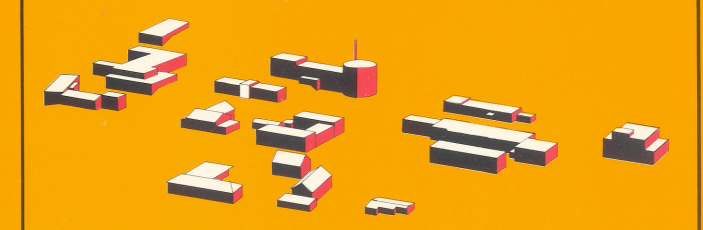
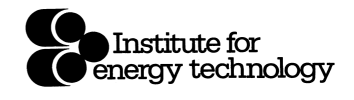
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REPORT ON STABLE ISOTOPES $(\delta^{13} \text{C AND } \delta \text{D})$ ON TEST GAS SAMPLES FROM 6406/3-1







Laur Garder

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APPROVED BY

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ANALYTICAL PROCEDURE

The samples were received in an aluminium bag (1156) and in a pressurized gas bottle (b 95). The natural gases were separated into the different gas components by a Carlo-Erba 4200 instrument. The hydrocarbon gas components were oxydized in separate Cu0-ovens, which enables us to collect several times when the concentration of a gas component is low. The combustion products CO $_2$ and H $_2$ 0 were frozen into collection vessels and separated. The isotopic measurements were performed on a Finnigan Mat 251 mass spectrometer. Our δ^{13} C-value on NBS-22 is -29.77 \pm .06 0 /oo.

RESULTS

The stable isotope results of the natural gases from well xx-1 are given in the following table.

Table 1. Isotopic composition of gas samples from well xx-1.

Sample	δ ¹³ C δ	5 D	C ₂	С ³	iC ₄	nC ₄	δ ¹³ C	0 ₂ δ ¹⁸ 0,220
1156 Al bag	-42.1			-30.3	-24.3	-29.1	-14.7	-12.2
b 95	-42.0	-231	-31.5	-31.1	-26.8	-30.0	-16.2	-16.0

Our uncertainty on the δ^{13} C-value is estimated to be \pm 0.3 0 /oo and includes all the different analysis steps. The uncertainty on the δD -value is estimated to be \pm 5 0 /oo.

The composition of the gas samples are given in table 2. The results have not been normalized to 100 %. The rest is air.

Table 2. Composition of gases from well xx-1. (%)

	1156 Al bag	b 95
c ₁	66.0	74.0
C ₂	. 8	2.1
C ₂ C ₃ iC ₄	. 4	. 4
iC ₄	.02	.02
nC ₄	.05	.05
nC ₄ CO ₂	10.0	13.0

INTERPRETATION

The δ^{13} C-values of methane, ethane, propane and n-butane for the two samples have been plotted on the maturity diagram by James (1983)* figure 1. A high maturity is indicated for both samples. Sample, 1156 Al-bag, however appear to be oil-associated, while sample, b 95, appear to have formed at somewhat higher maturity in the condensate field of the oil window. The same conclusion is derived when the samples are plotted in a δ^{13} C methane vs δ D methane cross plot (Schoell, 1983)** fig. 2. Sample, 1156 Al bag, appear to be oil associated while sample, b 95, appear to be condensate associated.

The relationship of the two samples from well xx-1 is not known to us. The more negative δ^{13} C Co $_2$ value of sample b 95 suggests, however, that this sample is collected closest to the source. The δ^{18} O CO $_2$ value of sample b 95 indicate a higher collection site temperature than sample 1156 Al-bag.

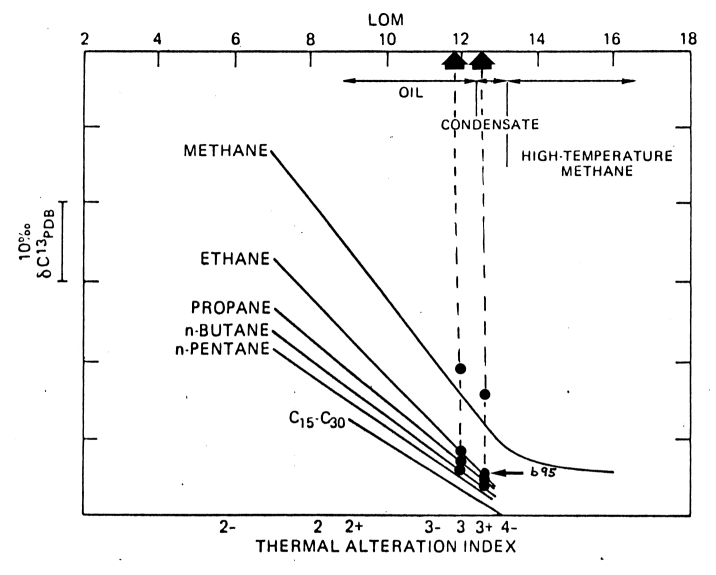
CONCLUSION

Both samples from well xx-1 appear to have formed at relatively high maturity in the oil window. Sample 1156 Al-bag appear to be oil associated while sample b 95 appear to be condensate associated.

- * James Alan T. (1983): Correlation of Natural Gas by Use of Carbon Isotopic Distribution Between Hydro carbon Components.

 A.A.B.G. Vol 67, No. 7. July 1983.
- ** Scholl, M. (1983): Genetic Characterization of Natural Gases.

 A.A.P.G. Vol 67, No. 12. Dec. 1983.



<u>Figure 1.</u> Carbon isotopic separations of the gas from well XX-1 is plotted on the maturity diagram (after James, 1983).

The calculated carbon isotopic separations between gas components are plotted on the vertical axis using a sliding scale that is simply the algebraic difference,in parts per mil, between the isotopic compositions of the natural gas components. The scale does not possess a fixed origin, but is oriented with the more depleted δ C values at the upper end. Use of this sliding scale allows the maturity of a gas to be assessed without prior knowledge of the isotopic composition of the gas' source.



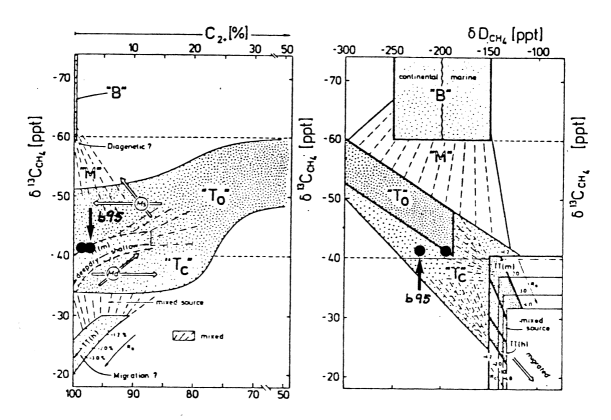


Figure 2 a. Variations of molecular composition in natural gases related to the isotope variations of methane.

<u>Figure 2 b</u>. Carbon and hydrogen isotope variations in methanes.

The principle for the genetic characterization of natural gases is that the primary gases (B-biogenic gas, T-associated gas, TT-non-associated gas) are defined by fields of compositional variations. These primary gases may become mixed and form various mixtures "M" of intermediate composition. "TT(m)" and "TT(h)" are non associated gases from marine source rocks and coal gases from N.W. Germany respectively, compositional shifts due to migration are indicated by arrows Md (deep migration) and Ms (shallow migration) respectively. "T" are gases associated with petroleum in an initial phase of formation. "T_f" are gases associated with condensates. (Schoell, 1983).