NEW MEXICO HYDROCARBON SOURCE ROCK EVALUATION PROJECT

BAR-S-BAR RANCH NO.1 FRE WELL SEC.23, T12N, R10E, SANTA FE CO., NEW MEXICO API NO. 30-049-20001 NORTHWEST AREA GEOCHEM JOB NO. 3820

Prepared

for

PROGRAM PARTICIPANTS

by

Dr. Geoffrey S. Bayliss and Dr. Rudy R. Schwarzer

GEOCHEM LABORATORIES, INC. 1143-C BRITTMOORE ROAD HOUSTON, TEXAS 77043 (713) 467-7011

> CONFIDENTIAL JUNE, 1989

NEW MEXICO HYDROCARBON SOURCE ROCK EVALUATION

WELL NAME:

BAR-S-BAR RANCH NO.1 FEE WELL

API NO.:

30-049-20001

AREA:

NORTHWEST

LOCATION:

SANTA FE COUNTY, NEW MEXICO SEC.23, T12N, R10E

GEOCHEM JOB NO.:

3820

TOTAL DEPTH:

4204 ft.

INTERVAL SAMPLED:

1940-2300 ft.

TOTAL NUMBER OF SAMPLES: 3

				AN	ALYSE	S	
GEOCHEM SAMPLE NUMBER	SAMPLE DEPTH	STRATIGRAPHIC INTERVAL	LITHO	TOC	ROCK-EVAL	KEROGEN	отнек
3820-001 3820-002 3820-003	1940-1990 2000-2020 2200-2300	Madera Madera Madera	X X	X X	X X X	X X X	

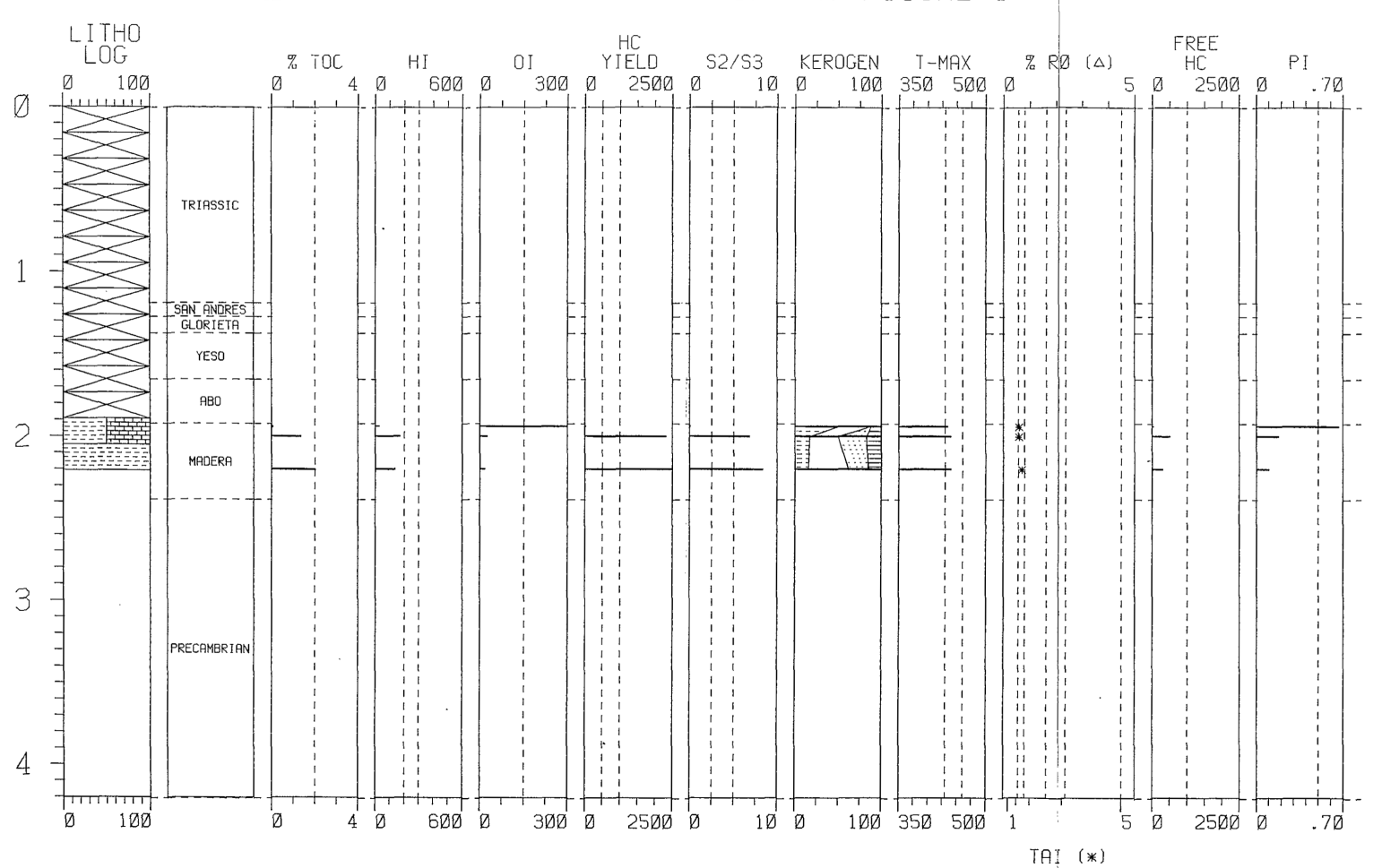


TABLE I

RESULTS OF TOTAL ORGANIC CARBON

NEW MEXICO HYDROCARBON SOURCE ROCK EVALUATION

BAR-S-BAR RANCH NO.1 FEE WELL SEC.23, T12N, R10E, SANTA FE COUNTY, NEW MEXICO API #30-049-20001

GEOCHEM SAMPLE NUMBER	DEPTH INTERVAL (feet)	TOTAL ORGANIC CARBON (% of Rock)
3820-001	1940-1990	0.07
3820-002	2000-2020	1.35/1.33
3820-003	2200-2300	2.02

TABLE II LITHOLOGICAL DESCRIPTIONS AND ORGANIC CARBON ANALYSES

NEW MEXICO HYDROCARBON SOURCE ROCK EVALUATION

BAR-S-BAR RANCH NO.1 FEE WELL SEC.23, T12N, R10E, SANTA FE COUNTY, NEW MEXICO API #30-049-20001

GEOCHEM SAMPLE NUMBER	DEPTH INTERVAL (feet)	LITHO DESCRIPTION	GSA NO.	ORGANIC CARBON (wt.%)
3820-001 -A	1940-1990	100% Limestone, fine crystalline,	10YR-6/	0.07
3820-002 -A	2000-2020	100% Shale, slightly calcareous, micaceous, grayish black.	101R-67	1.35/1.33
3820-003 -A	2200-2300	100% Shale, slightly calcareous, micaceous, grayish black.	N3	2.02

TABLE III

SUMMARY OF ORGANIC CARBON AND VISUAL REROGEN DATA

NEW MEXICO HYDROCARBON SOURCE ROCK EVALUATION

BAR-S-BAR RANCH NO.1 FEE WELL SEC.23, T12N, R10E, SANTA FE COUNTY, NEW MEXICO API #30-049-20001

GROCHEM SAMPLE	depth Interval	TOTAL ORGANIC	ORGANIC MATTER			AL ABUN LIZED P		ALTERATION	THERMAL ALTERATION	
number	IIMBER (feet) CARBON TYPE	Al	Am	H	¥	I	STAGE	INDEX		
3820-001 3820-002 3820-003	1940-1990 2000-2020 2200-2300	0.07 1.35/1.33 2.02	Am-H*;I H-W;Am-I:- H*:W;Am-I	0 0 0	50 17 15	.38 33 47	0 33 23	12 17 15	2 2 2 to 2+	2.2 2.2 2.3

LEGEND:

KEROGEN KEY

Predominant: Secondary: Trace 60-100% 20-40% 0-20%

Al = Algal

Am = Amorphous-Sapropel

Am* = Relic Amorphous-Sapropel

H = Herbaceous-Spore/Pollen

H* □ Degraded Herbaceous

W = Woody-Structured

U = Unidentified Material

I = Inertinite

C = Coaly

TABLE IV

RESULTS OF ROCK-EVAL PYROLYSIS ANALYSIS

NEW MEXICO HYDROCARBON SOURCE ROCK EVALUATION

BAR-S-BAR RANCH NO.1 FEE WELL SEC.23, T12N, R10E, SANTA FE COUNTY, NEW MEXICO API #30-049-20001

DEPTH INTERVAL (Feet)	TMAX (c)	S1 (mg/g)	S2 (mg/g)	S3 (mg/g)	PI	PC*	T.O.C. (wt.X)	Hydrogen Index	OXYGEN INDEX
1940-1990 2000-2020	435 440	0.04 0.51	0.02 2.32	0.34 0.34	0.67 0.18	0.05 0.23	0.07 1.34	28 173	485 25 16
	INTERVAL (Feet) 1940-1990 2000-2020	INTERVAL TMAX (Feet) (c)	INTERVAL (c) (mg/g) 1940-1990 435 0.04 2000-2020 440 0.51	INTERVAL (c) (mg/g) (mg/g) 1940-1990 435 0.04 0.02 2000-2020 440 0.51 2.32	INTERVAL (c) (mg/g) (mg/g) (mg/g) 1940-1990 435 0.04 0.02 0.34 2000-2020 440 0.51 2.32 0.34	INTERVAL (c) (mg/g) (mg/g) (mg/g) PI 1940-1990 435 0.04 0.02 0.34 0.67 2000-2020 440 0.51 2.32 0.34 0.18	INTERVAL (c) (mg/g) (mg/g) (mg/g) PI PC* 1940-1990 435 0.04 0.02 0.34 0.67 0.05 2000-2020 440 0.51 2.32 0.34 0.18 0.23	INTERVAL (Feet) (c) (mg/g) (mg/g) (mg/g) PI PC* (wt.X) 1940-1990 435 0.04 0.02 0.34 0.67 0.05 0.07 2000-2020 440 0.51 2.32 0.34 0.18 0.23 1.34	INTERVAL TMAX S1 S2 S3 T.O.C. HYDROGEN (Feet) (c) (mg/g) (mg/g) PI PC* (wt.7) INDEX 1940-1990 435 0.04 0.02 0.34 0.67 0.05 0.07 28 2000-2020 440 0.51 2.32 0.34 0.18 0.23 1.34 173

T.O.C. = Total organic carbon, wt.%

S2

■ Free hydrocarbons, mg Hc/g of rock

- Residual hydrocarbon potential (mg HC/g or rock)

S3 = CO2 produced from kerogen pyrolysis

(mg CO2/g of rock) PC* = 0.083 (S1 + S2)

Hydrogen

mg HC/g organic carbon Index

0xygen

= mg CO2/g organic carbon Index PΙ

= \$1/\$1 + \$2

- Temperature Index, degrees C. TMAX

TABLE V VISUAL KEROGEN ASSESSMENT WORKSHEET

	RANCH NO.1		ÆLL		M	DIGE	NOU:	S PC	PULATION (INT	ERF	RE	LED)				CH/			RAL ERIS	STIC	\$			RE	wo				ND/OR PULATION(S)	SUMMAR' ORGANIC
SEC.23, T	12N, R10E			0	TY	PE IC N	OF ATTE	R M	ATURATION IND	EX		•			ORG	OLOF	O MAT	ER	S ORG	ANK		TER			ORG	TYP ANI	E C N	OF IAT T	ER	M/	ATURATION INDEX	MATTER TYPE
T.D. 4,20			7777						REMARKS												//										REMARKS	
3820-001	1940-1990			Ш	П			П			T	V	\prod		1	\mathbb{T}				П	Ш	Ш				П	П	П				Am-H*;I
3820-002	2000-2020	•	+	Ш	Ш		$\top \top $			П	Ī	1		\mathbf{I}						П	\mathbf{H}	Ш			łΠ		П	П		\Box		H-W; Am-I;-
3820-003	2200-2300	!!!! !		###	TTI		Π			П	T	1	1	U	1			7	П	П	1				П	П	П	\sqcap	П			H*;W;Am-I
				Ш	Π	77						17	17	П	TΤ	\top	П	1	П	П	7111	тп		ПП		П	77	\top	П	\top		
		1 111111	<u> </u>	╫┸	111	11	+					Ħ	11	Ħ	77	11		╁	\sqcap	Ħ	1111	##		11111	\sqcap	Ħ	\top	11	77	\top		-
			*****	!!!!	111	77	#					##	11	\forall	11	11-		十	\vdash	Ħ	T 1 I I	ш		11111	Ħ	Ħ	Ħ	11	11	+		<u> </u>
			*****	₩.	111	#				\dashv	\top	#	11	11	11	11		╁	\vdash	††	† ##	###			11	†=†	77	++	Ħ	\top		<u> </u>
				ш	†††	-1-1-	+++	H		[-	+	††	++	++	++	++-	1	+	+	H	1111	Ш			†	H	++	++	17	\top		
				Ш	1 1	╅	+-1-1	++	 	-	-	╁	++	╁	+	╂╂╾	 	Н	H	╫	Ш	Ш			1	Н	++	╅╅	+	+		
			╂┼╂┾╬╂┾┫	╫╫╴	╀┼┼	++	++-	+++	 	+	+	₩	╁	╁	╅	╬	Н	+H	╌	╁┼	1111				 	╁	+	╁┼	╂	Н		
			4141111	 	} } }	+	++-	╌╂╍╂╌	 	-11	+	╁┼	╁╂	╌	++	╫		+	H	₩	1111	}}}}}	 		╁┼	₽	++	╅╋	+1	-		}
				##	+++		-	HH		$-\!$	+	₩	++	₩	-}-}	₩	Н-		Н-	₩	Ш	Ш		7777	╌	╌	++	╫	╂╉	-		
				##	$\sqcup \sqcup$	+	11.	- -	 	$-\!$	4	++	44	╌┦	-1-1	#	╀	4	Ш	11	-	Ш		4444	1-1-	₩	11	┵┵	- -	4		ļ
		1111111		TITI	$\bot \bot \downarrow$	44	11		<u> </u>	_11	1	Ц	11	44	-1-1	11			Ш	Ш	###	!!!!			ш	Ш	11	₩	11			<u> </u>
		,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,		TITI	\coprod	$\perp \perp \perp$				Ш			Ш	Ш		<u> 11</u>				Ш	3111	1111		111111	<u>1 L</u>	Ш		Ш	Ш			
				+};;[\prod				!		Т					\prod				\prod	HH	╫		++++	1]	П			\prod			
		 		11:1	Ш	\sqcap				П	T	П		П	11	П	П	П	П	П	H	1111				П	77	77	П	T		
				###	Ш		7-1-	- - -	 	┰	T	Π	11	77	71	T		\top	\vdash	П	1171	1111			1	11	$\dagger\dagger$	11	Ħ			<u> </u>
		11111111	.,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	1111	┼┼┼	++	11	HH	<u> </u>	- -{-{	1	††	††	11	++	++	1			H	1111	!!!!		11111	 - -	††	++	++	11			
		1111111		mr	╁═╁╌┠		+	++	 	-+1	+	╁	++	╫	++	╫	╁┼╴	+	+	╁	╁	 			 	Н	╁	╅╅	╁╉	+		
				Ш1.	╁┼╂	+	╂╂╸	┝╌┼╌╂╼		\dashv	+	₩	++	╁	++	╂╂╾	- -	+	- -	╀╀						 - -	4-1	11	┩┥	-		ļ
		33 H 11 H 11 H			ш	Ш	\perp		<u> </u>	$\perp \! \! \! \! \! \! \! \! \! \! \perp \! \! \! \! \! \! \! \!$		Ш	$\perp \perp$	ш		$\perp \perp$	LLi	\perp L		$\mathbf{L}\mathbf{L}$	Ш	ĦĦ	ШШ	Ш	<u> </u>	Ш.	$\perp \perp$	11.	\perp 1	.14	[1

LEGEND FOR SUMMARY DIAGRAM

DEPTH:

in feet

LITHO LOG:

see lithology symbols

STRATIGRAPHY:

by age

% TOC:

percent total organic carbon

HI: $\overline{\mathtt{OI}}$: Rock-Eval, Hydrocarbon Index = 100 S2(0/00 Wr)/TOC Rock-Eval, Oxygen Index = 100 S3 (0/00 Wt)/TOC

HC YIELD:

Rock-Eval, S2 peak (ppm)

<u>52/53:</u>

Rock-Eval, Ratio of S2 to S3 peak

KEROGEN:

see Kerogen symbols

T-MAX:

Rock-Eval, maximum temperature of S2 peak, in degrees Centigrade

%RO (△):

Vitrinite Reflectance (scale 0 to 5) Thermal Alteration Index (Scale 1 to 5)

TAI (*): FREE HC:

Rock-Eval, Si peak (ppm)

PI:

Rock-Eval, Productivity Index = S1/(S1+S2)

LITHOLOGIES

SHALE



SILICEOUS ROCKS



MUDSTONE



EVAPORITES



SILTSTONE



COAL



SANDSTONE



IGNEOUS ROCKS



CONGLOMERATE



VOLCANICS



BRECCIA



METAMORPHIC ROCKS



LIMESTONE



BASEMENT



DOLOMITE



OTHER





MISSING SECTION

KEROGE'I TYPES

AMOR?HOUS



HERBACEOUS MOODY



INERTINITE

APPENDIX A

Brief Description of Organic Geochemical analyses Carried Out by GeoChem

C1-C7 Hydrocarbon

The C_1 - C_7 hydrocarbon content and composition of sediments reflects source type, source quality and thermal maturity.

The C_1 - C_7 hydrocarbon content of well cuttings is determined by analyzing both a sample of the cuttings and the air space at the top of the can. The results of the two analyses are summed to give an inventory of the C_1 - C_7 hydrocarbon content of the well cuttings prior to any losses from the cuttings during the lapsed time period between collection at the wellsite and laboratory analysis.

The air space $C_1^-C_7$ hydrocarbon analysis involves taking a measured volume of the air space gas out of the can with a syringe and injecting same into a gas chromatograph. GeoChem uses a Varian Aerograph Model 1400 instrument equipped with a Porapec Q column. The gas sample is taken through the column by a carrier gas and before reaching the detector is separated into its various C_1 (methane), C_2 (ethane), C_3 (propane), iC₄ (isobutane), nC_4 (normal butane), and C_5 , C_6 , C_7 hydrocarbon components.

This particular analysis gives a complete separation of the C_1 - C_4 gas-range hydrocarbons and a partial separation of the C_5 - C_7 gasoline-range hydrocarbons. (A detailed C_4 - C_7 analysis, to be discussed later, involving a capillary column, effects a complete separation of this molecular range into its several individual molecular species.)

The electrical response of the various hydrocarbons as they reach the detector is recorded on a paper strip chart as a peak. This response is simultaneously fed to an integrator which computes the area of each peak. The concentration of C_1 - C_7 hydrocarbons in the air space, expressed as volumes of gas per million volumes of cuttings, is determined by a calculation involving the volume of cuttings, volume of air space in the can, volume of sample injected, volume of standard gas sample used in the calibration, calibration factor for C_1 , C_2 , C_3 , etc. determined by gc analysis of a standard gas sample, and the gc peak response.

The C_1 - C_7 hydrocarbon content of the cuttings is determined by degasification of a measured volume of cuttings (in a medium of a measured volume of water) in a closed blender, sampling of the air space at the top of the blender, and injection of a measured volume of gas into the gas chromatograph.

The C_1 - C_7 hydrocarbon data from the air space and cuttings gas analyses are summed to give a "restored" C_1 - C_7 hydrocarbon content of the cuttings.

Sample Washing and Hand-Picking of Uncaved Lithology Samples

The cuttings samples are washed to remove all drilling mud from the cuttings. Care is taken in the washing procedure not to remove any soft clays, claystones, etc. and any loose fine sand and silt. The washed cuttings are usually kept under water cover until picked, to prevent loss of any gasoline-range hydrocarbons. Using the C_1 - C_7 hydrocarbon data profile and the electrical well log supplied to us and our visual examination of the cuttings material under the binocular microscope, we carefully hand-pick and describe a suite of uncaved lithologies representative of the various strattgraphic zones penetrated by the well. The lithological data is used to compile a gross litho percentage log which is shown on all Figures. The 2-4 gram picked lithology samples are stored under water in small glass vials in those instances where we wish to run detailed C_4 - C_7 hydrocarbon analyses. This sample set is used not only for the C_4 - C_7 hydrocarbon analysis, but also for the visual kerogen and total organic carbon analyses. All remaining cuttings material is dried and packaged in labelled plastic bags for possible C_{15+} soxhlet extraction and/or eventual return to the client. Sample material from this study will be retained at GeoChem until advised of disposition.

Detailed C4-C7 Hydrocarbon

The C_4 - C_7 gasoline-range hydrocarbon content of sediments reflects source quality, thermal maturation and organic factes. Compositional data can be used in crude oil-parent rock correlation work.

The C_4 - C_7 hydrocarbon content and detailed molecular composition of hydrocarbon, in hand-picked lithologies, is determined by a gc analysis of the light hydrocarbon extracted from 1-2 gram cuttings samples macerated in a microblender. A measured volume of sample is placed in a scaled microblender along with a measured volume of hot water. The rock sample is pulverized by the blades of the blender. A sample of the liberated light hydrocarbons which collect in the air space at the top of the blender is injected into our Varian Aerograph 1400 gc unit which is equipped with a capillary column. Data recording, computations, etc. are comparable to those used for the C_1 - C_7 analysis discussed previously in this report. Hydrocarbon concentration is expressed as volume gas per million volumes of cuttings.

Organic Çarbon

The total organic carbon content of a rock is a measure of its total organic richness. This data is used, in conjunction with visual kerogen and C_1 - C_4 , C_4 - C_7 and C_{15+} hydrocarbon content of a rock, to indicate the hydrocarbon source quality of roci's.

The procedure for determining the total organic carbon content of a rock involves drying the sample, grinding to a powder, weighing out 0.2729 gram sample into a crucible, acidizing with hot and cold hydrochloric acid to remove calcium and magnesium carbonate, and carbon analysis by combustion in a Leco carbon analyzer.

We run several blank crucibles, standards (from rings of known curbon content) and duplicate rock samples in this analysis at no additional charge to the client for purposes of data quality control.

$C_{{f 15+}}$ Soxhlet Extraction, Deasphaltening and Chromatographic Separation

The amount and composition of the organic matter which can be solvent-extracted from a rock reflects source quality and source type. C^{13}/C^{12} carbon isotopic, high mass spectrometric and gc analyses of the paraffin-naphthene and aromatic hydrocarbon fractions of the soluble extract gives data which is used in crude oil-parent rock correlations.

This analysis involves grinding of a dry rock sample to a powder and removal of the soluble organic matter by soxhlet extraction using a co-distilled toluene-methanol azeotrope solvent. Where the amount of available sample material permits, we like to use at least 100 grams of rock for this analysis.

The extracted bitumen is separated into an asphaltene (ASPII) and a pentane soluble fraction by normal pentane precipitation. The pentane soluble components are separated into a C_{15} , paraffin-naphthene (P-N) hydrocarbon, C_{16} , aromatic hydrocarbon (AROM) and C_{15} , nitrogen-sulfur-oxygen containing fraction (NSO) by adsorption chromatography on a silica gel-alumina column using pentane, toluene and toluene-methanol azeotrope chants.

APPENDIX A (continued)

GC Analysis of C15+ Paraffin-Naphthene (P-N) Hydrocarbons

The content and molecular composition of the heavy C_{15+} paraffin-naphthene (P-N) hydrocarbons of rocks, as determined by ge analysis, reflects source quality, source type and degree of thermal maturation.

In this analysis, we subject a very small fraction of the total amount of the P-N fraction extracted from a rock sample to gc analysis. The gas chromatograph is a Varian Aerograph Model 1400 equipped with a solid rod injection system and a cutectic column.

The calculated C.P.I. (carbon preference index) values for the normal paraffin data is defined as the mean of two ratios which are determined by dividing the sum of concentrations of odd-carbon numbered n-paraffins by the sum of even-carbon numbered n-paraffins. The C.P. Indices A and B were obtained by the formulas:

C. P. Index A =
$$\frac{C_{21} + C_{23} + C_{26} + C_{27}}{C_{22} + C_{26} + C_{28}} + \frac{C_{21} + C_{23} + C_{26} + C_{27}}{C_{20} + C_{22} + C_{24} + C_{26}}$$
C. P. Index B =
$$\frac{C_{25} + C_{27} + C_{29} + C_{31}}{C_{26} + C_{28} + C_{30} + C_{32}} + \frac{C_{25} + C_{27} + C_{29} + C_{31}}{C_{24} + C_{26} + C_{28} + C_{30}}$$

Visual Kerogen

A visual study of kerogen, the insoluble organic matter in rocks, can indicate the relative abundance, size, and state of preservation of the various recognizable kerogen types and thereby indicate the hydrocarbon source character of a rock. The color of the kerogen can be used to indicate the state of thermal maturity of the sediments (i.e. their time-temperature history). Thermal maturation plays an important role in the generation of hydrocarbons from organic matter, and also affects the composition of reservoired hydrocarbons.

Our procedure for visual kerogen slide preparation involves isolation of the organic matter of a rock by removal of the rock material with hydrochloric and hydrofluoric acid treatment and heavy liquid separation. This procedure is comparable to that used by the palynologist except it does not include an exidation stage. (The exidation treatment is deleted from our procedure because it removes a great deal of kerogen and blanches any remaining kerogen to an extent whereby it is useless for our kerogen color observations.) The kerogen residue is mounted on a glass slide and is examined visually under a high power microscope.

Vitrinite Reflectance

Measurement of the reflectivity of vitrinite particles (KRo) present in the kerogen isolated from sedimentary rocks provides a method of determining the state of maturation, and the diagenetic (time-temperature) history of the organic matter present in the sediments.

The kerogen, obtained from a 25 gram aliquot of crushed rock by the acid procedure previously discussed, is dried and embedded in a Bioplastic plug. The surface of the plug is polished using 0.05 micron alumina and the reflectivity determined under oil using a Ziess high resolution microscope. A minimum of 40 values are required to adequately determine the Maturation Rank.

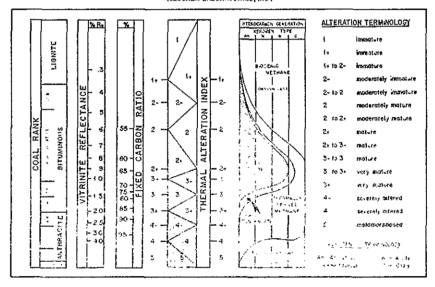
Fluorescence Spectrophotometric Analysis

Fluorescence spectrophotometry can be used to characterize and fingerprint crude oils, establish crude oil-source rock relationships, and to measure the hydrocarbon source potential of fine-grained sediments.

A one (1) microliter aliquot of either (i) a crude oil or (ii) the solvent extractable rock bitumen, is passed through an alumina/silica gel micro column and the C₁₀₊ aromatic hydrocarbons isolated. The aromatic hydrocarbon is diluted and the emission and excitation spectra determined at 240 nm and 420 nm using a Perkin-Elmer Model 512 Double Beam Fluorescence Spectrophotometer.

GEOTHERMAL DIAGENETIC CRITERIA

(GEOCHEM LABORATORIES, INC.)



```
:DEPTH: QTV :TMAX: S 1 : S 2 : S 3 : P I :S2/S3 : P C : TOC : H I : O I
                                  0.04: 15.93: 0.59: 5.53: 123: 7:
               0.32: 6.85: 0.43:
            289e: 06-13089 IME = 5 ВТЕЙК GRADIENÇŽСЬЕ: 4 ТЕЙРГЕТОР 1432 390
INIT TEMP =
              S 1 = 000000000 S 2 = 00000000 S 3 ≠ 00000618
                                             "GYCLE: 4 SCALE = 1/32
                                  STANDARD 
                  06-13-89
            DATE:
                                                        TRAP STOP T = 390
                  ISO TIME = 5 TEMP GRADIENT=25
INIT TEMP =
            250
                   6.00 53= 0.43 54 = 0.00
                                                              TMAX = 43
OTHER STD : S2=
STD DTY = 100.0 S 1 = 00001002  S 2 = 00019542  S 3 = 00002643
                                                             = 439
                                                        TMAX
```

```
The Theory
:DEPTH: QTY :TMAX: S 1 : S 2 : S 3 : P I :S2/S3 : P C : TOC : H I : O I
3920-05: 98.3: 440: 0.30: 2.74: 0.33: 0.10: 8.30: 0.25: 2.02: 135: 16:
          06-13-89 ANALYSIS CYCLE: 4 SCALE = 1/32
       DATE:
           ISO TIME = 5 TEMP GRADIENT=25 TRAP STOP T = 390
INIT TEMP = 250
:DEPTH: QTV :TMAX: S 1 : S 2 : S 3 : P I :S2/S3 : P C : TOC : H I : O I :
3920-002: 95.5: 440: 0.51: 2.32: 0.34: 0.18: 6.82: 0.23: 1.34: 173: 25:
          06-13-89 ANALYSIS CYCLE: 4 SCALE = 1/32
       DATE:
INIT TEMP = 250 ISO TIME = 5 TEMP GRADIENT=25 TRAP STOP T = 390
:DEPTH: QTV:TMAX: S 1 : S 2 : S 3 : P I :S2/S3 : P C : TOC : H I : O I :
3%20-co1: 98.8: 435: 0.04: 0.02: 0.34: 0.67: 0.05: 0.00: 0.07: 28: 485:
```