

GEOCHEMICAL SERVICE REPORT

Prepared for STATOIL

ANALYTICAL DATA
HALTENBANKEN STUDY

March 1984

CHESTER STREET : CHESTER CH4 8RD - ENGLAND

COMPANY PROPRIETARY

SUPPLIMENTARY GEOCHEMICAL DATA

FOR WELL 6407/1-2.

DATA REPORT ONLY

DATA SUPPLIED BY GEOCHEM LABORATORIES (U.K.) LIMITED

MARCH 1984.

SAMPLE IDENTIFICATION AND ANALYTICAL FORMAT

WELL	DEPTH (METRES)	STATOIL SAMPLE CODE	SAMPLE TYPE	ANALYTICAL SCHEME A B C D E
6407/1-2	3551.5 3595 3661.4 3688.16 4210 4285 4285 4360 4366 4510 4560 DST 1	E 17199 E 17147 E 17804 E 17805 E 17725 E 17730 E 17730 E 17748 E 17763 E 17762	EXTRACT " " " " POWDER " CUTTINGS " POWDER " CONDENSATE	

- A TOC, ROCKEVAL PYROLYSIS
- B C_{15+} EXTRACTION AND CHROMATOGRAPHY, PYROLYSIS-GC, CARBON ISOTOPES ON KEROGEN
- CAPILLARY CHROMATOGRAMS OF PARAFFIN-NAPHTHENES AND AROMATICS, GC/MS (6 ION), CARBON ISOTOPE COMPOSITION OF PARAFFIN-NAPHTHENES AND AROMATICS
- D CARBON ISOTOPE COMPOSITION OF TOTAL EXTRACT
- E API GRAVITY, WHOLE OIL CHROMATOGRAM, CARBON ISOTOPE COMPOSITION OF WHOLE OIL

TABLE 1
ORGANIC CARBON RESULTS AND GROSS LITHOLOGIC DESCRIPTIONS

	OndAir	IC CARBON NEGOLIO AND CHOOS II III CE COM		
GEOCHEM SAMPLE NUMBER	DEPTH	GROSS LITHOLOGIC DESCRIPTION	G S A Colour Code	TOTAL ORGANIC CARBON (Wt. % of Rock)
WELL 640	7/1-2		-	
867-015	4285m	POWDER		7.38
867-016	4285m	POWDER		4.59
867-012	4360m	A 75% Shale, platy, mod. hard, non- calc., minor caving, greyish brownish black, turbo drilled	5YR3/:	1 1.65
		B 20% Sandstone, blocky, mod. hard, non calc., matrix, fine grained, yellowish grey	5Y8/1	
		C 5% LCM - bitumen Minor coal	-	
867-014	4366m	A 70% Shale, as 867-012A turbo drilled B 25% Sandstone, as 867-012B C 5% LCM - bitumen Minor sand, coal and LCM - paint	5YR3/ 5Y8/1	
867-017	4510m	POWDER		29.50
867-013	4560m	A 60% Shale, as 867-012A B 35% Coal, blocky, mod. hard, black C 5% Quartz sand, unconsolidated, fine grained, white Minor sandstone	5YR3/ N1 N9	1 8.50 7.51

TABLE 2

ROCKEVAL PYROLYSIS

SAMPLE NUMBER	DEPTH	<u>S1</u> (mg/g)	<u>S2</u> (mg/g)	<u>S3</u> (mg/g)	HYDROGEN INDEX	PRODUCTION INDEX	Tmax (°C)
Well 6407/	1-2						
867-015	4285m	2.85	16.00	3.37	0.22	0.15	437
867-016	4285m	0.36	5.02	2.76	0.11	0.07	443
867-012A	4360m	0.29	0.53	0.49	0.03	0.35	445
867-014A	4366m	0.36	0.81	0.80	0.05	0.31	442
867-017	4510m	1.90	36.80	1.60	0.12	0.05	462
867-013A	4560m	1.48	12.72	0.84	0.15	0.10	458
867-013B	4560m	7.66	165.16	7.83	0.22	0.04	457

TABLE 3a
CONCENTRATION (PPM) OF EXTRACTED C₁₅₊ MATERIAL IN ROCK

			Н	DROCARBO	NS .	NO	N HYDRO	CARBONS	5
GEOCHEM SAMPLE NUMBER	DEPTH	TOTAL EXTRACT	Paratin Parating	Aromatics	TOTAL	Preciped res	Elused 5	HOT HEO'S	Sulphur
WELL 6407/1	2								
867-015	4285	19341	2714	2934	5648	11044	1747	901	0
867-016	4285	3332	329	270	599	2614	78	41	0
867-017	4510	4190	468	509	9 77	3029	123	60	0
867-013A	4560	1528	65	134	199	1093	116	120	0

TABLE 3b COMPOSITION (NORMALISED %) OF C15+ MATERIAL EXTRACTED FROM ROCK

GEOCHEM		HYDROC	CARBONS		NON HYDROCARBONS			
SAMPLE NUMBER	DEPTH	Paraffin — Naphthenes	Aromatics	Preciptd. Asphaltenes	Eluted NSO's	Non eluted NSO's	Sulphur	
WELL 6407/1-2				۸			•	
867-015 867-016 867-017 867-013A	4285 4285 4510 4560	14.03 9.87 11.18 4.24	15.17 8.12 12.15 8.79	57.10 78.46 72.29 71.52	9.03 2.33 2.94 7.58	4.66 1.22 1.44 7.88	0.00 0.00 0.00 0.00	
CONDENSATES 867-022 6407/	/1-2 DST1	72.69	24.17	0.91	1.16	0.83	0.24	

TABLE $^{3\text{C}}$ SIGNIFICANT RATIOS (%) OF C₁₅₊ FRACTIONS AND ORGANIC CARBON

GEOCHEM SAMPLE NUMBER	DEPTH	ORGANIC CARBON (wt. %)	HYDROCARBONS TOTAL EXTRACT	HYDROCARBONS ORG. CARBON	TOTAL EXTRACT	P-NAPHTHENES
WELL 6407/1-	2					
WELL 6407/1-3	<u>2</u> 4285	4.31	29.20	13.11	44.87	0.93
	_	4.31 5.40	29.20 17.99	13.11 1.11	44.87 6.17	0.93 1.22
867-015	 4285					

TABLE 4 COMPOSITION (NORMALISED %) OF C₁₅₊ PARAFFIN — NAPHTHENE HYDROCARBONS

GEOCHEM SAMPLE NUMBER	-021	-020	-018	-019	-024	-015
DEPTH	3551.5m	3595m	3661.4	3688.16m	4210m	4285m
SAMPLE TYPE			WELL 64	07/1-2		,
nC ₁₅	14.79	12.98	9.54	10.68	18.52	2.67
nC ₁₆	13.81	12.60	10.57	10.95	15.18	6.89
nC ₁₇	11.37	11.43	10.41	9.89	14.21	9.82
nC ₁₈	9.17	9.21	8.66	8.93	11.98	10.42
nC ₁₉	8.44	8.72	8.74	8.41	9.75	10.59
nC ₂₀	7.33	7.56	8.03	7.36	7.80	9.39
nC ₂₁	5.99	5.91	7.00	6.57	4.87	8.35
nC ₂₂	5.38	5.91	6.52	5.87	3.06	6.55
nC ₂₃	4.40	4.75	5.56	. 5.08	2.65	6.03
nC ₂₄	3.79	4.36	4.85	4.90	2.23	5.17
nC ₂₅	2.93	3.78	4.37	4.20	1.95	4.74
nC ₂₆	2.81	3.20	3.90	3.77	1.67	4.13
nC ₂₇	2.32	2.81	3.26	3.50	1.53	3.70
nC ₂₈	2.08	1.94	2.70	2.71	1.11	2.84
nC ₂₉	1.71	1.74	2.15	2.28	1.11	2.50
nC ₃₀	1.22	1.07	1.43	1.75	0.70	1.98
nC ₃₁	1.10	0.87	0.87	1.14	0.56	1.55
nC ₃₂	0.61	0.48	0.64	0.79	0.42	1.03
nC ₃₃	0.24	0.29	0.40	0.53	0.28	0.69
nC ₃₄	0.24	0.19	0.24	0.44	0.28	0.60
nC ₃₅	0.24	0.19	0.16	0.26	0.14	0.34
PARAFFIN	.39.42	49.05	45.14	39.35	44.46	27.22
ISOPRENOID	6.80	8.51	6.71	6.10	6.32	3.16
NAPHTHENE	53.78	42.44	48.15	54.55	49.23	69.62
CPI INDEX A	0.96	0.97	1.00	1.00	1.05	1.06
CPI INDEX B	1.01	1.12	1.03	1.04	1.11	1.07
PRISTANE/PHYTANE	1.14	2.20	1.25	1.27	1.62	1.45
PRISTANE/nC	0.81	1.04	0.79	0.88	0.62	0.70

TABLE 4 COMPOSITION (NORMALISED %) OF C₁₅₊ PARAFFIN — NAPHTHENE HYDROCARBONS

GEOCHEM SAMPLE NUMBER	-016	-017	-013A	022
DEPTH	4285m	4510m	4560m	DST 1
SAMPLE TYPE	(WELL 6407/1-2	.)	6407/1-2
nC ₁₅	8.86	11.21	0.39	15.01
nC ₁₆	12.00	14.27	1.07	13.56
nC ₁₇	11.62	12.90	6.30	11.54
nC ₁₈	11.43	12.37	11.35	10.25
nC ₁₉	8.86	10.25	13.00	8.66
nC ₂₀	6.76	8.56	12.22	8.08
nC ₂₁	5.71	6.87	11.64	6.20
nC ₂₂	4.86	5.60	9.99	5.92
nC ₂₃	4.10	4.02	9.02	4.76
nC ₂₄	4.38	3.49	7.18	4.04
nC ₂₅	3.90	2.75	6.40	3.03
nC ₂₆	3.43	2.22	4.36	2.45
nC ₂₇	3.33	1.80	3.20	1.88
nC ₂₈	2.57	1.16	1.65	1.44
nC ₂₉	2.29	0.95	0.97	1.15
nC ₃₀	1.43	0.53	0.29	0.72
nC ₃₁	2.19	0.32	0.29	0.43
nC ₃₂	0.86	0.21	0.19	0.29
nC ₃₃	0.67	0.21	0.19	0.29
nC ₃₄	0.48	0.21	0.19	0.14
nC ₃₅	0.29	0.11	0.10	0.14
PARAFFIN	29.56	48.94	59.73	43.48
ISOPRENOID	3.52	5.43	3.19	6.52
NAPHTHENE	66.92	45.63	37.08	50.00
CPI INDEX A	1.00	1.01	1.10	0.96
CPI INDEX B	1.20	1.10	1.24	1.04
PRISTANE/PHYTANE	1.91	3.04	0.96	1.36
PRISTANE/nC ₁₇	0.67	0.65	0.42	0.75

TABLE 5

CARBON ISOTOPES (PDB O/00)

GEOCHEM SAMPLE NUMBER	<u>DEPTH</u>	OIL	TOTAL EXTRACT	PARAFFIN- NAPHTHENE	AROMATICS	KEROGEN
WELL 6407/1-2 867-021 867-020 867-018 867-019 867-024 867-015A 867-016 867-017 867-013A	3551.5m 3595m 3661.4m 3688.16m 4210m 4285m 4285m 4510m 4560m	- - - - - -	- -28.95 -29.76 -27.22 - -	-28.64 -30.23 -29.34 -29.77 -27.83 -27.98 -28.17 -27.98 -28.37	-26.86 * -28.91 -29.28 -26.71 -27.08 -26.73 -27.21 -27.94	-28.23 -26.22 -27.14 -27.18
			CONDENSAT	<u>es</u>		
GEOCHEM SAMPLE NUMBER	DEPTH	OIL	TOTAL EXTRACT	PARAFFIN- NAPHTHENE	AROMATICS	KEROGEN
WELL 6407/1-2 867-022	DST-1	-26.94	-	-29.99	-29.08	

^{*} insufficient material

TABLE 6

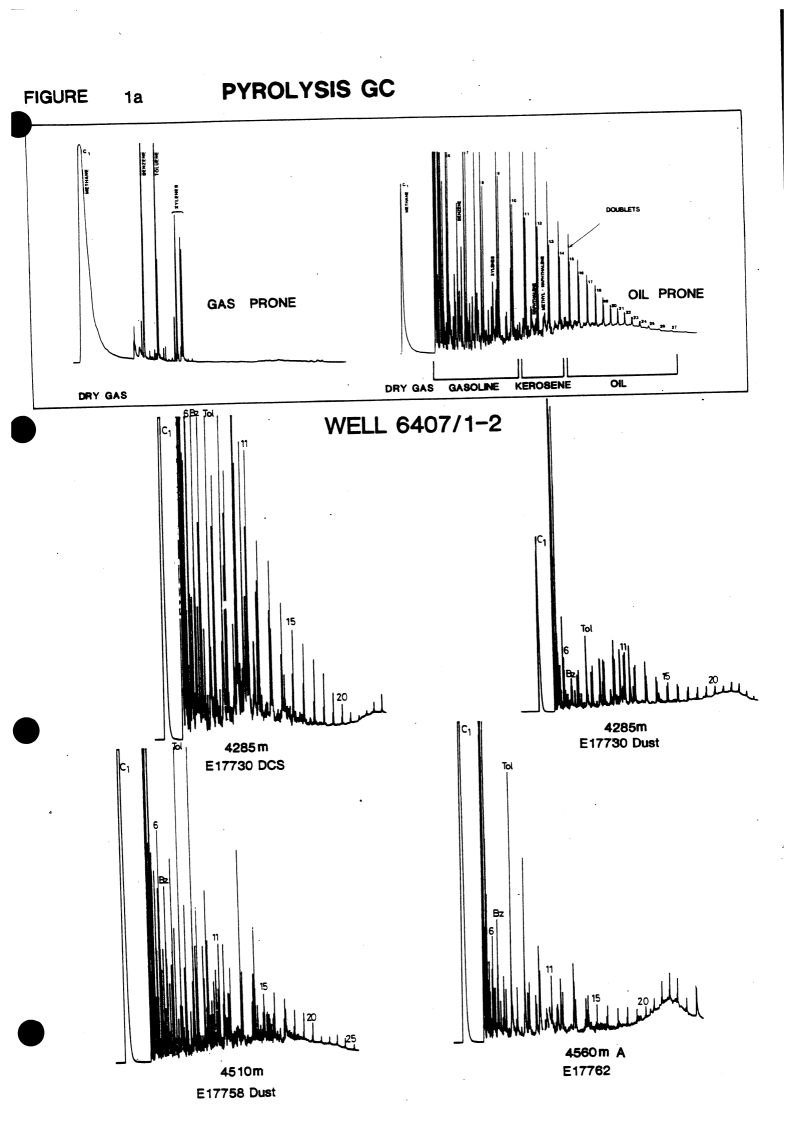
CONDENSATE SAMPLES - API GRAVITIES

GEOCHEM SAMPLE NUMBER	WELL	DEPTH	<u>°API</u>	
867-022	6407/1-2	DST 1	48.53	

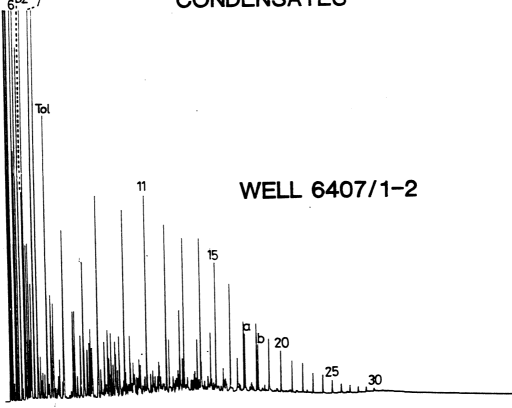
TABLE 7

MOLECULAR MATURATION PARAMETERS

GEOCHEM SAMPLE NUMBER	DEPTH	C ₂₉ 20S (coc) C ₂₉ 20R (coc)	C ₂₉ 20R(\$\beta\$) C ₂₉ 20R(\$\beta\$)	C ₂₇ (20S) Diasteranes C ₂₇ (20R) Diasteranes	$\frac{C_{31}}{C_{31}}$ (22S) homohopane x 100 $\frac{C_{31}}{C_{31}}$ (22S) + $\frac{C_{31}}{C_{31}}$ (22R) homohopane	C ₃₀ moretane	Tm Ts	MPI = $\frac{1.5 (2mp+3mp)}{p + 1mp + 9mp}$
Well 6407/	1-2							
867-021	3551.5m	1.00	1.18	1.40	57%	0.07	0.78	0.80
867-020	3595m	0.93	0.91	1.60	57%	0.12	2.35	0.62
	3661.4m	1.40	1.57	1.23	. 60 %	0.09	0.74	0.69
867-018	3688.16m	0.86	1,10	1.58	54%	0.10	1.08	0.74
867-019		0.81	1.10	1.38	53%	0.11	2.18	0.83
867-024	4210m		1.08	1.33	54%	0.09	2.30	0.77
867-015	4285m	0.85	1.02	0.66	55%	0.12	2.04	0.78
867-016	4285m	0.66		0.97	50%	0.13	0.76	0.91
867-017	4510m	0.75	1.18		60%	0.14	_	1.19
867-013	4560m	0.90	1.08	1.36		0.09	0.75	0.57
867-022	DST 1	1.27	1.51	1.17	60%	0.09	0.75	

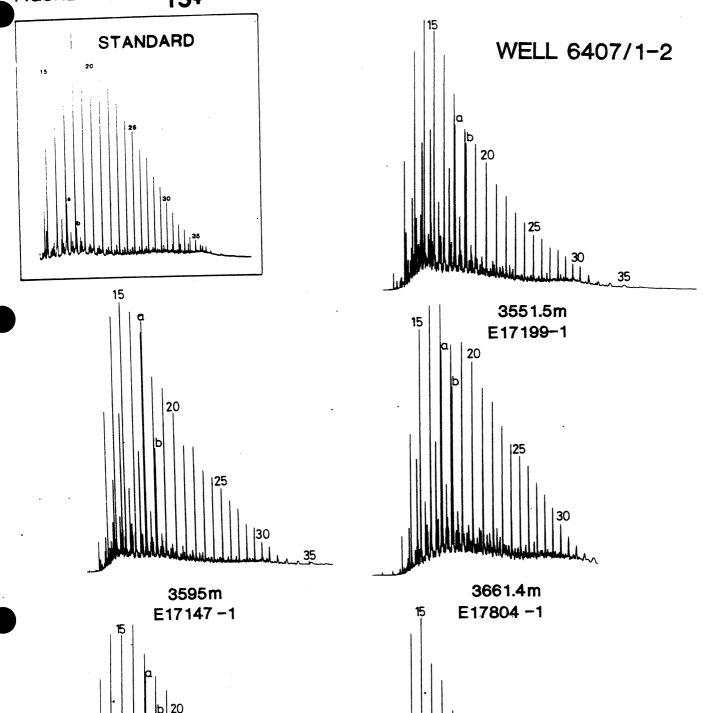






DST 1





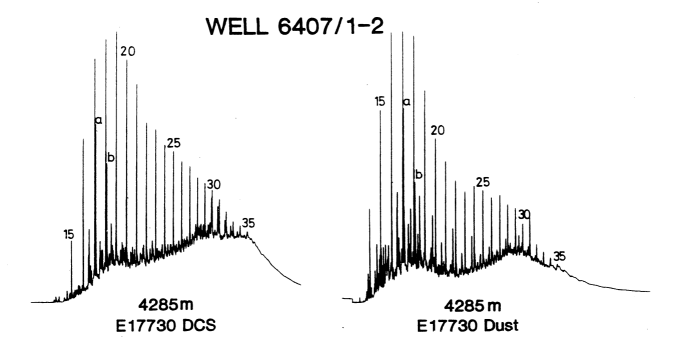


3688.16m

E17805-1

4210m E17725-1

FIGURE 3b C₁₅₊ PARAFFIN - NAPHTHENES



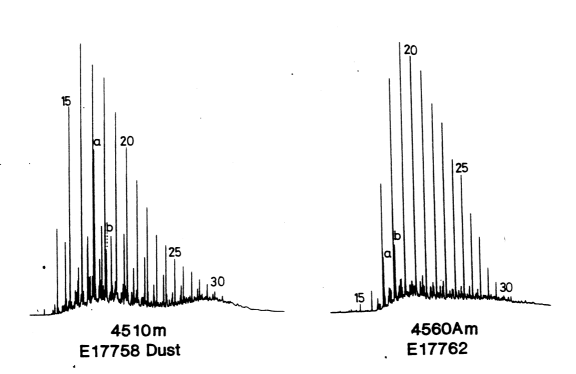
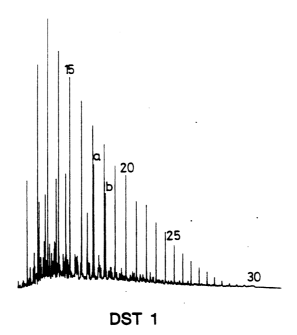


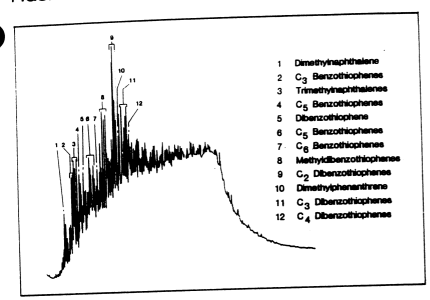
FIGURE 3d C15+ PARAFFIN - NAPHTHENES

CONDENSATES

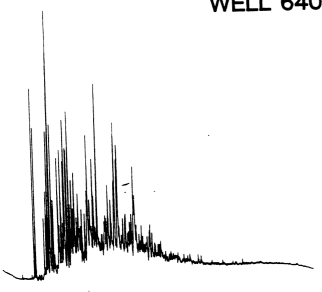
WELL 6407/1-2



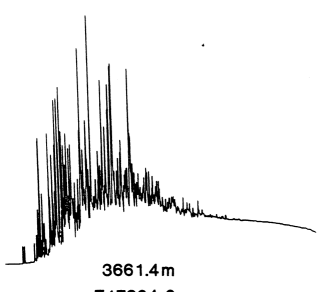
AROMATIC CHROMATOGRAMS



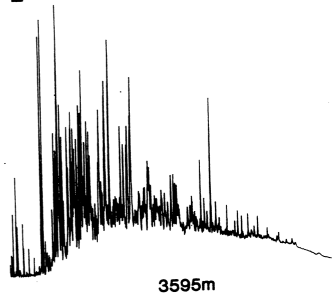




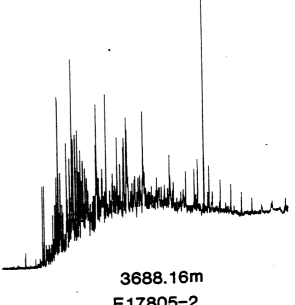
3551.5m E17199-2



E17804-2



E17147-2

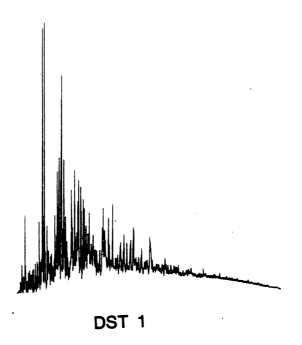


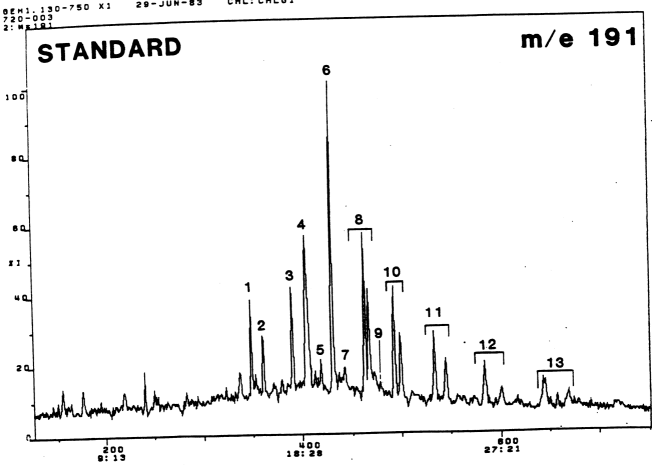
E17805-2

AROMATIC CHROMATOGRAMS

CONDENSATES

WELL 6407/1-2





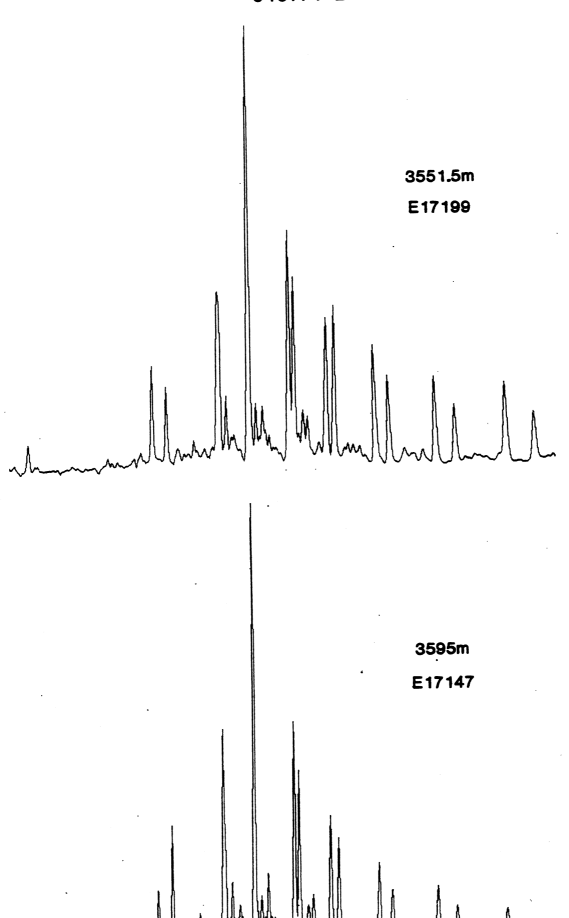
LIST OF IDENTIFIED TRITERPANES

1		176H TRISNORHOPANE (C27)
2	•	176H TRISNORHOPANE (C27)
3		BISNORHOPANE (C ₂₈)
4		174 H NORHOPANE (C29)
5		NORMORETANE (C ₂₉)
6		176H HOPANE (C30)
7		17∝H MORETANE (C ₃₀)
8	(22S) (22R)	17xH HOMOHOPANES (C31)
9		GAMMACERANE
	(22S) (22R)	BISHOMOHOPANE (C ₃₂)
	(22S) (22R)	TRISHOMOHOPANES (C ₃₃)
	(22S) (22R)	TETRAHOPANES (C ₃₄)
13	(22S) (22R)	HOPANES (C ₃₅)

FIGURE 5a

MASS FRAGMENTOGRAMS

TRITERPANES m/e 191



MASS FRAGMENTOGRAMS **FIGURE** 5b TRITERPANES m/e 191 6407/1-2



3661.4m

E17804

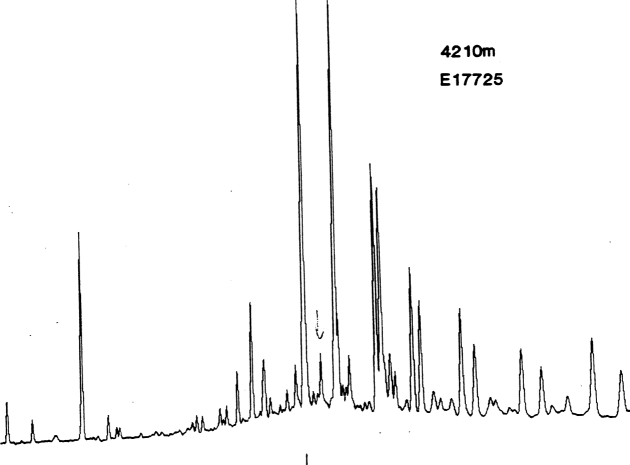
E17805

FIGURE 5c



TRITERPANES m/e 191

6407/1-2



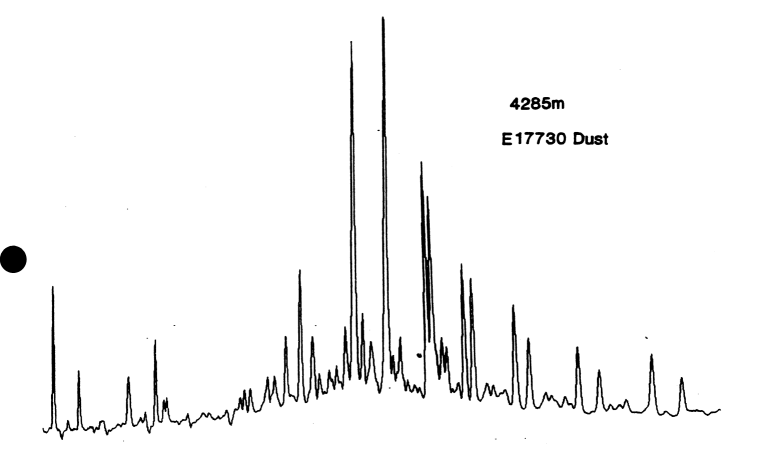
4285m

E17730 DCS

FIGURE 5d

MASS FRAGMENTOGRAMS

TRITERPANES m/e 191



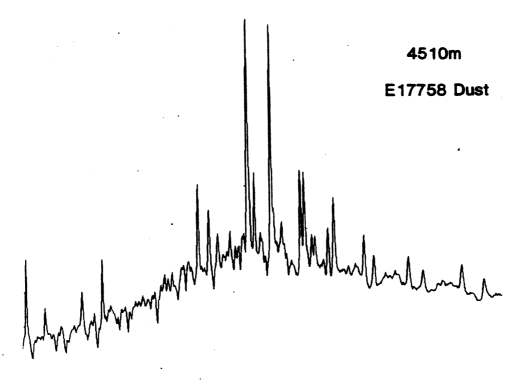


FIGURE 5e

MASS FRAGMENTOGRAMS

TRITERPANES m/e 191

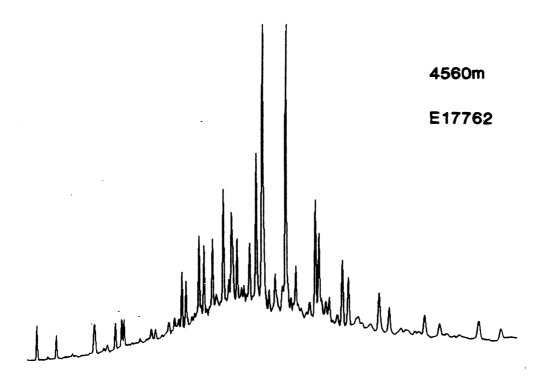
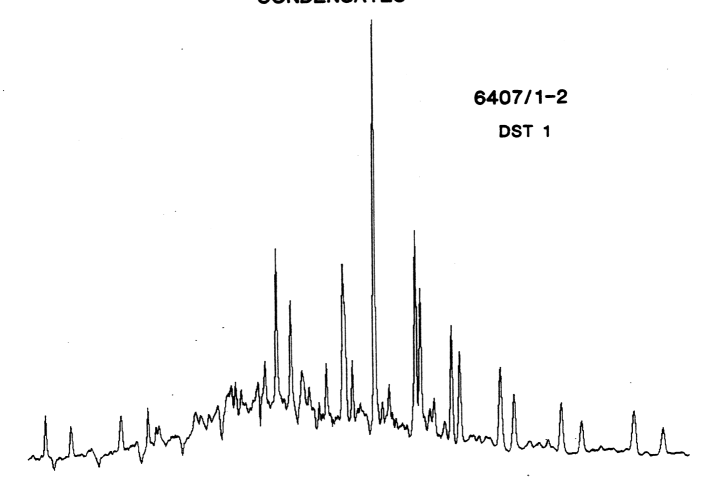
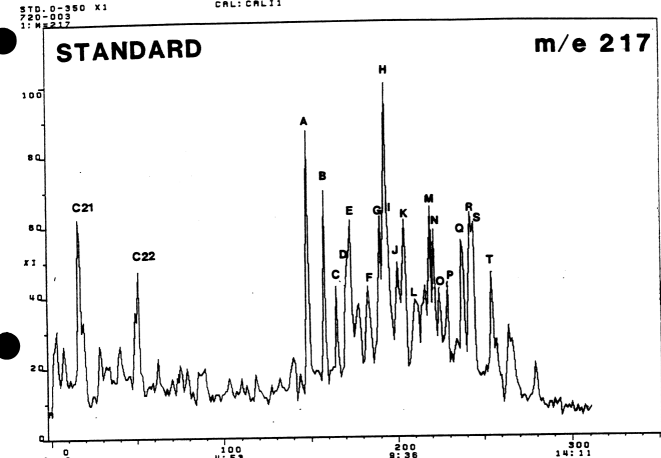


FIGURE 5i

MASS FRAGMENTOGRAMS TRITERPANES m/e 191 CONDENSATES





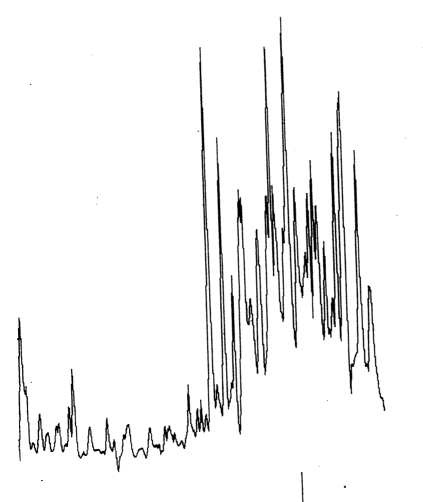
LIST OF IDENTIFIED STERANES

- A C27 DIACHOLESTANE (20S)
- B C₂₇ DIACHOLESTANE (20R)
- C C₂₇ DIACHOLESTANE (20S)
- D C₂₇ DIACHOLESTANE (20R)
- E C₂₈ METHYL DIACHOLESTANE (20S)
- F C28 METHYL DIACHOLESTANE (20R)
- G C₂₈ METHYL DIACHOLESTANE (20S)
- H C₂₉ ETHYL DIACHOLESTANE (20S)
- I C27 CHOLESTANE (20S) & C28 METHYL DIACHOLESTANE (20R)
- J C₂₇ CHOLESTANE (20R)
- K C29 ETHYL DIACHOLESTANE (20R)
- L C₂₉ ETHYL DIACHOLESTANE (20S)
- M C₂₈ METHYL CHOLESTANE (20S)
- N C₂₉ ETHYL DIACHOLESTANE (20R) .
- O C₂₈ METHYL CHOLESTANE (20S)
- P C₂₈ METHYL CHOLESTANE (20R)
- Q C₂₉ ETHYL CHOLESTANE (20S) $[\sim \sim]$
- R C_{29} ETHYL CHOLESTANE (20R) [/3/3]
- S C₂₉ ETHYL CHOLESTANE (20S) [/3/3]
- T C29 ETHYL CHOLESTANE (20R) [~ ~]

FIGURE 6a

MASS FRAGMENTOGRAMS
STERANES m/e 217

6407/1-2



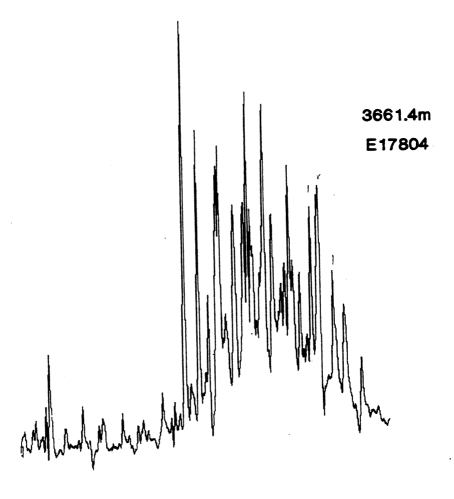
3551.5m E17199

3595m

E17147

FIGURE 6b

MASS FRAGMENTOGRAMS STERANES m/e 217



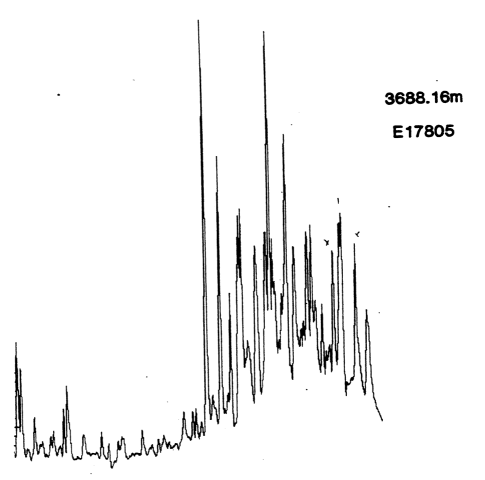
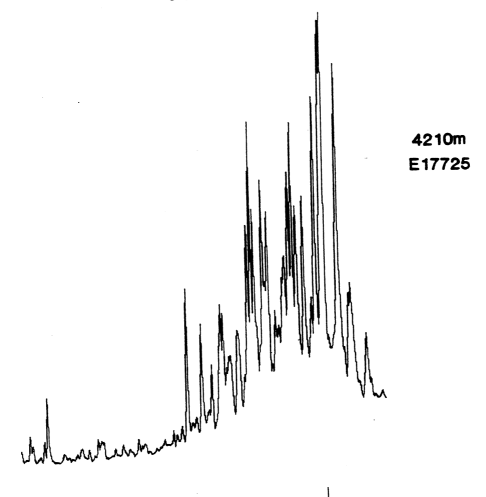


FIGURE 6c MASS FRAGMENTOGRAMS
STERANES m/e 217



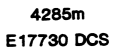
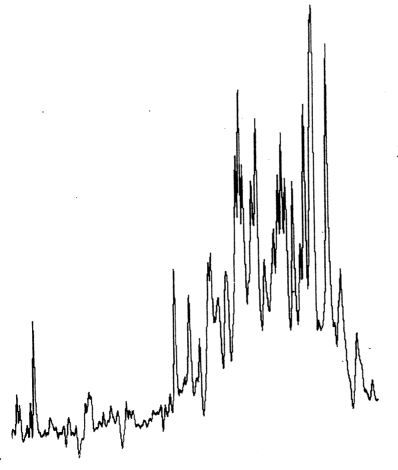


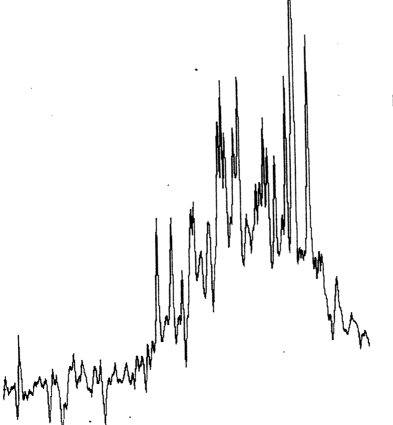
FIGURE 6d

MASS FRAGMENTOGRAMS STERANES m/e 217

6407/1-2



4285m E17730 Dust



4510m E17758 Dust FIGURE 6e

MASS FRAGMENTOGRAMS
STERANES m/e 217

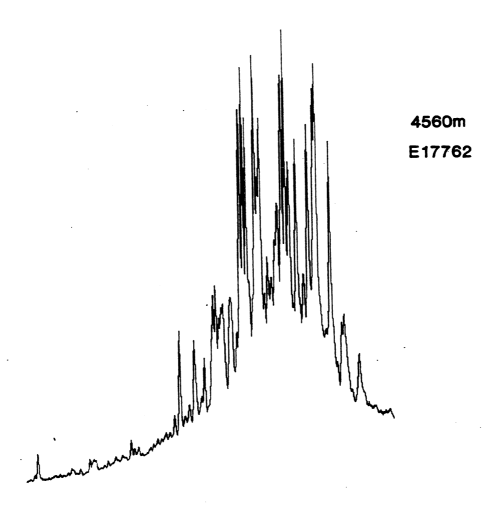
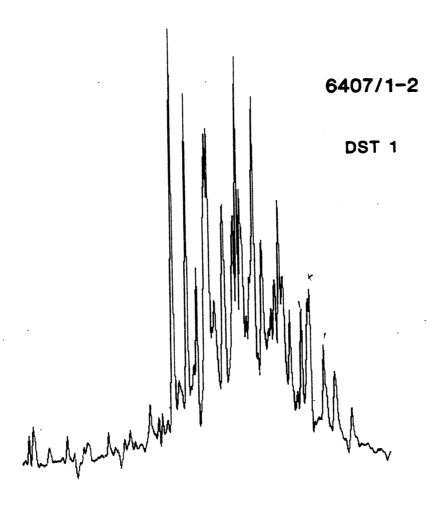


FIGURE 6i

MASS FRAGMENTOGRAMS
STERANES m/e 217
CONDENSATES



MASS FRAGMENTOGRAMS 7a FIGURE STERANES m/e 218 6407/1-2

3551.5m E17199

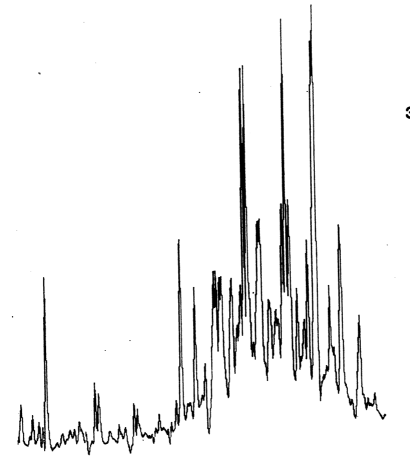
3595m

E17147

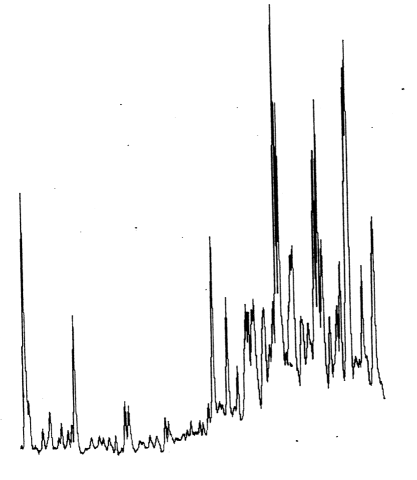
FIGURE 7b

MASS FRAGMENTOGRAMS STERANES m/e 218

6407/1-2



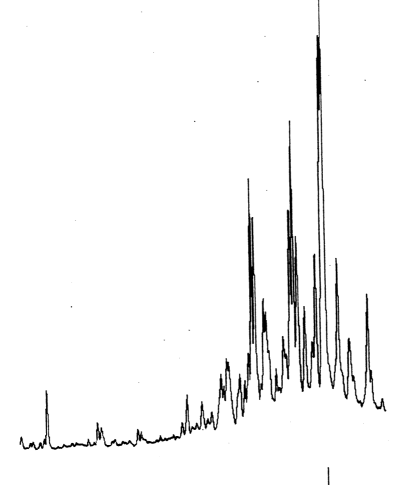
3661.4m E17804



3688.16m E17805 FIGURE 7c

MASS FRAGMENTOGRAMS STERANES m/e 218

6407/1-2



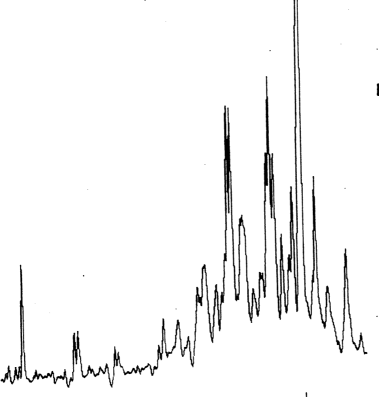
4210m E17725

4285m E17730 DCS FIGURE 7d

MASS FRAGMENTOGRAMS

TRIAROMATIC STERANES m/e 231

6407/1-2



4285m E17730 Dust



FIGURE 7e

MASS FRAGMENTOGRAMS

STERANES m/e 218

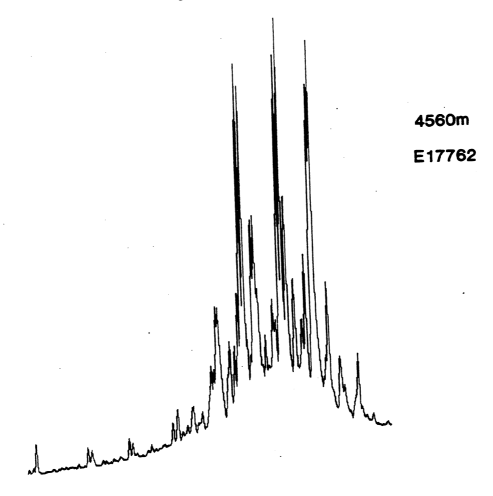


FIGURE 7i

MASS FRAGMENTOGRAMS STERANES m/e 218 CONDENSATES

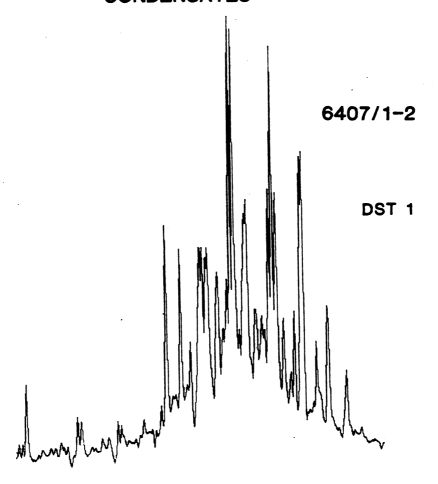
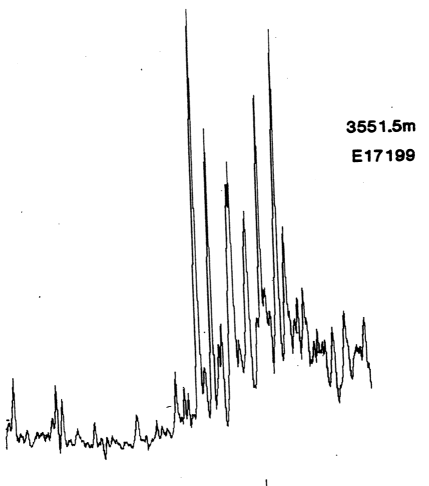


FIGURE 8a

MASS FRAGMENTOGRAMS

REARRANGED STERANES m/e 259

6407/1-2



3595m

E17147

FIGURE 8b

MASS FRAGMENTOGRAMS
REARRANGED STERANES m/e 259

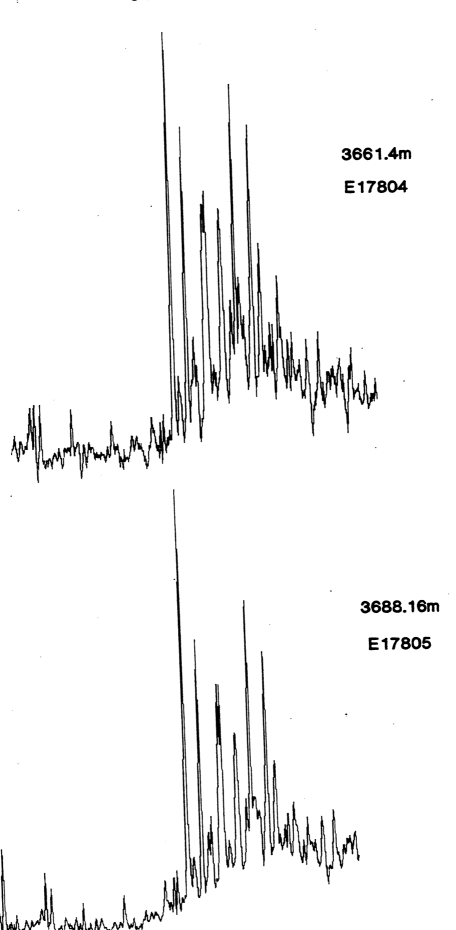
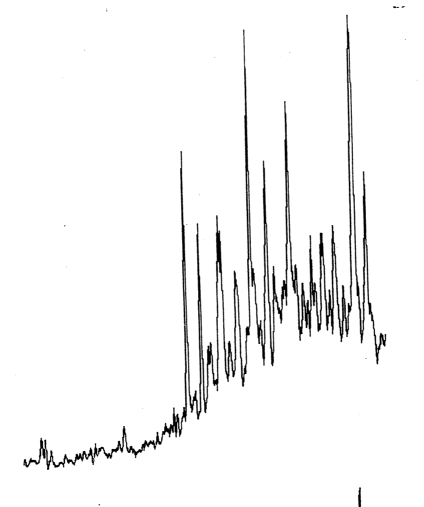


FIGURE 8c

MASS FRAGMENTOGRAMS

REARRANGED STERANES m/e 259

6407/1-2



4210m

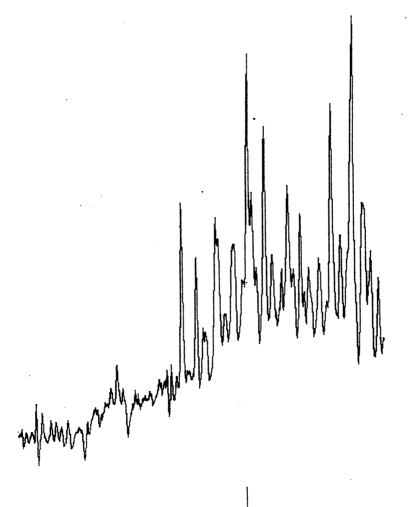
E17725

4285m E17730 DCS FIGURE 8d

MASS FRAGMENTOGRAMS

REARRANGED STERANES m/e 259

6407/1-2



Mind by the state of the state

4285m E17730 Dust

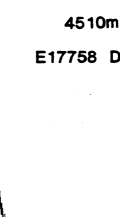


FIGURE 8e

MASS FRAGMENTOGRAMS
REARRANGED STERANES m/e 259
6407/1-2

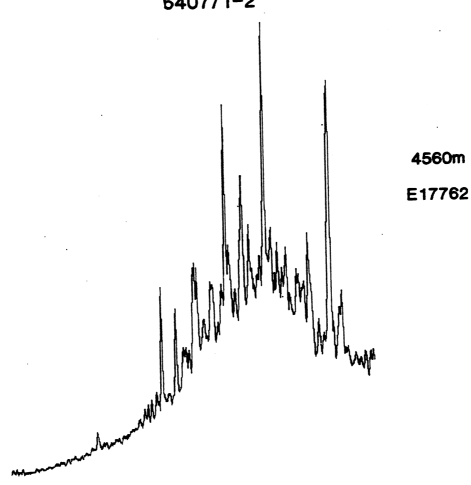


FIGURE 8i

MASS FRAGMENTOGRAMS REARRANGED STERANES m/e 259 CONDENSATES

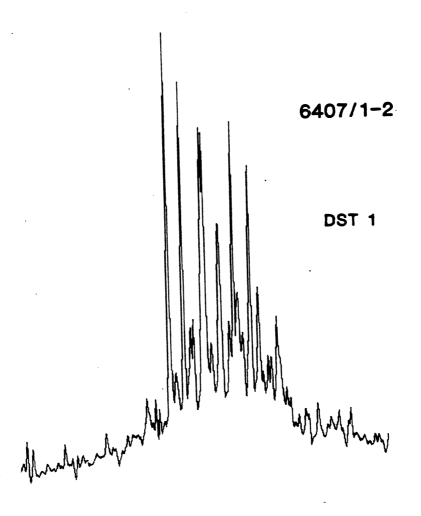


FIGURE 9a

MASS FRAGMENTOGRAMS

TRIAROMATIC STERANES m/e 231

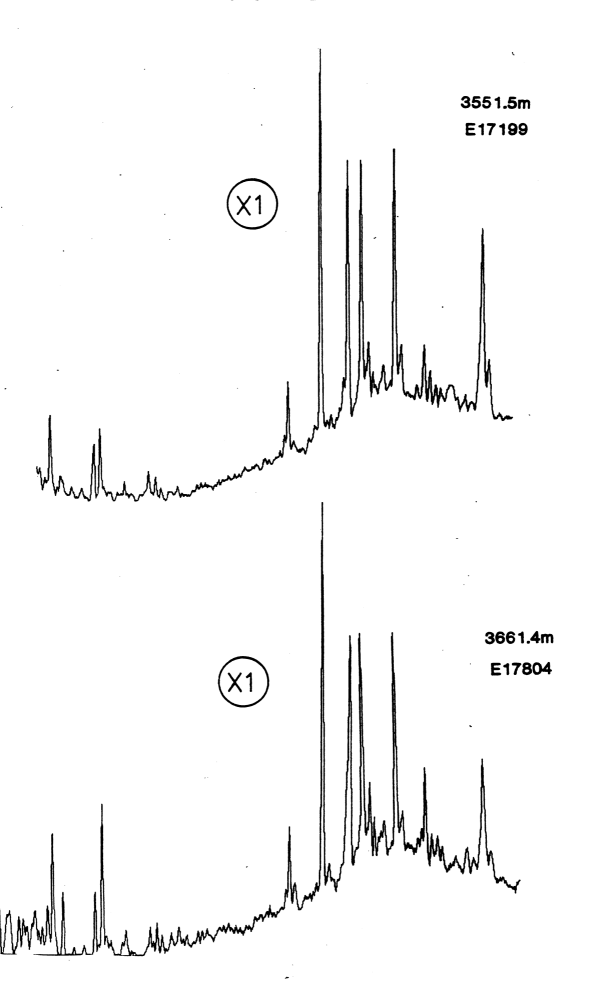
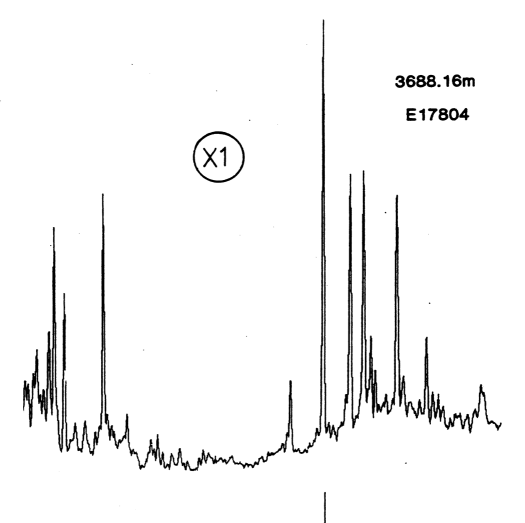


FIGURE 9b

MASS FRAGMENTOGRAMS

TRIAROMATIC STERANES m/e 231 6407/1-2



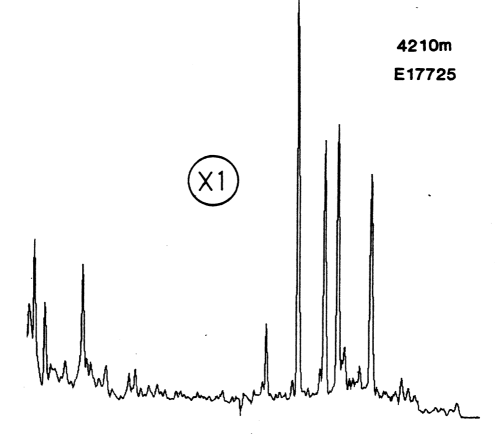


FIGURE 9c

MASS FRAGMENTOGRAMS

TRIAROMATIC STERANES m/e 231

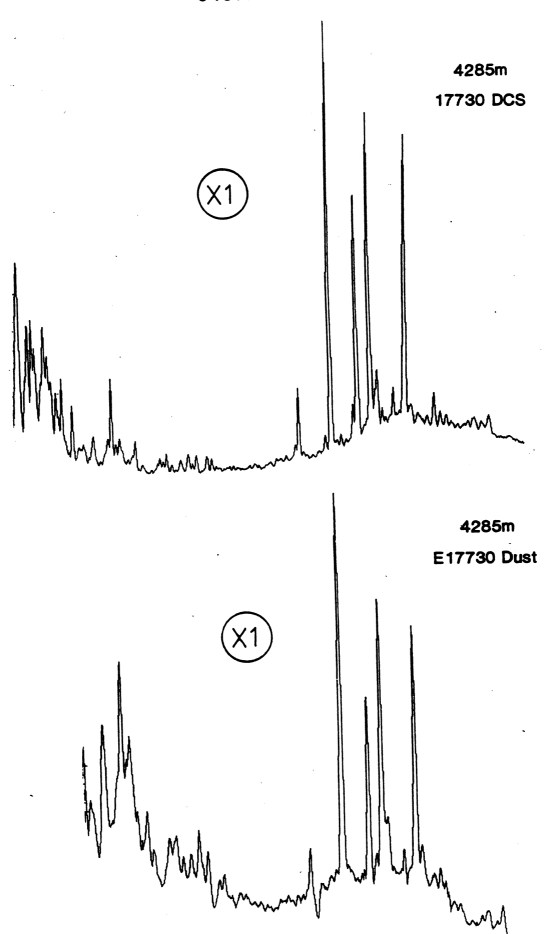


FIGURE 9d

MASS FRAGMENTOGRAMS

TRIAROMATIC STERANES m/e 231

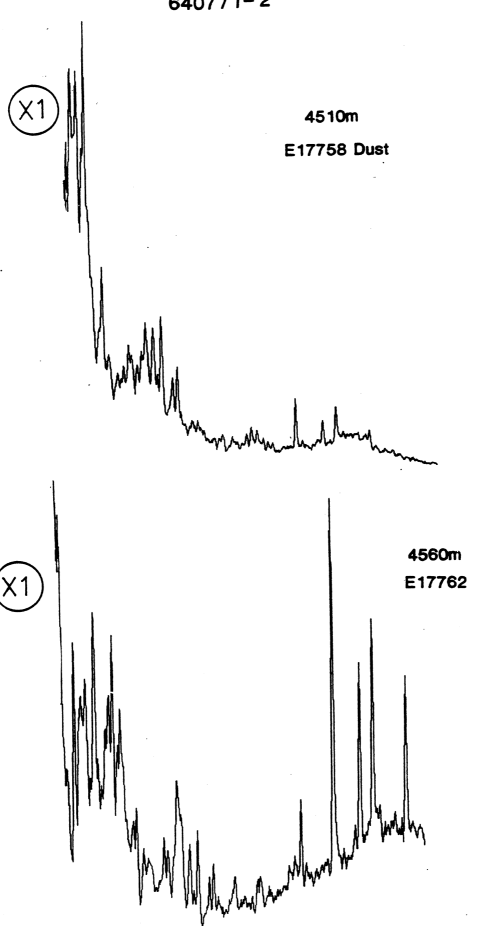
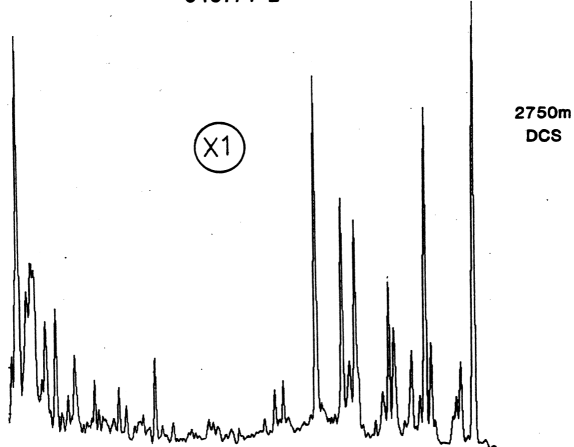


FIGURE 9f

MASS FRAGMENTOGRAMS

TRIAROMATIC STERANES m/e 231





IGURE 9h

MASS FRAGMENTOGRAMS

TRIAROMATIC STERANES m/e 231

CONDENSATES

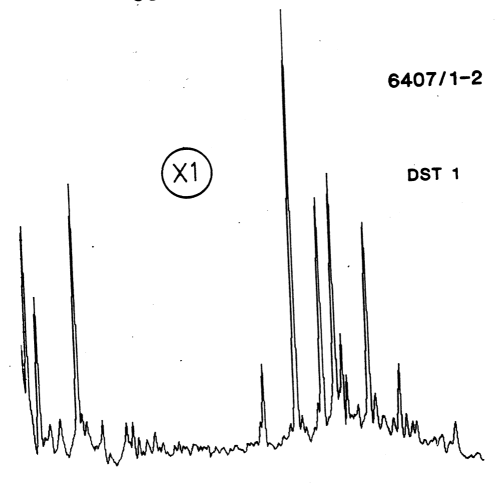
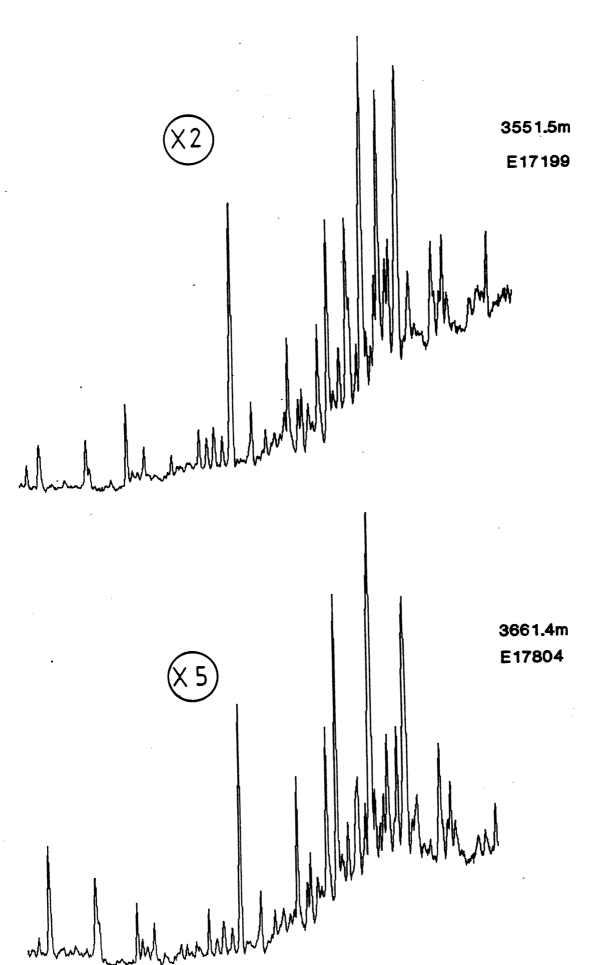


FIGURE 10a

MASS FRAGMENTOGRAMS

MONOAROMATIC STERANES m/e 253

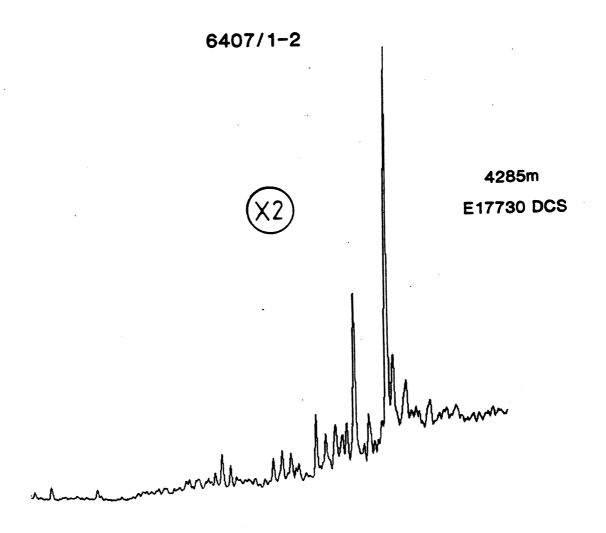


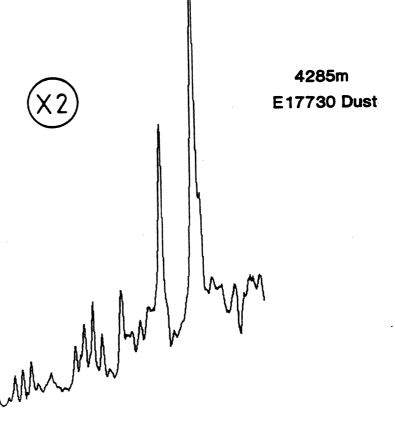
MASS FRAGMENTOGRAMS 10b FIGURE MONOAROMATIC STERANES m/e 253 6407/1-2

E17805 4210m

MASS FRAGMENTOGRAMS

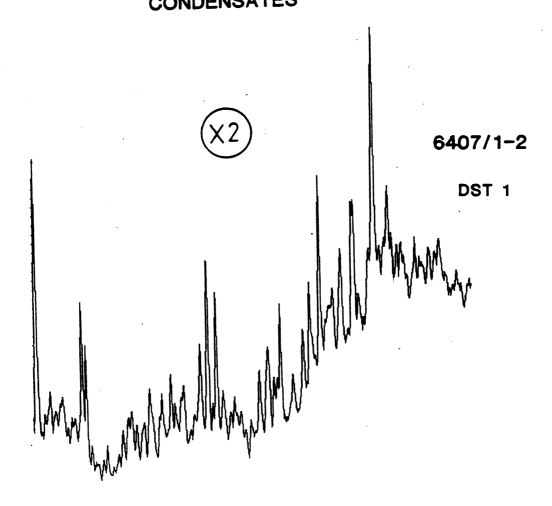
MONOAROMATIC STERANES m/e 253





MASS FRAGMENTOGRAMS 10d FIGURE MONOAROMATIC STERANES m/e 253 6407/1-2 4510m 4560m E17762 FIGURE 10h

MASS FRAGMENTOGRAMS MONOAROMATIC STERANES m/e 253 CONDENSATES



BRIEF DESCRIPTION OF THE ANALYSES PERFORMED BY GEOCHEM

"Screen Analyses" are described in sections A, C and D, "Sample Preparation" in section B, "Follow-up Analyses" in sections E through K and "Correlation Studies" in section L. The analyses can be run on either core or cuttings material with the proviso that samples must be canned for the C_1 - C_7 analysis and should be canned (or at least wet) for the C_4 - C_7 analysis. The other analyses can be run on both canned and bagged samples.

A) C1-C7 LIGHT HYDROCARBON ANALYSIS

The abundance and composition of the C_1 - C_7 hydrocarbons in sediments reflects their source richness, maturity and the character of the hydrocarbons they can yield. Most importantly, it is extremely sensitive to the presence of migrated hydrocarbons and is an excellent method for their detection. As it provides the information on most of the critical parameters and is also economical, this analysis is excellent for screening samples to decide which of them merit further analysis.

During the time which elapses between the collection of the sample at the wellsite and its analysis in the laboratory, a fraction of the total gas passes from the rock to the air space at the top of the can. For this reason, both the air space and the cuttings are analysed.

The analysis involves the gas chromatographic separation of the individual C_1 - C_4 gaseous hydrocarbons (methane, ethane, propane, isobutane and normal butane) and a partial resolution of the C_5 - C_7 gasoline-range hydrocarbons (for their complete resolution see Section E). The ppm abundance of the five gases and of the total C_5 - C_7 hydrocarbons are calculated from their electronically integrated peak areas (not from peak height) by comparison with a standard.

In the report, the following data are tabulated: the abundance and composition of the air space gas, of the cuttings gas and of the combined air space and cuttings gases. The combined results are also presented graphically.

B) SAMPLE WASHING AND HAND PICKING

All of the analyses described in subsequent sections are run on washed and hand picked samples.

Cuttings are washed to remove the drilling mud, care being taken not to remove soft clays and fine sand during the washing procedure. Using the C_1 - C_7 hydrocarbon data profile of the well, or the organic carbon profile (if this analysis is used for screening), electric logs (if supplied) and the appearance of the cuttings under the binocular microscope, samples are selected to represent the lithological and geochemical zones penetrated by the well. These samples are then carefully hand picked and the lithology of the uncaved material is described. It is these samples which are submitted for further analysis.

Sample material remaining after analysis is retained for six months. Unless instructions are received to the contrary, Geochem Laboratories may then destroy the samples.

Our reports incorporate a gross lithological description of <u>all</u> the samples which have been analysed and litho percentage logs. As screen analyses are recommended at narrow intervals, a complete lithological profile is obtained.

C) ORGANIC CARBON ANALYSIS

The organic carbon content of a rock is a measure of its total organic richness. Combined with the visual kerogen, C_1 - C_7 , C_4 - C_7 , pyrolysis and C_{15+} analyses, the organic carbon content is used to evaluate the potential (not necessarily actual) hydrocarbon source richness of the sediment. This analysis is an integral part of a total evaluation and it can also be used as an economical screen analysis for dry samples (when the C_1 - C_7 analysis cannot be used).

Hand picked samples are dried, crushed and then acidised to remove the inorganic calcium and magnesium carbonates. The actual analysis involves combustion in a Leco carbon analyser. Blanks, standards and duplicates are run routinely for purposes of quality control at no extra cost to the client.

The data are tabulated and presented diagramatically in our reports in a manner which facilitates comparison with the gross lithology (see Section B) of the samples.

D) MINI-PYROLYSIS

An ideal screen analysis which provides a definitive measure of potential source richness upon those samples whose organic carbon contents suggest fair or good source potential. This is described in detail in section K.

E) DETAILED C4-C7 HYDROCARBON ANALYSIS

The abundance and composition of the C_4-C_7 gasoline-range hydrocarbons in sediments reflects their source quality, level of thermal maturation and organic facies. In addition, the data also reveal the presence of migrated hydrocarbons and can be used for crude oil-parent source rock correlation studies.

This powerful analysis, performed upon hand picked lithologies, is employed as a follow-up to confirm the potential of samples which have been selected using the initial screen analysis. It is used in conjunction with the organic carbon, visual kerogen and C_{15+} analyses.

The individual normal paraffins, isoparaffins, naphthenes and aromatics with between four and seven carbon atoms in the molecule (but also including toluene) are resolved by capillary gas chromatography and their peak areas electronically integrated.

Normalised compositions, selected ratios and the ppm abundance of the total gasoline-range fraction are tabulated in the report and also presented graphically.

F) KEROGEN TYPE AND MATURATION

Kerogen is the insoluble organic matter in rocks. Visual examination of the kerogen gives a direct measure of thermal maturity and of the composition of the organic matter (organic facies) and indicates the source quality of the sediment – which is confirmed using the organic carbon, light hydrocarbon, pyrolysis and C_{15+} analyses.

The type of hydrocarbon (oil or gas) generated by a source rock is a function of the types and level of thermal maturation of the organic matter which are present. Both of these parameters are measured $\underline{\text{directly}}$ by this method.

Kerogen is separated from the inorganic rock matrix by acid digestion and flotation methods which avoid oxidation of the organic matter. It is then mounted on a glass slide and examined at high and low magnifications with a Leitz microscope. Chemical methods measure the total kerogen population but, with this technique, individual particles can be selected for examination and spurious material identified. This is particularly valuable in reworked, contaminated and turbodrilled sediments.

The following data are generated: the types of organic matter present and their relative abundances, an estimate of the proportion of reworked material, preservation state, the thermal maturity of the non-reworked organic matter using the spore colouration technique.

Our maturation scale has been developed to digitise small but recognisable changes in organic matter colouration resulting from increasing maturity and to place particular emphasis upon the immature to mature transition. In the absence of a universal colouration scale, the most significant points on our scale have been calibrated against equivalent vitrinite reflectance values. The following maturation stages are recognised at the low end of the scale:-

- a) immature; thermal index less than 2- (0.45% Ro)
- b) marginally mature; indices between 2- and 2. Minor hydrocarbon generation from amorphous and herbaceous ([±] algal) organic matter
- c) mature; indices between 2 (0.53% Ro) and 2 to 2+ (0.72% Ro), significant generation from amorphous, algal and herbaceous organic matter but wood only marginally mature
- organic matter but wood only marginally mature
 d) oil window; indices of 2 to 2+ (0.72% Ro) through to
 3 (1.2% Ro). Peak hydrocarbon generation.

The condensate zone starts at a thermal index of 3 whilst indices of $3+(2.0\ Ro)$ and higher indicate the eometamorphic dry gas stage.

A total of fourteen types of organic matter are sought based upon the major categories of algal, amorphous, herbaceous (spore, pollen, cuticle), wood, inertinite and resin. This detail is essential for a proper understanding of hydrocarbon source potential as the different sub-groups within each category have different properties.

Upon completion of the study, the kerogen slides are sent to the client.

G) VITRINITE REFLECTANCE

Vitrinite reflectance is an alternative/confirmatory method for evaluating thermal maturation which is used in conjection with the <u>visual kerogen</u> analysis. The reflectivity of vitrinite macerals increases in response to thermal alteration and is used to define maturation levels and, by projection, to predict maturity at depth or the thicknesses of section removed by erosion.

Measurements are made upon kerogen separations in conjunction with polished whole rock samples. In general, this analysis is performed upon the same samples as the visual kerogen analysis, thus facilitating a direct comparison of the two sets of results.

If possible, forty to fifty measurements are taken per sample - unless the sediments are organically lean, vitrinite is sparse or only a single uniform population is present. The data are plotted in a histogram which distinguishes the indigenous vitrinite from possible reworked or caved material. Averages are calculated for each population. Comments upon exinite fluorescence and upon the character of the phytoclasts are noted on the histograms. The reports contain the tabulated data, histograms and the reflectivities plotted against depth.

The vitrinite and visual kerogen techniques provide mutually complementary information upon maturity, organic matter type and diagenesis.

H) C15+ EXTRACTION, DEASPHALTENING AND CHROMATOGRAPHIC SEPARATION

Sections "A" and "E" dealt with analyses covering the light end of the hydrocarbon spectrum. This section is concerned with the solvent extractable organic material in the rock with more than fourteen carbon atoms in the molecule (i.e. the heavy end). The amount and composition of this extract indicates source richness and type, the level of thermal maturation and the possible presence of migrated hydrocarbons.

These results are integrated with those derived from the pyrolysis, visual kerogen, organic carbon and light hydrocarbon analyses.

The techniques involved in this analysis employ pure solvents and have been designed to give reproducible results. Hand picked samples are ground and then solvent extracted in a soxhlet apparatus, or by blending, with dichloromethane (the solvent system can be adapted to client's specifications). After asphaltene precipitation, the total extract is separated by column chromatography or high pressure liquid chromatography into the following fractions: paraffin-naphthene hydrocarbons, aromatic hydrocarbons, eluted NSO's (nitrogen-, sulphur-, and oxygen- containing non-hydrocarbons) and non-eluted NSO's. Note that the non-hydrocarbons are split into three fractions and not reported as a gross value. These fractions can be submitted for further analyses (carbon isotopes, gas chromatography, high mass spectroscopy) including correlation studies.

For convenience and thoroughness, the data are reported in three formats: the weights of the fractions, ppm abundances and normalised percentage compositions. The data are also presented diagramatically.

J) GC ANALYSIS OF C15+ PARAFFIN-NAPHTHENE HYDROCARBONS

The gas chromatographic configurations of the heavy C_{15+} paraffinnaphthene hydrocarbons reflect source type, the degree of thermal maturation and the presence and character of migrated hydrocarbons or contamination.

Not only is this analysis an integral part of any source rock study but it also provides a fingerprint for correlation purposes and helps to define the geochemical/palynological environmental character of the source rocks from which crude oils were derived.

The paraffin-naphthene hydrocarbons obtained by column chromatography are separated by high resolution capillary chromatography. Excellent resolution of the individual normal paraffins, isoprenoids and significant individual isoparaffins and naphthenes is achieved. Runs are normally terminated at nC35. A powerful in-house microprocessor system is being introduced to correct for the change in response factor with chain length.

The normal paraffin carbon preference indices (C.P.I.) indicate if odd (values in excess of 1) or even (values less than 1) normal paraffins are dominant. Strong odd preferences (* strong pristane peaks) are characteristic of immature land plant organic matter whilst even preferences (* strong phytane peaks) suggest a reducing environment of deposition. With increasing maturity, values approach 1.0 and oils are typically close to 1.0. The indices are calculated using the following formulae:

Chromatograms are reproduced in the report for use as visual fingerprints and in addition, the following data are tabulated: normalised normal paraffin distributions; proportions of paraffins, isoprenoids and naphthenes in the total paraffin-naphthene fraction; C.P.I $_{\rm A}$ and C.P.I $_{\rm B}$; pristane to phytane ratio; pristane to nC17 ratio.

K) PYROLYSIS

The process of thermal maturation can be simulated in the laboratory by pyrolysis, which involves heating the sample under specified conditions and measuring the oil-like material which is freed/generated from the rock. With this analysis, the potential richness of immature sediments can be determined and, by coupling the pyrolysis unit to a gas chromatograph, the liberated material can be characterised. These results are correlated with those obtained from the organic carbon, kerogen and C_{15+} analyses.

Small amounts of powdered sample are heated in helium to release the thermal bitumen (up to 340°C) and pyrolysate (340-550°C). The thermal bitumen correlates with the solvent extractable material (see above) whilst the pyrolysate fraction does not exist in a "free" state but is generated from the kerogen, thus simulating maturation in the subsurface. Abundances (weight ppm of rock) are measured with a flame ionisation detector against a standard. Thermal bitumen includes source indigenous, contaminant and migrated hydrocarbons but the pyrolysate abundance is a measure of ultimate source richness. The capillary gas chromatogram of the pyrolysate is used to evaluate the character of the parent organic matter and whether it is oil or gas prone. Peak temperature(s) of pyrolysate evolution is recorded. Carbon dioxide can be measured if requested but is normally ignored as the separation of the organic and inorganic species has been found to be artificial and unreliable.

Pyrolysate yields provide a definitive measure of potential source richness which avoids the ambiguities of the organic carbon data and the problem of contamination. This analysis is also used to evaluate the quality and character of the organic matter and the degree to which it has realised its ultimate hydrocarbon potential. Geochem does not employ the pyrolysis technique to evaluate maturation, preferring the kerogen and vitrinite reflectance analyses which avoid the problem of reworking and hence, are more reliable.

Capillary chromatograms produced for the pyrolysate hydrocarbons range from C_1 (methane) out towards C_{35} but exhibit considerable variations. They are used to define whether a source rock will yield oil, condensate or gas. With this new technique, it is now possible to complete the evaluation of a source rock.

The data are tabulated and presented graphically. MINI-PYROLYSIS includes ppm thermal bitumen and ppm pyrolsate. PYROLYSIS also provides the above together with the temperature of peak pyrolysate evolution. The capillary chromatograms of the pyrolysate obtained by PYROLYSIS-GC are reproduced in the report. The Mini-Pyrolysis analysis is recommended as a screening technique.

L) CORRELATION STUDY ANALYSES

Oil to oil and oil to parent source rock correlation studies require high resolution analytical techniques. This requirement is satisfied by some of the analyses discussed above but others have been selected specifically for correlation work. Many of these analyses also provide information upon the character of the environment of deposition of the parent source rocks.

- detailed C_4 - C_7 hydrocarbon (gasoline range) analysis. See Section E. Although these hydrocarbons can be affected by migrational/alteration processes, they commonly provide a very useful correlation parameter.
- capillary gas chromatography of the $C_{15}+$ paraffin-naphthenes. See section J. The branched normal paraffin distributions are used to "fingerprint" the samples.
- capillary chromatograms of whole oils and of the C_{8+} fraction of source rocks.
- capillary gas chromatography of C_{15+} aromatic hydrocarbons. Separate chromatograms of the hydrocarbons and of the sulphurbearing species are reproduced.
- high pressure liquid chromatograms.
- mass spectrometric carbon isotope analyses of crude oil and rock extract fractions and of kerogen separations. A powerful tool for comparing hydrocarbons and correlating hydrocarbons to organic matter. With this technique the problem of source rock contamination can be avoided. The data are recorded on x-y or Galimov plots.
- mass fragmentograms (mass chromatograms) of fragment ions characteristic of selected hydrocarbon groups such as the steranes and terpanes. The fragmentograms provide a convenient and simple means of presenting detailed mass spectrometric data and are used as a sophisticated fingerprinting technique. This provides the ultimate resolution for correlating hydrocarbons and facilitates the examination of hydrocarbon classes.
- vanadium and nickel contents.

Suites of (rather than single) analyses are employed in correlation studies, the actual selection depending upon the complexity of the problem. See also section N.

M) ANALYSES FOR SPECIAL CASES

M-1) ELEMENTAL KEROGEN ANALYSIS

This analysis evaluates source quality, whether the sediments are oil or gas prone, the character of the organic matter and its level of thermal maturation. It is the chemical equivalent of the visual kerogen analysis. The pyrolysis analysis is generally preferred to this technique, both methods providing similar information.

M-2) SULPHUR ANALYSIS

The abundance of sulphur in source rocks and crude oils.

M-3) CARBONATE CONTENT

The mineral carbonate content of sediments is determined by acid treatment. These data are particularly useful when used in conjunction with organic carbon contents as a screening technique.

M-4) NORMAL PARAFFIN ANALYSIS

Following the removal of the branched paraffins and naphthenes from the total paraffin-naphthene fraction, a chromatogram of the normal paraffins is obtained. The resulting less complicated chromatogram facilitates the examination of normal paraffin distributions.

M-5) SOLID BITUMEN EVALUATION

Residual solid bitumen after crude oil is generated by three prime processes: the action of waters, gas deasphalting, thermal alteration. Thus it provides a means of determing the reservoir history of a crude and of evaluating whether adjacent traps will or will not be prospective for oil. In carbonate sections, where organic matter is sometimes sparse, this technique is also used to evaluate thermal maturation levels.

The analysis involves the determination of the solubility (in CS_2) of the solid bitumen and of the atomic hydrogen to carbon ratio of the insoluble fraction.

N) CRUDE OIL ANALYSIS

N-1) API GRAVITY

This can be performed upon large (hydrometer) and small (SG bottle, pycnometer) samples and even upon stains extracted from sediments (refractive index).

- N-2) SULPHUR CONTENTS (ASTM E30-47)
- N-3) POUR POINT (ASTM D97-66, IP15/67)
- N-4) VISCOSITY (ASTM D445-72, IP71/75)
- N-5) FRACTIONAL DISTILLATION

Graph of cumulative distillation yield against temperature. Five percent cuts taken for further analysis. Mass spectrometric studies of these fractions provide a detailed picture of the distribution of paraffins and of the various naphthene and aromatic groups within a crude, which is useful both for correlation and for refinery evaluation purposes.