis System

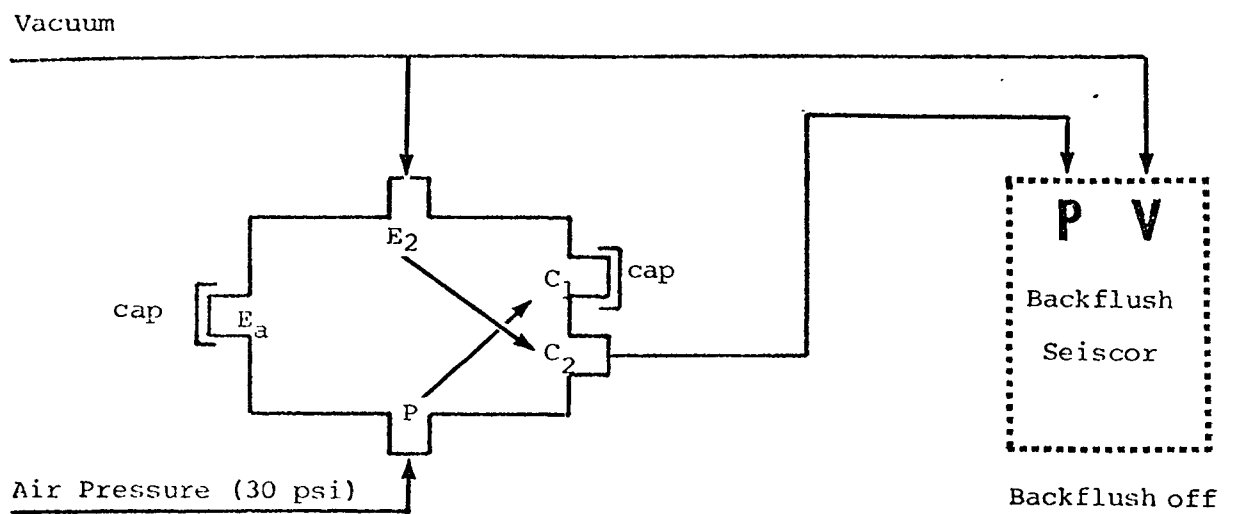
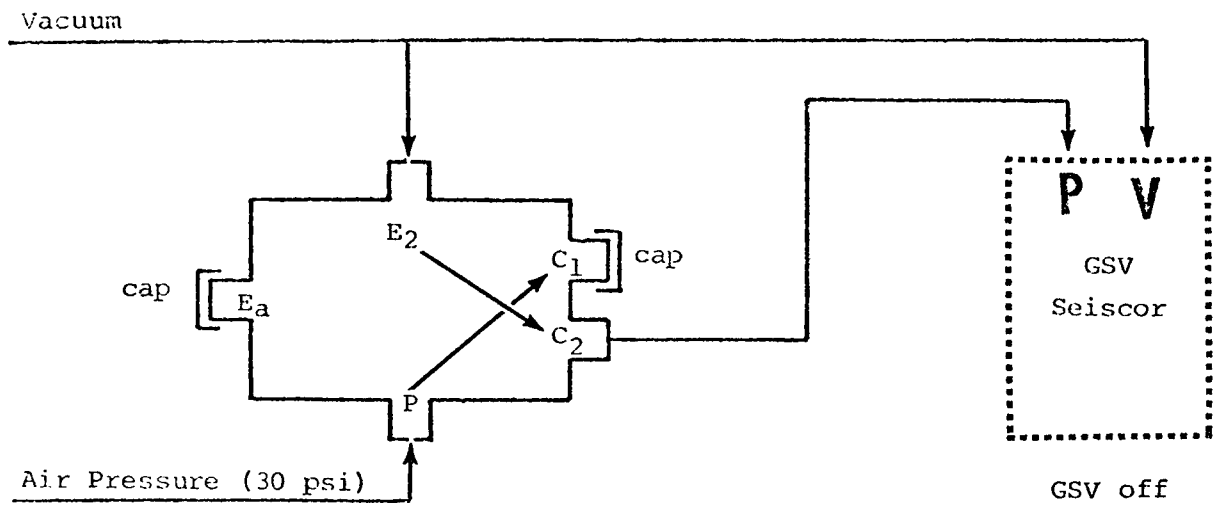


Figure 4. Flow Schematic in Electric Solenoid Valves
(Both valves de-energized)

Gas Chromatograph

1. Perkin-Elmer Model 900 chromatograph
2. Soltec Model B-281 recorder
3. Hewlett-Packard Model 3354C GC computer system
4. Hewlett-Packard Model 1865A A/D Converter
5. Analytical column, 10' x 1/8" SS, 15% TCEP on 60/80 Chromosorb P-AW

Control Console System

1. Seiscor valve - gas sampling configuration
2. Seiscor valve - backflush configuration
3. ATC timers, Model 3254A346A10PX (2 ea)
4. ACSO solenoid valve, Model 834501 (2 ea)
5. Brook flowmeter, R-2-15-A w/SS float, 0-150 scale
6. Metal Bellows MB-155 pump
7. Female quick-connect, stainless steel
8. Nupro Model 2M stainless steel regulating valve
9. Stainless steel tubing (0.01"ID) for capillary restrictor
10. Miscellaneous stainless steel, copper and Teflon tubing (1/8" and 1/16")
11. Miscellaneous stainless steel and brass unions, tees, etc.
12. Bud Classic II control console cabinet, 14" x 19" panel
13. Miscellaneous electrical on-off switches

SHED Bag Sample Acquisition System

1. Sample pump, Thomas Model 917 CA TFE
2. Brooks flowmeter, Model 1555, R-6-15-A, sapphire float, 0-11 lpm
3. Regulating valve, Mupro 4MG, SS
4. Teflon tubing, 1/4" OD x 1/8" ID
5. SS quick connects, male and female
6. Tedlar bag ~ one cubic foot capacity
7. Miscellaneous nuts, females and assorted unions, tees, etc.

Sample Calculations

It was initially planned to use ethyl alcohol blends in normal calibration cylinders. During the validation of the calibration technique, it became apparent that high pressure ethyl alcohol blends were not stable and other calibration techniques were investigated. Benzene was selected to be used as the external standard and the relative response to ethyl alcohol was established. Experiments determined that 1 ppm C (benzene) was equivalent to 0.05 mg ethanol per cubic foot. Using this relationship, the following equations were used to determine grams of ethanol per test.

1. General Calculation Equation

$$\text{ppm C Ethanol} = \frac{\text{GC area of Ethanol} \times \text{ppm C Standard (Benzene)}}{\text{GV area standard (benzene)}}$$

2. Ethanol Evaporative Losses

Ethanol g/test - ppm C (0.050 mg/ppm C ft³) (SHED vol. ft³) (1 g/10³mg)

If a test used a 68.3 ppm C benzene standard that gave area counts and the SHED volume was 1708 ft³, then the following rate would be calculated.

$$\text{ppm C Ethanol} = \frac{(3560 \text{ area counts, C}_2\text{H}_5\text{OH}) (68.3 \text{ ppm C C}_6\text{H}_6)}{(17521 \text{ area counts, C}_6\text{H}_6)}$$

$$\text{ppm C Ethanol} = 13.88 \text{ ppm C}$$

$$\text{Ethanol, g/test} = \text{ppm C C}_2\text{H}_5\text{OH} \times \text{Response factor} \times \text{SHED vol.}$$

$$\text{Ethanol, g/test} = 13.88 \text{ ppm C} \times 0.050 \text{ mg/ppm C ft}^3 \times 1708 \text{ ft}^3 \times 1 \text{ g/1000mg}$$

$$\text{Ethanol, g/test} = 1.19 \text{ g}$$

THIS PROCEDURE IS REPRINTED FROM A FINAL REPORT TO THE ENVIRONMENTAL PROTECTION AGENCY UNDER CONTRACT NO. 68-03-2377, "GASOHOL, TBA, MTBE EFFECTS ON LIGHT-DUTY EMISSIONS." (Reference 6)

APPENDIX B

B-1. DERIVATION OF EQUATIONS FOR CALCULATION OF EXHAUST EMISSIONS

APPENDIX B-1

DERIVATION OF EQUATIONS FOR CALCULATION OF EXHAUST EMISSIONS

The derivations are given in the order of occurrence of the equation in Section IV.B. of this report. Atmospheric values and atomic weight utilized are:

Standard Atmosphere (11)

Molecular Wt. = 28.966
 O_2 % by Vol. = 20.95 (23.14 by weight)
 O_2 Mol. Wt. = 32.000
 N_2 + % of Vol. = 79.05 (76.86 by weight)
 N_2 + Mol. Wt. = 28.162

Where N_2 + = Sum of all elements other than oxygen

Atomic Weights (11)

Carbon (C) = 12.011
 Hydrogen (H) = 1.0079 (use 1.008)
 Oxygen (O) = 15.9994 (use 16.000)

Exhaust Organic Matter Density:

Density_{HC} = 16.33 g/ft³ for carbon-to-hydrogen ratio of 1:1.85
 Fuel Fraction Carbon for 1:1.85 = $(1 \times 12.011) / (1 \times 12.011 + 1.85 \times 1.008) = 0.8656$
 HC Molecular Weight per Carbon Atom = 12.011/FFC
 Density OM = 16.33 × Ratio of Mol. Wts.
 = 16.33 ((12.011/EFC) / (12.011/0.8656))
 = 14.135/EFC g/ft³ (0.4493/EFC kg/m³)

(c) (3) CO_{mass} = CARBON MONOXIDE EMISSIONS

Hydrogen-to-Carbon Ratio of Fuel (HCR)

HCR = Atoms of H/Atoms of C
 = Ratio of weight amounts in fuel/Ratio of mol. wt.
 = (FFC/FFC) / (1.008/12.011)
 HCR = (FFH/1.008) / (FFC/12.011)

CO Correction for Water Vapor

Total H₂O = Water Due to Combustion + Humidity

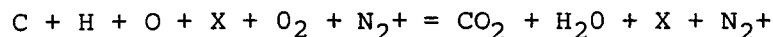
$$H_2O_{comb} \approx HCR \times CO_{2C} / 2$$

Therefore:

$$CO_C = (1 - 0.01 + 0.005 \times HCR \times CO_{2C} - 0.000323F) \times CO_{cm}$$

(c) (5) DF = Stoichiometric Percent CO₂/Summation of Carbon Containing Exhaust Components

Combustion Balance



N₂+ = Summation of all components in the atmosphere except oxygen

Number of Atoms \propto Relative Weight/Atm. Wt.

Number of Molecules \propto Relative Weight/Mol. Wt.

Therefore:

$$\frac{FFC}{12.011} + \frac{FFH}{1.008} + \frac{FFO}{16.000} + \frac{FFX}{MWX} + \frac{SAFR \times 0.2314}{32.000} + \frac{SAFR \times 0.7686}{28.162} =$$

$$\frac{FFC}{12.011} + \frac{0.5 \times FFH}{1.008} + \frac{FFX}{MWX} + \frac{SAFR \times 0.7686}{28.162}$$

SAFR = Stoichiometric Air to Fuel Ratio by Weight

Oxygen Balance (Number of Atoms)

$$\frac{2 \times FFC}{12.011} + \frac{0.5 \times FFH}{1.008} - \frac{FFO}{16.000} = \frac{2 \times SAFR \times 0.2314}{32.000}$$

$$SAFR = 11.514 \times FFC + 34.298 \times FFH - 4.322 \times FFO$$

Stoichiometric Percent CO₂

$$\begin{aligned} SPCO_2 &= (\text{Exhaust CO}_2 / \text{Total Exhaust}) \times 100 \\ &= ((FFC/12.011) / (FFC/12.011 + 0.5 \times FFH/1.008 + FFX/MWX + \\ &\quad SAFR \times 0.7686/28.162)) \times 100 \\ &= (FFC / (FFC + 5.958 \times FFH + 12.011 \times FFX/MWX + 0.3278 \times SAFR)) \times 100 \end{aligned}$$

Dilution Factor

$$DF = SPCO_2 / [CO_{2C} + (OM_C + CO_C) \times 10^{-4}]$$

Where:

OM_C = Organic matter in ppm carbon equivalent

APPENDIX C

- C-1. DERIVATION OF ENERGY BASED FUEL CONSUMPTION
- C-2. DERIVATION OF EQUATIONS FOR CALCULATION OF
EVAPORATIVE EMISSIONS

APPENDIX C-1

DERIVATION OF ENERGY BASED FUEL CONSUMPTION

$$\begin{aligned}\text{EBFC} &= (1/\text{FE}) \times (3785.4 \times \text{FDEN} / 453.6) \times \text{LHV} \\ &= 8.345 \times \text{FD} \times \text{LHV} / \text{FE}\end{aligned}$$

Where:

ESFC = Energy based fuel consumption, Btu/mile
FDEN = Fuel density, g/ml
LHV^a = Lower heating value of the fuel, Btu/lb
FE = Fuel economy, mpg
3785.4 = Conversion from gallon to ml
453.6 = Conversion from pound to gram

^aThe lower heating value has been used so as to be in agreement with the normal usage for engines.

APPENDIX C-2

DERIVATION OF EQUATIONS FOR CALCULATION OF EVAPORATIVE EMISSIONS

$$M_{OM} = MW_{OM}/MW_{AIR} \times ppmC \times 10^{-6} \times V_n \times DEN_{AIR} \times C_T \times C_P$$

Where:

MW_{OM} = Molecular Weight of organic matter per carbon atom

MW_{AIR} = 28.966

V_n = Net Volume of the Enclosure, ft^3

DEN_{AIR} = 34.69 g/ ft^3

C_T = 518.7/T; T = °R

C_P = $P_B/29.92$ P_B = in. Hg

$$M_{OM} = (0.208 MW_{OM}) \times V_n \times (ppmC \times Baro/Temp) \times 10^{-4}$$

Let:

K = $0.208 \times MW_{OM}$

C = $ppmC \times Baro/Temp$

Then:

$$M_{OM} = KV_n \times 10^{-4} (C_f - C_i)$$

| TECHNICAL REPORT DATA <i>(Please read Instructions on the reverse before completing)</i> | | |
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| 16. ABSTRACT <p>This report provides methods for the calculation of vehicle emissions and fuel consumption when nonstandard fuels are used. Methods of analyses, required for evaluation of alternate fuels, are included by reference or as Appendices to this report.</p> | | |
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