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THE UNIVERSITY OF ALBERTA

A FRACTAL ANALYSIS OF HETEROGENEITY

IN MISCIBLE DISPLACEMENT

by Daniel Joseph Giesbrecht

A THESIS

SUBMITTED TO THE FACULTY OF GRADUATE STUDIES AND RESEARCH IN PARTIAL FULFILMENT OF THE REQUIREMENTS FOR THE DEGREE OF MASTER OF SCIENCE

IN

PETROLEUM ENGINEERING

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Effect of Pore Structure on Miscible Displacement in Laboratory Cores

R.E. Bretz, SPE, New Mexico Inst. of Mining and Technology R.M. Specter,* Arizona State U. F.M. Orr Jr., SPE, Stanford U.

Summary. Observations of pore structure in thin-sections are related to the performance of stable, lirst-contact-miscible displacements in reservoir cores and then to simulations of displacement performance of CO₂ corefloods. Results of effluent composition measurements are reported for miscible displacements in seven core samples—three sandstones and four San Andres carbonates from west Texas or eastern New Mexico. Those displacements are interpreted by fitting the measured effluent compositions to the Coats-Smith (C-S) model, which represents the flow as occurring in flowing and stagnant fractions with mass transfer between them. Observations of thin-sections, including measurements of pore-size distributions and a simple measurement of spatial correlation of pore sizes, are also reported. Comparison of displacement results and thin-section data indicates that wide pore-size distributions and preferential flow paths are characterized in the C-S model by high dispersion coefficients and low flowing fractions. Simulations of the interactions of phase behavior and flow in nonuniform pore structures indicate that wide pore-size distributions and preferential flow paths can significantly increase residual oil saturations (ROS's) in CO₂ floods over those for uniform pore structures. Thus, heterogeneities observable at the scale of a thin-section have significant effects in laboratory core but much smaller effects in displacements at field scale. Large-scale heterogeneities present in field floods, however, probably cause similar increases in residual saturation in some fields.

introduction

Mixing between injected fluid and that present in a reservoir plays an important role in many EOR processes. In CO2 floods and other multiple-contact-miscible gas injection processes, for example, it is the transfer of components between phases that leads to high local displacement efficiency.¹⁻³ If the zone in which mixing takes place is confined to a narrow region, as is the case in slim-tube displacements, then the displacement is efficient³⁻⁵ as long as the pressure is high enough that CO₂ extracts hydrocarbons relatively efficiently from the oil.^{6.7} Comparison of numerical simulations of CO₂ floods^{3,5} with Helfferich's analytic solution f_{-} the interactions of phase behavior and flow in the absence of dispersive mixing⁴ indicates that when the transition zone is broad, as when the level of dispersion is high, displacement efficiency is reduced. Gardner and Ypma⁸ argued, on the basis of numerical simulations of the growth of a viscous finger, that mixing between CO_2 in a finger with oil in adjacent unswept regions also reduces local displacement efficiency. Dai and Orr⁹ used simulations of the effects of phase behavior on flow in a porous medium consisting of flowing and stagnant fractions to show that the broadening of the transition zone caused by the presence of the stagnant fraction has a similar effect. They used their model to interpret the CO2 coreflood experiments performed by Spence and Watkins, 10 who found that cores with a wide pore-size distribution showed higher ROS's after the CO₂ displacements. Thus, there is both experimental and theoretical evidence that mixing effects have a significant impact on CO₂ flood performance at the laboratory scale.

In this paper, we examine the influence of pore structure on the mixing that occurs during miscible displacements in laboratory cores. We present the results of San Andres carbonates. Of these samples, one sandstone and one carbonate are outcrop samples; the remainder are reservoir core samples. The displacements were performed with fluids at matched density and viscosity to eliminate the effects of gravity segregation and viscous instability and hence to isolate the effects of the pore space. We report the displacement results in terms of the parameters of the C-S model, ¹¹ which represents the pore space as flowing and stagnant fractions with mass transfer between them.

To characterize the geometry of the pore space, we present poresize distributions obtained from thin-sections from the same rock samples for which the displacements were performed. We argue

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SPB Reservoir Engineering, August 1988

that long transition zones, characterized in the C-S model by flowing fractions less than one, require not only a wide pore-size distribution but also that the pores be connected in such a manner that preferential flow paths are formed. We present a simple method for qualitative detection of the existence of preferential flow paths. The method is based on measurements of the mean size of pores neighboring randomly selected reference pores. Thus, we use the comparison of thin-section observations and coreflood results to argue that features of the pore structure observable at thin-section scale offer clues to the causes of miscible displacement behavior at coreflood scale.

To illustrate the influence of mixing on displacement performance of CO_2 corefloods, we report results of one-dimensional simulations made with the model developed by Dai and Orr.⁹ Results of those calculations confirm that pore structures that lead to preferential paths also produce lower displacement efficiency in laboratory corefloods. Finally, we comment on the problems associated with scaling laboratory coreflood results to field scale.

Displacement Apparatus and Procedure. Fig. 1 is a schematic of the apparatus used for displacement experiments. The fluids used and the apparatus varied slightly over the course of the experiments. In a typical experiment, the positive displacement pump pushes brine [1% NaNO3, 1% KNO3, 1% Ca(NO3)2, 0.1% NaN3, CaSO4 to saturation] through a sample valve, the core, and a differential refractometer and into a collection vessel placed on an electronic balance. Injection of a slug of miscible fluid is achieved by redirecting the flow by means of the sample valve through a sample loop of known volume filled previously with the same brine containing 0.4% sucrose as a tracer. The tracer concentration exiting the core is monitored continuously by the refractometer. Concentration and fluid weight data are acquired automatically with a microcomputer. The smallest available tubing, fittings, end caps, and refractometer cell were used to minimize dead volume in the system. Additional details of the apparatus and procedure may be found in Orr and Taber. 13

Least-Error Fits to the C-S Model. Effluent composition data were fit to the C-S readel, which has the form

$$\int \frac{\partial C}{\partial t_D} + \frac{\partial C}{\partial X_D} - \frac{1}{N_{\text{Pe}}} \frac{\partial^2 C}{\partial X_D^2} + (1-f) \frac{\partial C^*}{\partial t_D} = 0;$$

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$$(1-f)\frac{\partial C^*}{\partial t_D} = N_{Da}(C-C^*). \quad \dots \quad (1)$$

In that model, the PV is divided into flowing and stagnahl fractions. Thus, the pore space is characterized by three parameters: (1) the flowing fraction, f, the fraction of the pore space that contributes to flow; (2) the Peclet number, $N_{Pe} = vL/K_{d}$, the ratio of characteristic times for dispersion and flow; and (3) the Damköhler numer, $N_{Da} = K_m L/v$, the ratio of characteristic times for mass transfer and flow. Thus, the model represents contributions of convection and dispersion in the flowing fraction and mass transfer between the flowing and stagnant fractions. The composition of fluid at any point within the flowing and stagnant fractions is taken to be uniform, and the rate of mass transfer is assumed to be linear in the difference in concentration between the two fractions.

The appearance of calculated effluent composition curves depends strongly on all three parameters.¹³ For pulse injections of the type used here, reduction of f below 1 causes the injected fluid to appear at the outlet before 1 PV has been injected (see, for example, Figs. 4.16 through 4.18 of Ref. 13). Increasing the Damköhler number above zero causes a long tail on the effluent composition curve, and decreasing the Peclet number broadens the produced peak. If the flowing fraction is 1, the C-S model reduces to the convectiondispersion equation with its single parameter, the Peclet number. In that case, the 50% normalized concentration arrives at 1 PV injected (PVI). The C-S model is the simplest differential model that reflects typical observations of corefloods in which breakthrough occurs before 1 PVI and hence has been used extensively to interpret miscible displacements in laboratory cores. $10.11.13 \cdot 17$

The C-S model equations are first order in time and second order in space and hence require one initial condition and two boundary conditions for solution. For the experiments described here, the initial condition is

 $C(X_D, 0) = 0, \ 0 \le X_D \le 1.$ (2)

The inlet boundary condition for a pulse input experiment is

$$C(0,t_D) = C_0 - \frac{1}{N_{\text{Pe}}} \frac{\partial C}{\partial X_D} \bigg|_{X_D = 0}, \quad \dots \quad \dots \quad (3a)$$

 $C_0 = 1, \ 0 \le t_D \le \beta, \ldots$ (3b)

and

 $C_0 = 0, t_D > \beta.$ (3c)

The exit boundary condition is

 $C(X_D \rightarrow \infty, t_D) = 0$



$$C' = C(1, t_D) - \frac{1}{N_{\text{Pe}}} \frac{\partial C}{\partial X_D} \Big|_{X_D = 1.}$$

where C' is the exit concentration.

A discussion concerning the choice of boundary conditions may be found in Ref. 18.

The C-S model was solved numerically with a fully explicit finitedifference algorithm.¹³ The value of physical dispersion used was corrected to account for numerical dispersion caused by truncation errors. The least-error parameter determination was based on a form of Powell's¹⁹ method. The flowing fraction was adjusted first to minimize the sum of the absolute deviations between the calculated solution and data for the leading edge of the effluent pulse. Then, the Peclet number was varied to adjust calculated peak width. Finally, the Damköhler number was adjusted to match data for the tail portion of the peak. For the remainder of the fitting procedure, Powell's method was used to select and minimize along new search vectors until the best fit was achieved. The final result minimized the sum of the absolute deviations.

Thin-Section Analysis. Outcrop core plugs were cut paraliel to bedding planes. Reservoir core plugs were cut horizontally from vertical core samples, and where bedding planes were evident, the horizontal plugs were aligned with those planes. A commercial laboratory prepared epoxy-impregnated thin-sections, which were cut parallel to the axis of the plugs. Pore-size distributions were obtained by measuring the pare-body sizes of 300 randomly selected pores. Pore-body size was defined as the smallest-diameter circle that completely circumscribes the pore body. For irregular or ciongated pores, that definition exaggerates pore size for pores cut through their longest axis. That exaggeration is partly compensated for by the fact that the plane of the section does not coincide with the longest axis for many pores. In any case, that definition allows comparison between thin-sections, although it does not necessarily produce the same pore-size distribution as might be obtained by other techniques. For additional details of the procedures for measurement of pore sizes and size distributions, see the description given by Specter 20

Several investigators have argued that wide or multimodal poresize distributions cause the early breakthrough and tailing characMultiple-Contact Experiment," SPE? (April 1983) 272-80.

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A Review of Diffusion and Dispersion in Porous Media

T. K. PERKINS O. C. JOHNSTON MEMBERS AIME

THE ATLANTIC REFINING CO. DALLAS, TEX.

ABSTRACT

Because of the influence of dispersion on miscibledisplacement processes, diffusion and dispersion phenomena in parous rocks are of current interest in the oil industry. This paper reviews and summarizes a great deal of pertinent information from the literature.

Porous media (both unconsolidated packs and consolidated rocks) can be visualized as a network of flow chambers, having random size and flow conductivity, connected together by openings of smaller size. In such a porous medium, the apparent diffusion coefficient D is less than the molecular diffusion coefficient D₀, as measured in the absence of a porous medium. For packs of unconsolidated granular material the ratio D/D₀ is about 0.6 to 0.7. For all porous rocks, both cemented and unconsolidated, the ratio of diffusion coefficients can also be represented as D/D₀ = $\frac{1}{F\phi}$, where F is the formation electrical resistivity factor and ϕ is the porosity.

If fluids are flowing through the porous medium, dispersion may be greater than that due to diffusion alone. At moderate flow rates the porous medium will create a slightly asymmetrical mix zone (trailing edge stretched out), with the longitudinal dispersion coefficient approximately proportional to the first power of average fluid velocity (if composition is nearly equalized in pore spaces by diffusion). If the velocity in interstices is large enough, there will be insufficient time for diffusion to equalize concentration within pore spaces. In this region, longitudinal dispersion increases more rapidly than fluid velocity.

At low velocities in interstices, transverse dispersion is characterized by a region in which transverse di/fusion dominates. If the fluid velocity gets high enough, there will be a transition into a region where there is stream splitting with mass transfer but with insufficient residence time to completely damp-out concentration variations within pore spaces.

There are several variables that must be con-

trolled to get consistent longitudinal and transverse dispersion results, viz., (1) edge effect in packed tubes, (2) particle size distribution, (3) particle shape, (4) packing or permeability beterogeneities, (5) viscosity ratios, (6) gravity forces, (7) amount of turbulence, and (8) effect of an immobile phase.

INTRODUCTION

Diffusion and dispersion in porous rocks are of current interest to the oil industry. This interest arises because of the influence of dispersion on miscible-displacement processes.

In a recovery process utilizing a zone of miscible fluid, there is the possibility of losing miscibility by dissipating the miscible fluid or by channeling or "fingering" through the miscible zone. Diffusion and dispersion are two of the mechanisms that may lead to mixing and dissipation of the slug. On the other hand, dispersion may tend to damp-out viscous fingers which may be channeling through the miscible slug.⁵⁸ Hence, dispersion may be detrimental or beneficial (if it prevents fingering through the miscible zone). Therefore, it is doubly important that we understand these processes.

In this paper we review, summarize and interpret a gleat deal of information from the literature. In particular, we will briefly discuss molecular diffusion in miscible fluids. Then we will discuss what differences to expect for diffusion in a porous rock. If there is movement of the fluid through the rock, then there may be an additional mixing or "dispersion". Furthermore, the dispersion longitudinally (in the direction of gross fluid movement) and transverse to the direction of fluid movement will not be equal. We will discuss both types of dispersion as well as several variables which can affect dispersion (viscosity differences, density differences, turbulence, heterogeneity of media, etc.). This group of variables has sometimes led to difficulty when comparing literature data.

DIFFUSION OF MISCIBLE FLUIDS

If two miscible fluids see in contact, with an initially sharp interface, they will slowly diffuse into one another. As time passes, the sharp interface between the two fluids will become a diffuse

⁵⁸References given at end of paper.

Original manuscript received in Society of Petroleum Engineers office Sept. 10, 1962. Revised manuscript received Jan. 15, 1963.

A derivation of this equation has been shown oy Fowler and Brown,²⁴ Rifai,⁴⁴ and others.

In the actual case, molecular diffusion will cause mixing along the interface. The net result will be a mixed zone growing at a more rapid rate than would obtain from diffusion alone, but less than the rate predicted by Eq. 8. Taylor⁵² and Aris¹ have studied the case where the time necessary for appreciable concentration changes to appear, owing to convection transport, was long compared with the "time of decay" during which radial variations of concentration were reduced to a fraction of their initial value through the action of molecular diffusion. Theoretical equations derived by these investigators showed that, if one fluid were displaced by another fluid under these conditions where diffusion could nearly damp-out radial concentration variations, then a symmetrical longitudinal mixed zone would be established. The mixed zone would travel with the mean speed of the injected fluid and would be dispersed as if there were a constant dispersion coefficient given by Eq. 9.

$$K_{g} = D_{o} + \frac{U^{2} \sigma^{2}}{48 D_{o}} \cdot \cdots \cdot \cdots \cdot \cdots \cdot \cdots \cdot (9)$$

where $K_l =$ longitudinal dispersion coefficient, sq cm/sec,

U = average velocity, cm/sec, and

a = radius of the capillary, cm.

The effluent concentration, for the capillary tube, is given by the diffusion equation, Eq. 10.

$$C = \frac{1}{2} \left[1 \pm \frac{1}{\sqrt{K_{f}}/UL} \right] \left(\frac{1-V/V_{\rho}}{\sqrt{V/V_{\rho}}} \right] \dots (10)$$

Under what conditions will Eqs. 8 or 9 be valid? Taylor has shown that the ratio of time necessary for damping of radial concentration variations to the time to get a significant change in concentration is proportional to the dimensionless group DL

Furthermore, he has shown that diffusion effects



FIG. 4-LONGITUDINAL DISPERSION COEFFICIENTS FOR A RANDOM NETWORK OF CAPILLARIES (FROM SAFFMAN, REF. 45).

MARCH, 1963

are negligible, and therefore Eq. 8 is applicable, if $\frac{Ua^2}{D_oL} > 250.$

An intermediate region where neither Eq. 8 nor Eq. 9 is valid has also been studied by Bosworth⁸ and Var. Deemter, et al,56 but their studies did not result in a quantitative representation of behavior over the full intermediate range. Recently, Bailey and Gogarty⁴ have studied dispersion over the full range by solving the diffusion and flow equations numerically.

Dispersion in a Network of Capillary Tubes

Again we should note that a bundle of straight capillary tubes is not a very good representation of a porous rock. de Jong¹⁷ and Saffman⁴⁵⁻⁴⁷ have studied a random network of capillaries. Their mathematical model is a more accurate description of a porous rock than is a bundle of straight capillaries, but the results are still not quantitatively correct for rock. The analysis is valuable, however, from a conceptual point of view. Saffman's results can be sketched as shown on Fig. 4.

Dispersion in Packs of Granular Material

Let us now proceed to a discussion of longitudinal dispersion in packs of granular material. Most investigators studying this problem have filled a packed column with one fluid, displaced it with another fluid, and measured fluid composition at the exit end of the tube as a function of displacement. Brigham, et al,⁹ have shown a convenient method for determining the dispersion coefficient from data of this type. In a slight modification of their method, the function $\frac{V/V_p - 1}{\sqrt{V/V_p}}$ is plotted vs the per cent of displacing fluid (on arithmeticprobability paper) as shown on Fig. 5. The dispersion coefficient can then be calculated with the aid of Eq. 11.

$$\kappa_{g} = UL \quad \left(\frac{\lambda_{90} - \lambda_{10}}{3.625}\right)^{2} \cdots \cdots \cdots \cdots (11)$$





USE Ref. 9 as original source or this paper and include Ref 9 in 73 references

- a = radius of a capillary, cm
- C = concentration
- D = apparent diffusion coefficient in a porous medium based on the average area open for diffusion and the over-all length
- D_o = molecular diffusion coefficient, sq cm/sec
- d_p = particle diameter, cm
- d_T = tube diameter, cm
- E = longitudinal convective dispersion coefficient, sq cm/sec
- F = formation electrical resistivity factor
- G = quantity of material diffusing across a plane (see Eq. 1)
- g = acceleration of gravity
- K_l = total longitudinal dispersion coefficient, sq cm/sec
- K_t = total transverse dispersion coefficient, sq cm /sec
- k = permeability
- L =length of a tube or a porous block, cm
- Pe = Peclet number (see Eq. 23)
- S = slope of a log-normal particle size distribution (see Eq. 21)
- t = time, seconds
- U = average interstitial velocity, cm/sec
- V = volume injected, cc
- V_p = total volume of tube or total pore volume
- X = distance, cm
- x = volume fraction of injected fluid in the effluent (see Eq. 8)
- β_t = factor to account for wall effect (see Fig. 14)

$$\lambda = \frac{(V/V_p) - 1}{V/V_p}$$

- μ = fluid viscosity
- ν = turbulence weighting factor (see Fig. 23)
- $\rho =$ fluid density
- σ = a measure of the inhomogeneity of the porous pack
- $\phi = \text{porosity}$
- ψ = sphericity of a particle (see Fig. 17)

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where $\lambda_{90} = \frac{V/V_p - 1}{\sqrt{V/V_p}}$ when effluent contains 90

per cent displacing fluid.

Nearly all literature data show that the longitudinal dispersion coefficients for unconsolidated sand or bead packs can be represented as shown by Eq. 12.

where E = longitudinal convective dispersion coefficient, sq cm/sec, and

 d_{b} = particle diameter, cm.

Furthermore, in the region where both diffusion and convective dispersion are important, the total dispersion coefficient is the sum of these two coefficients. Hence, the total longitudinal dispersion coefficient in an unconsolidated sand pack can be represented as shown by Eq. 13, and is sketched on Fig. 6.

Can the dispersive behavior of granular packs be represented fairly accurately by an "equivalent" bundle of capillary tubes? A comparison of Eqs. 9 and 12 shows that in packed columns the longitudinal dispersion coefficient is proportional to the first power of average velocity (if longitudinal dispersion is large compared to longitudinal molecular diffusion), whereas for capillary tubes the dispersion coefficient is proportional to the second power of average velocity. How can we explain this significant difference in behavior?

A clue is furnished in work reported by Aris and Amundson.³ These investigators have studied the dispersion to be expected in mixing chambers of



FIG. 6-LONGITUDINAL DISPERSION COEFFICIENTS FOR UNCONSOLIDATED, RANDOM PACKS OF UNI-FORM SIZE SAND OR BEADS (DATA FROM REF. 11, 42, 53 AND 59).

A BEF this paper but also cite where authors obtained data to construct figure

uniform size connected in series. The chambers are filled with one fluid and a second fluid is injected at one end at a constant rate. The composition within each chamber is maintained uniform at all times by complete mixing (in the case of granular packs and for some flow conditions, the concentration in each pore space is maintained essentially uniform by diffusion). Eq. 14 gives the concentration of displaced fluid in each cell of a very long series of cells.

$$C_{i'j} = 1 - e^{-qt/\nu} \left[\sum_{n=1}^{n=n} \frac{(qt/\nu)^n}{n!} + 1 \right] \cdots (14)$$

where $C_n = \text{concentration of the displaced fluid in the nth cell,}$

q = injection rate,

t = time, and

v = volume of each cell.

Fig. 7 shows the concentration of displaced fluid calculated from Eq. 14 when a pore volume of 100 cells has been injected (i.e., $qt/\nu = 100$).

This figure shows that a mixed zone is established which moves with the mean speed of the injected fluid; that is, the 50 per cent point is essentially at the hundredth cell (if the displacement had been piston-like, there would have been an abrupt change in composition after the hundredth cell). Furthermore, the growth of the mix zone, relative to the 50 per cent composition point, can be represented by a constant dispersion coefficient (i.e., plots as an essentially straight line on probability paper). Fig. 7 is equivalent to Fig. 1. The apparent dispersion coefficient can be calculated with Eq. 3. By noting that the time to inject 100 cell volumes is equal to the length of 100 cells divided by the average fluid velocity, it follows that the apparent dispersion ccefficient varies as the first power of the mean velocity (the same behavior as exhibited by porous packs, see Eq. 12).

A pack of granular material, of course, can be thought of as a series of chambers or pore spaces connected by smaller openings. This is indicated graphically on Fig. 8, which shows the shapes of void spaces for various types of symmetrical packing of spheres as reported by Graton and Fraser.²⁷

Study of dispersion data from packed columns reveals that the concentration profile in the mixed zone is not typically a perfect S-shaped probability





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- a = radius of a capillary, cm
- C = concentration
- D = apparent diffusion coefficient in a porous medium based on the average area open for diffusion and the over-all length
- $D_o =$ molecular diffusion coefficient, sq cm/sec
- d_p = particle diameter, cm
- d_T = tube diameter, cm
- E = longitudinal convective dispersion coefficient, sq cm/sec
- F = formation electrical resistivity factor
- G = quantity of material diffusing across a plane (see Eq. 1)
- g =acceleration of gravity
- K₁ = total longitudinal dispersion coefficient, sq cm/sec
- K_t = total transverse dispersion coefficient, sq cm /sec
- k = permeability
- L =length of a tube or a porous block, cm
- Pe = Peclet number (see Eq. 23)
- S = slope of a log-normal particle size distribution (see Eq. 21)
- t = time, seconds
- U = average interstitial velocity, cm/sec
- V = volume injected, cc
- V_p = total volume of tube or total pore volume
- X = distance, cm
- β_t = factor to account for wall effect (see Fig. 14)

$$\lambda = \frac{(V/V_p) - 1}{V/V_p}$$

- $\mu \approx$ fluid viscosity
- ν = turbulence weighting factor (see Fig. 23)
- $\rho =$ fluid density
- σ = a measure of the inhomogeneity of the porous pack
- ϕ = porosity
- ψ = sphericity of a particle (see Fig. 17)

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DR. D.L. FLOCK (SUPERVISOR)

WHITING

AMBASTHA DR. A.K.

DR. J.D. SCOTT (EXTERNAL EXAMINER)

DATE: . June. 18. 1990....

ABSTRACT

The description of heterogeneity in porous media has always been limited by the impracticality of describing the property in question in complete deterministic detail. The result of this difficulty is that the spatial distribution of the property is often neglected. Recent work has indicated that fractal theory may provide an avenue for the estimation of spatial distribution over many scales of measurement. This knowledge would facilitate the extrapolation of laboratory measured dispersion values to dispersion on a field-wide scale.

The purpose of this investigation was to study the possibility of using the fractal dimension of permeability to describe heterogeneity for a variety of rock types. This study was undertaken by comparing the effluent concentration profiles of first contact miscible displacements in various rock types to the fractal dimensions calculated for permeability, porosity and mean pore throat size. The fractal dimension was calculated through the use of autocorrelation function and variogram techniques using a series of equally spaced permeability, porosity and mean pore throat size values obtained from each core tested.

The fractal dimension for permeability calculated using an autocorrelation function technique provided a valid means

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of estimating heterogeneity which compared favorably with other, more traditional heterogeneity indicators. The advantage in this approach is that the fractal dimension describes heterogeneity over all scales it is measured over and provides a means of extrapolating convective dispersion coefficients measured in the laboratory to field scale applications.

V

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Nomenclature

A	cross-sectional area cm ²
ANOVA	analysis of variance test.
a	range of semi-variogram (L)
b	scale multiplier.
С	relative solvent concentration.
C_D, C_D^*	dimensionless concentrations
	in mobile and stagnant fractions ,respectively.
$C_u(x, x+h)$	autocovariance function.
C(x, x+h)	autocorrelation function.
D	fractal dimension.
Dacf	fractal dimension calculated from
	autocorrelation functions.
D _{var}	fractal dimension calculated from
	variogram analysis.
D _a	Damkohler number = $\frac{\kappa L}{v}$
Dı	longitudinal dispersion coefficient (cm^2/s) .
D,D。	molecular diffusion coefficient (cm^2/s) .
đ	mean pore throat diameter (nM).
d _p	average particle diameter (cm).
Е	mathematical expectation operator.
erfc	complementary error function.
F	formation resistivity factor.

F	length of a line.
f	mobile fluid fraction.
G	quantity of solute diffused across a sharp interface (moles).
Н	fractal co-dimension = $\eta - D$
Н	Koval's heterogeneity factor.
h	lag distance.
K	mass transfer coefficient (cm^2/s) .
K.	effective dispersion coefficient (cm ² /s).
K,	coefficient of longitudinal dispersion (cm²/s).
K,	coefficient of transverse dispersion (cm²/s).
k	permeability (md).
Ke _{lab}	dispersivity = $\frac{\kappa_{\bullet}}{v}$.
L	length.
М	mobility ratio.
N	number of measurements.
N(h)	number of data pairs separated by lag value h.
n	number of observations.
P _c	capillary pressure (psi).
P.	Peclet number = $\frac{\sqrt{L}}{D_l}$.
P.V.	pore volume.

R.F. recovery factor.

r	correlation coefficient.
S	fraction of porespace invaded by mercury.
t _D	dimensionless time = $\frac{\nabla t}{L}$.
t	time (s).
V	pore volumes of fluid injected.
V _p	pore volume of the core.
V _{DP}	Dykstra-Parson's permeability variation.
X _D	dimensionless distance = $\frac{x}{l}$.
x	distance.
Z(x)	variable value at location x.
Z(x+h)	variable value at location x+h.
γ(h)	semi-variance.
η	dimension in euclidean space.
θ	contact angle (degrees).
λ	lambda function.
λι	lithology factor.
μ	viscosity (cp).
$\bar{\mu}_x$	mean value.
v,U	Darcian velocity (cm/s).
ξ	length of a measurement.
σ	heterogeneity coefficient.
σ²	variance.

.

I. Introduction

A major focus in petroleum engineering since its inception has been an attempt to maximize oil recovery in the most economical manner possible. Primary depletion, using natural reservoir energy as the driving force, will typically recover only a few percent of the original oil in place. Secondary recovery methods utilizing voidage replacement techniques have been successful in raising overall recovery to 50% in favorable cases. Despite these advances, a large percentage of the original oil in place remains unrecoverable due, in part, to the effects of interfacial tension and capillary forces which exist between phases within the porous media.

Tertiary recovery methods were developed in an attempt to increase recovery by minimizing or eliminating the effects of interfacial tension and capillary forces. Commonly, a displacing fluid which is completely miscible with the resident oil under reservoir conditions is injected into the formation. The choice of the solvent used is dictated by economic and operational considerations.

The efficiency of the miscible flooding process depends on two main groups of factors.

1

1.) Flood Instabilities

An unstable miscible front is detrimental to the efficiency of the displacement process. Factors important to the attainment of a stable front include the velocity of the front, the production rate of fluids from the wellbore as well as the effect of geological heterogeneities within the porous media.

2.) Fluid Mixing

Fluid mixing is an important factor in the maintenance of miscibility conditions. Factors such as molecular diffusion and convective dispersion have been quantified on a microscopic and macroscopic scale by many investigators. These phenomena affect the efficiency of the displacement process.

It has long been recognized that the severely unstable behavior of a miscible displacement front is a result of both microscopic and macroscopic heterogeneities within the porous media. A number of investigators have attempted to quantify the effect on a microscopic level by using either a Koval heterogeneity factor (H), or a Dykstra-Parsons V_{DS} variance factor. While these studies tend to be based upon porous media having a relatively homogeneous pore structure, little work has been directed toward quantifying the relationship between flood front instabilities and the more
heterogeneous pore structures which exist in carbonate rocks.

The primary objective of this study was to further investigate flood front mixing processes as influenced by heterogeneous pore systems. Any estimator of the degree of heterogeneity in a porous medium must account for the following factors (Warren and Price, 1961).

- The nature of the property variation (frequency distribution).
- 2. The spatial distribution of the property.
- 3. The inherent stability of the mechanism under study.

While many estimators of heterogeneity have been developed which account for factors one and three, the spatial distribution of properties as they relate to mixing processes is often ignored. The difficulty in quantifying the spatial correlation structure is that the properties of interest seem to exhibit different patterns of variability over different scales of measurement. Recently, many of these apparently random variations at different scales of measurement have been found to display characteristics of similarity across different scales of measurement (Gleick, 1987; Hewett, 1986). The concept of this apparent "order within chaos" is the basis for the emerging study of fractal theory and its application to processes in nature. In the current study, an examination was carried out of the distribution of porosity, permeability and pore throat sizes using mercury porosimetry techniques. Spatial correlation of these properties were examined using fractal methods and an attempt was made to provide an improved estimator of heterogeneity which incorporated a fractal component to account for spatial distribution. Specifically, the following topics were investigated:

- 1. Relationship between floodrate, convective dispersion (K_*) and recovery factor.
- 2. Effect of lithology on convective dispersion
 (K.) and recovery efficiency.
- 3. Effect of permeability, porosity and mean pore throat size variances on convective dispersion and recovery efficiency.
- Relationship between fractal dimension of permeability, porosity and mean pore throat size and dispersion and recovery behavior.
- 5. Comparison of V_{DP} ,Koval's H factor and a fractal estimator of heterogeneity (D_{acf}) with dispersion and recovery factor.
- 6. Scale dependent dispersivity and the fractal heterogeneity estimator D_{acl} .

II. Review of Diffusion and Dispersion

A. Introduction

Factors affecting the mixing of fluids at the flood front can have a profound effect upon the efficiency of the displacement process. In this context, mixing may be considered to be a microscopic process arising from velocity contrasts within the porous medium (convective dispersion) as well as the effects of molecular diffusion (Blackwell, 1962). Studies by Spence and Watkins (1980), among others, have found that there is an inverse relationship between the magnitude of convective dispersion and displacement efficiency. This finding has important implications not only for the oil industry, but in the area of groundwater hydrology as well. A large amount of research has been conducted in both fields aimed at quantifying the mechanisms associated with dispersion. Of particular interest has been the study of the effect of spatial arrangement of pore space as it relates to convective dispersion.

Figure II-1 illustrates a typical effluent concentration profile obtained in the current study for a first contact miscible displacement in Indiana limestone.



In this case, the mobility ratio was favorable and the porous medium relatively homogeneous. Typically, mixing between the solvent and the resident fluid results in the formation of a transition zone of finite thickness which separates 100% solvent from 100% resident fluid. Under conditions of favorable mobility ratio, eg. no viscous fingering, the thickness of this transition zone is the result of three main processes:

- 1. Molecular Diffusion
- 2. Convective Dispersion
- 3. Channeling

Convective dispersion as described by the effective dispersion coefficient K, used in this study is a function of these processes to varying degrees. It is possible to vary the parameters of the miscoble displacement in order to emphasize the effect of individual processes. This aspect will be discussed further in Chapter V.

Mixing caused by molecular diffusion in a porous medium is often represented by a modification of the Fick diffusion equation (Stalkup, 1983).

$$\frac{dG}{dt} = -DFA\phi \frac{\partial C}{\partial x}$$
(11-1)

Where: G = quantity of solute diffused across a sharp interface.

D =molecular diffusion coefficient.

- C = concentration.
- A = cross-sectional area.

F = formation resistivity factor.

- t = time.
- x = distance.
- $\phi = porosity.$

For miscible fluids flowing in a porous medium, more mixing occurs than can be accounted for by molecular diffusion alone. Convective dispersion and channeling play a major role in mixing processes at both the laboratory and field scale (Perkins and Johnston, 1963; Coats and Smith, 1964). Variations in the interstition velocity field at microscopic scales result in a combination of streamlines having markedly different solvent concentrations (Figure II-2).

Figure II-2 illustrates mixing of streamlines 1 and 2 in pore A. The equalized solvent concentration then proceeds to pore C where mixing with streamline 3 takes place. Convective dispersion may occur as longitudinal dispersion parallel to flow and/or as transverse dispersion normal to flow (Figure II-3).

This figure was removed due to the unavailability of copyright authorization. This figure showed a schematic diagram of convective dispersion on a pore scale in a in a porous medium. The original reference is fig. 3.3 on pg. 33 of the reference cited below.

Figure II-2 Convective Dispersion in Porous Media (from Stalkup, 1983) This figure was removed due to the unavailability of copyright authorization. This figure showed a schematic diagram of mixing in longitudinal and transverse directions in a porous medium. The original reference is fig. 3.2 on pg. 32 of the reference cited below.

Figure II-3 Mixing by Longitudinal and Transverse Dispersion in a Porous Medium. (from Stalkup, 1983)

Based upon a review of published data, Perkins and Johnston (1963) proposed empirical equations for longitudinal and transverse dispersion coefficients. These equations are based on miscible floods using fluids of equal density and viscosity:

$$K_{l} = D_{o} \left\{ \frac{1}{F\phi} + 0.50 \frac{v\sigma d_{P}}{D_{o}} \right\}, \qquad \frac{v\sigma d_{P}}{D_{o}} < 50 \qquad (11-2)$$

$$K_{i} = D_{o} \left(\frac{1}{F\phi} + 0.0157 \frac{v \sigma d_{p}}{D_{o}} \right), \qquad \frac{v \sigma d_{p}}{D_{o}} < 10^{4}$$
(11-3)

Where: $K_i = \text{coefficient of longitudinal dispersion}$.

- K_t = coefficient of transverse dispersion.
- v = average darcian velocity.
- $D_o =$ molecular diffusion coefficient.
- d_p = average particle diameter.

An examination of these two equations suggests several important results. At low fluid velocities the molecular diffusion term will tend to dominate and $K_i \cong K_i$. Under stable displacement conditions in the absence of gravity effects K_i will tend to dominate the mixing process with increasing fluid velocity. An examination of the dispersion terms of Equations II-2 and II-3 indicates that K_i will increase at a rate approximately 30 times that of K_i under these conditions.

Channeling of solvent through conduits of increased or decreased permeability is not addressed explicitly in the formulation of Equations' II-2 and II-3 but is implicitly included in most calculation methods used for determining K_1 and K_1 .



Brigham (1974) describes a simple method for determining K_i . The λ function method is shown in Figure II-4. The λ function is plotted vs the percent displacing fluid (on an arithmetic probability scale). The dispersion coefficient may then be calculated from Equation II-4:

$$K_{l} = vL \left\{ \frac{\lambda_{10} - \lambda_{10}}{3.625} \right\}^{2}$$
(11-4)
Where: $\lambda = \frac{\frac{v}{V_{p}} - 1}{\sqrt{\frac{v}{V_{p}}}}$
 $\frac{V}{V_{p}}$ = pore volumes injected.
 $v =$ darcian velocity.

L = length of the core.

 K_{l} = longitudinal dispersion coefficient.

Equation II-4 uses the width of the transition zone $(\lambda_{90} - \lambda_{10})$ as an indicator of K_i . The transition zone length will be significantly influenced by channeling or viscous fingering if it occurs. As a result, K_i will be a function of other factors in addition to the irregularity of the pore space under consideration. In general, as K_i increases, the recovery factor associated with the miscible flood will decrease. This result has been reported by several workers and occurs as a direct result of the adverse effect of heterogeneity upon the thickness of the transition zone (Koval, 1963; Spence and Watkins, 1980).

For the purposes of the current study, only the effects of longitudinal dispersion and channeling will be examined in detail.

A review of the literature reveals numerous attempts to model miscible mixing behavior through the use of mathematical models (Aronofsky and Heller, 1957; Coats and Smith, 1964). These models were developed by solving second order differential equations describing mass transport by dispersion within a porous medium. The following discussion will center on several models used to describe miscible flooding behavior.

B. <u>Convective</u> - <u>Dispersion Model</u>

The generalized form of the convective-dispersion (C-D) equation is given by Equation II-5 (Brigham, 1974).

$$K_{t}\frac{\partial^{2}C}{\partial x^{2}} - v\frac{\partial C}{\partial x} = \frac{\partial C}{\partial t}$$
(II-5)

Where: $K_i =$ longitudinal dispersion coefficient.

- x = distance from the inlet end of the core.
- v = darcian velocity.
- C = relative solvent concentration.

This equation may be solved by Laplace transform techniques under the appropriate boundary conditions. The choice of boundary conditions is not critical when the width of the transition zone is short compared with the length of the porous medium (Brigham, 1974). Under laboratory coreflood conditions the following boundary conditions and solution result:

Boundary Conditions: x = 0, C = 1 $x \to \infty$, $C \to 0$

$$C = \frac{1}{2} \operatorname{erfc}\left(\frac{x - vt}{2\sqrt{K_{i}t}}\right) + \frac{1}{2} \operatorname{e}^{\frac{vx}{K_{i}}} \operatorname{erfc}\left(\frac{x + vt}{2\sqrt{K_{i}t}}\right)$$
(11-6)

Where: C = relative solvent concentration at the outlet.

v = darcian velocity.

t = time.

 K_{i} = longitudinal dispersion coefficient.

erfc = complementary error function.

This equation may be history matched to effluent concentration profiles by setting x equal to L and expanding the second term of Equation II-6 using a Taylor series approach:

$$C = \frac{1}{2} \operatorname{erfc}\left(\frac{L - vt}{2\sqrt{K_{i}t}}\right) + \frac{1}{2\sqrt{\pi}} e^{-\left(\frac{L - vt}{2\sqrt{K_{i}t}}\right)^{2}} \cdot \left(\frac{2\sqrt{K_{i}t}}{x + vt} - \frac{1}{2}\left(\frac{2\sqrt{K_{i}t}}{x + vt}\right)^{3}\right)$$
(II-7)

Where: L = length of the core.

The derivation of this equation makes the following assumptions:

- 1. One dimensional flow.
- 2. First contact miscibility.

- 3. Dispersion is only significant parallel to flow.
- 4. Incompressible fluids.
- 5. Favorable mobility ratio.

The mobility ratio referred to in assumption five is simply the ratio of the viscosity of the resident oil divided by the viscosity of the displacing fluid. The preceeding conditions will be met in a properly designed laboratory coreflood. Brigham (1974) found that determination of K_i by the λ factor method (Equation II-4) is valid even under conditions where the boundary conditions applied to the convective-dispersion equation change eg. an infinite length boundary condition is not necessary. This implies that, for all practical purposes, only the first error function term of Equation II-7 is important when determining K_i . This reduces Equation II-7 to:

$$C = \frac{1}{2} erfc \left\{ \frac{L - vt}{2\sqrt{K_{i}t}} \right\}$$
(11-8)

This result is simply the solution to the C-D equation when an infinite length boundary condition is applied. The C-D model allows for the direct calculation of K_i as all of the other quantities will be available as parameters in a coreflood. The C-D equation described has the disadvantage of not accounting for dead-end pore volume in the porous medium and also does not address the problem of scale dependence in convective dispersion. In cases where a significant portion of the pore space consists of dead-end pores, a more sophisticated approach will be required (Baker, 1977).

C. Coats - Smith Model

A more sophisticated approach to the modelling of effluent concentration profiles was proposed by Coats and Smith in 1964. In this model, resident oil is assumed to be trapped in dead-end or occluded pore space. This dead-end pore space is only accessible to the flowing solvent at a single point with mass transfer occurring by diffusional processes only. The Coats-Smith model is a modification of the convective-dispersion (C-D) model in which terms accounting for a stagnant volume are added. All assumptions implicit in the development of the convective-dispersion model also apply to the Coats-Smith model. Equation II-9 illustrates the one-dimensional dimensionless form of their equation.

$$\frac{1}{P_{\bullet}} \frac{\partial^2 C_D}{\partial X_D^2} - \frac{\partial C_D}{\partial X_D} = f \frac{\partial C_D}{\partial t_D} + (1 - f) \frac{\partial C_D^*}{\partial t_D}$$
(11-9)
and: $(1 - f) \frac{\partial C_D}{\partial t_D} = D_a (C_D - C_D^*)$
Boundary conditions: $X_D \rightarrow 0.0$ $C_D \rightarrow C_D (t_D)$
 $X_D \rightarrow \infty$ $C_D \rightarrow 0.0$

Where: $P_{\bullet} = \text{Peclet number} = \frac{\nabla L}{D_{\downarrow}}$ D_a = Damkohler number = $\frac{KL}{v}$ f = mobile fraction. C_D, C_D^{\bullet} = dimensionless concentration in mobile and stagnant fractions, respectively. $t_p = \text{dimensionless time} = \frac{tv}{I}$ X_D = dimensionless distance = $\frac{x}{I}$ K = mass transfer coefficient. $D_l =$ longitudinal dispersion coefficient.

L = core length.

Equation II-9 is represented by three dimensionless parameters, the Damkohler number (D_a) . the mobile fraction (f) and the Peclet number (P_{\bullet}) which control the behavior of the Coats-Smith model and its ability to match effluent concentration profiles. As pointed out by Jasti et al (1987), D_a and P. are highly velocity dependent. For sufficiently small velocities, D_a will be large and mass transfer by diffusive processes will be essentially instantaneous (Coats and Smith, 1964). Under these conditions, Equation II-9 reduces to the simple C-D equation discussed earlier with $\frac{1}{P_1} \cong \frac{D_1}{L}$. In physical terms, this has the effect of reducing the relative contribution

of convective dispersion and describes a narrower transition zone in the effluent concentration profile. For cases where \vee is large, D_a becomes insignificant and Equation II-9 once again reduces to the C-D equation. In this case, convective dispersion is dominant over diffusion and $\frac{1}{P} \cong \frac{D_1}{N}$. Under these conditions, Perkins and Johnston (1963) show that D_i is approximately proportional to \vee so that $\frac{1}{P_1} \cong D_1$. The flowing fraction f appears to be relatively insensitive to variations in the Darcian velocity. Studies by Baker (1977) and Jasti et al (1987) suggest that f is a function of the porous addium only. The flowing fraction f has been defined as the break country pore volume observed in an experiment conducted at high fluid velocities. This method of defining f is conceptually identical to the phenomena of channeling implicitly accounted for in the longitudinal dispersion coefficient D_{i} which forms part of the convective-dispersion equation.

The prime advantage afforded by the Coats-Smith model is its ability to fit effluent concentration profiles which exhibit "tailing" of the concentration curve as a result of gradual mass transfer from dead-end pore space by diffusion processes. Figure II-5 shows that in the case of capacitance where dead-end pore space causes stagnant regions within the porous medium, tailing of the

concentration curve is much more pronounced and dispersion alone cannot account for it. This figure uses data from the current study. The curve depicting capacitance was from a miscible flood in Golden Spike limestone while the other curve is the result of a displacement at an equivalent rate in Berea sandstone.



Another advantage is the explicit treatment of the contribution of channeling through the use of the flowing or mobile fraction, f. The major disadvantage to the Coats-Smith model is that it is a multiple parameter model and, as such, encounters the problem of non-uniqueness of parameters. Typically, the parameters (D_a) , f and (P_a) are

chosen to provide the best fit to the observed effluent concentration profile from a laboratory coreflood. As Baker (1977) and Stalkup (1983) have pointed out; however, there are different combinations of model parameters which will yield an equally good fit to the experimental data. As a result of this, model parameters estimated solely through best-fit techniques may lack practical significance.

D. Other Related Work

Most related work in the area of miscible displacement modeling has centered on refinements to the original C-D and Coats-smith models. Brigham (1974) correctly recognized that the original analytical solutions obtained for these equations described *in-situ* solvent concentrations and were inappropriate for matching effluent concentration profiles. The use of the original solutions in modelling laboratory data resulted in serious material balance errors and new solutions for the original equations applicable to effluent concentrations were derived.

Baker (1977) provided a means of relating the effective dispersion coefficient, which governs field-scale floods, to the Coats-Smith parameters obtained from laboratory corefloods. Capacitance caused by dead-end pore space tended to increase the length of the transition zone above the length that would be expected from longitudinal

dispersion alone.

In general, neither the convective-dispersion model or the Coats-Smith model are able to account for the effect of viscous fingering. As a result, their use has been confined to miscible displacements with favorable mobility ratios. Vossoughi et al (1984) describe a method of modifying the C-D equation to account for the effect of viscous instabilities. This method was used with success in the modelling of polymer flow by introducing dead-end pore volume and polymer retention terms to the original differential equation.

Correa et al (1987) used simplified solutions to the Coats-Smith, porous sphere and transverse matrix diffusion models to interpret effluent concentration profiles from heterogeneous cores. Using simplified solutions in Laplace space, they developed a means of estimating a unique set of parameters (eg. (D_a) . f, and (P_*)) which apply to any of the three parameter models they studied.

III. Pore Structure Effects

A. Introduction

The most commonly measured properties of a porous medium are porosity and permeability. For the case of immiscible displacement, studies have indicated a positive correlation between porosity and recovery efficiency as well as a lack of correlation between permeability and recovery efficiency (Wardlaw, 1976). Although in common use, recent studies have indicated that porosity and permeability alone may not be good indicators of producibility (Jodry, 1972; Wardlaw, 1976; Bretz et al, 1988). Another approach to this question involved determining the relationship between pore size and recovery efficiency. Wardlaw (1976) found an inverse relationship between pore to throat size ratios and non-wetting phase displacement efficiency. Also recognized was a positive correlation between pore throat to pore coordination number and recovery efficiency. These studies were conducted under conditions of immiscible displacement and may not be valid under miscible conditions.

The relationship between pore structure and dispersion in miscible flooding has been investigated by several workers (Spence and Watkins, 1980; Bretz et al, 1988). A common approach to the problem has been to examine the

relationship between recovery efficiency and permeability distribution (Perkins and Johnston, 1963; Warren and Price, 1961; Warren and Skiba, 1964). This approach, while having the advantage of simplicity, does not address the difficulties inherent in the non-uniqueness of the permeability frequency distribution chosen. Figure III-1 illustrates this effect. Both sides of the figure have "similar frequency distributions" but "different spatial distributions" of permeability.



Bretz et al (1988) interpreted the effect of differing frequency and spatial distributions of pore size in terms of the Coats-Smith model for miscible displacement. While a broadening of the frequency distribution curve could be correlated with an increase in convective dispersion, the spatial correlation of the pore sizes also played a large role in the behavior of the displacement. This finding is not surprising in that spatial correlation of large and small pore sizes will often lead to systematic variations in permeability which can lead to channeling of the solvent and an increase in the apparent dispersion coefficient. Despite the fact that spatial correlation structure is of importance to fluid flow most measures of heterogeneity make the assumption that porosity and permeability are randomly distributed (McCaffery et al, 1978). Porosity and permeability development in nature occurs in response to geological processes which are "non-random" in both a lateral and vertical sense. Thus, the assumption of randomness of porosity and permeability on an areal scale may not be valid.

B. Geological Aspects

Studies of heterogeneity often assume that porosity and permeability are randomly distributed throughout the porous medium (Wyllie, 1962; Warren and Price, 1961). While this assumption may be reasonable on the scale of individual pores, it does not account for the influence of geological processes upon porosity and permeability distributions at

larger scales.

From an areal standpoint, the distribution of porosity and permeability have been shown to be related to the type of reservoir rock present in a given location (McCaffery, 1978; Blatt et al, 1980). Langton and Chin (1968), and McCulloch et al (1969) showed a distinct correlation between porosity and permeability and the occurrence of depositional environments in a carbonate reservoir at Rainbow Lake. The distribution of these depositional environments in space is predictable using current geological methods. In general, the lateral distribution of reservoir rocks is determined by the relationship between sediment supply and the energy level of the depositional environment. Under conditions of a marine transgression, where water depths are gradually increasing, the depositional environments shown in Figure III-2 will gradually prograde inland. The high-energy, shallow-water environments will tend to deposit more coarse-grained and well-sorted sediments which tend to form rocks having high porosity and permeability. Conversely, low-energy, deep-water environments result in the deposition of finer grained clays and mudstones with poor porosity and permeability characteristics.

This figure was removed due to the unavailability of copyright authorization. This figure showed the relationship between the vertical geological record and depositional environments. The original reference is fig 19-1 on pg. 620 of the reference cited below.

Figure III-2 Application of Walther's Law to Lateral Depositional Environments (from Blatt et al, 1980)

A fundamental principle in the geological sciences is Walther's law of succession of facies. In it's simplest form, it states that rocks which are deposited in a lateral sequence will also show the same sequence in cross-section (Figure III-2). This principle is valid provided that the depositional record is continuous, eg. subsequent erosion has not removed any of the rock layers. If the vertical succession of rock these is related to areal environments of deposition which on an non-random physical laws, then it is reasonable to assume that vertical rock sequences may also be non-random. Studies by Cant and Walker (1976) used Markov chain analysis to demonstrate spatial correlation of rock sequences in a vertical sense. Hence, porosity and permeability distributions are non-random if a sufficiently large scale of investigation is applied. Recent studies (Burrough, 1981; Hewett, 1986; Goggin, 1988) suggest that spatial correlation may also exist at smaller scales of measurement and that the behavior of spatial correlation may be fractal in nature. This possibility will be discussed in a later section.

The vast majority of oil bearing reservoirs are composed of either carbonates or sandstones. Both of these rock types develop porosity and permeability distributions characteristic of their environments of deposition, but through different processes. The fact that carbonate reservoirs are organic in nature tends to complicate studies of their patterns of heterogeneity. Modern analogs to ancient carbonate environments indicate that water depth, temperature and salinity are the controlling factors. Carbonate rocks exhibit a much more variable nature than do sandstones. Carbonates exhibit their greatest porosity and permeability immediately after deposition. Subsequent burial and compaction causes a sharp reduction in the original porosity and permeability in temperature and chemistry of meteoric waters. The particles which comprise limestones are primarily biological in origin and may contain a significant amount of intra-particle porosity and permeability. These rocks are; by definition, composed almost exclusively of calcium carbonate and aragonite. These minerals are highly soluble in meteoric waters leading to dissolution of the matrix in some locations and a reprecipitation of carbonate as pore-filling cements in other areas. In addition to dissolution and cementation of pore space, replacement of calcite is also a common occurrence. The circulation of Mg² -rich groundwaters through the carbonate matrix causes the recrystalization of calcite to form dolomite by the following reaction (Blatt *et al.*, 1980):

 $2CaCo_3 + Mg^{2*} \Leftrightarrow CaMg(CO_3)_2 + Ca^{2*}$

This reaction is significant in that dolomitization improves the producibility of the rock due to a transformation of the pore structure and an increase in porosity.

Diagenetic alteration is a complex process related to the geochemical stability of the matrix under local conditions of pressure, temperature and cation saturation

within circulating groundwaters. Interplay of these factors in carbonate rocks results in an extremely heterogeneous pore structure which may greatly increase dispersion and channeling under miscible displacement conditions. Chilinger (1972) states that the effect of diagenetic alteration of carbonates is such that quantitative measures of porosity and permeability may no longer offer a satisfactory indicator of the producibility of the rock. Under these conditions, a study of the pore to pore correlation structure is essential to a proper determination of producibility (Jodry, 1972; Wardlaw, 1976; Wardlaw and Cassem, 1978).

Sandstones are, in general, much more areally continuous than carbonates and display significantly different characteristics. Like carbonates, most sandstones experience a sharp reduction in porosity and permeability upon lithification. Unlike carbonates, however, sandstones are composed of dense, volcanically derived quartz and feldspar particles which contain little or no intra-particle porosity or permeability. As a result porosity and permeability is inter-granular in nature in these rocks. Post-depositional diagenetic alteration of sandstones by circulating meteoric waters has been found to produce only minor alteration of the pore structure (Blatt et al, 1980). Alteration of grain boundaries through pressure dissolution and mineral cementation will occur under conditions of high temperature and pressure however, the relative insolubility of quartz and feldspar serves to minimize this effect. Hence, the pore structure in sandstones is fairly homogeneous resulting in a fairly direct influence of porosity and permeability upon the producibility of the rock (Jodry, 1972).

IV. Methods of Spatial Correlation

The use of deterministic methods in the assessment of spatial correlation structure requires a complete description of the parameter of interest with respect to space. While this requirement is satisfied for properties which are homogeneous over a given sample space, it is difficult to describe heterogeneous systems in complete deterministic detail (Mishra, 1987). The intermittent nature of sampling in the current study thus rules out deterministic methods in the study of spatial correlation. As a result, a stochastic approach is required whereby the observed properties will be treated as random variables with a known mean and variance (Journel and Huijbregts, 1978).

The current study is concerned with defining an improved estimator for heterogeneity at the core level incorporing a component of spatial correlation. One such component is available through the use of fractal analysis. Fractal analysis is a mathematical technique which is useful in defining the "irregularity" or spatial heterogeneity of a particular property. In this study, the irregularities of permeability, porosity and pore throat size were studied using this technique. Determining spatial heterogeneity in this context requires a determination of the Haussdorff -Besicovitch (fractal) dimension of the data series. Recent

studies (Hewett, 1986; Goggin, 1988) indicate that large scale reservoir properties such as permeability distributions also display fractal characteristics. The remainder of this chapter outlines the basics of fractal theory and describes methods of application employed in the current study.

A. Fractal Theory

1. Introduction

Fractal theory, in it's present form, may be traced to studies conducted by Benoit Mandelbrot two decades ago. Mandelbrot (1971) provided a mathematical framework for the description of irregular and complex shapes found in nature. Fractal geometry has since been used to fit a variety of natural property distributions. Natural phenomena ranging from annual flowrates in rivers and the geometry of blood vessels (Gleick, 1987) to permeability distributions in porous media (Hewett, 1986; Goggin, 1988) have been nodeled with fractals. The search for fractal behavior in sedimentary rocks is closely linked to the fractal nature of hydrological cycles. Since most reservoir rocks were deposited under aqueous conditions and were subsequently subjected to diagenetic alteration in response to varying hydrogeological conditions, it would seem reasonable that the distribution of properties

within the rock may also be fractal in nature.

The term "fractal" may be described through the concept of "self-similarity". Self-similarity implies symmetry across scales of measurement. Hewett (1986) provides the following description:

"Fractals are characterized by the fact that they exhibit variations at all scales of observation and have partial correlations over all scales. Every attempt to divide such a geometry into smaller, more uniform regions results in the resolution of even more structure or roughness. The closer you look, the more detail you see."

By way of illustration, consider a coastline. Observed from an orbiting satellite, it appears to be very irregular. Move closer and the scale of observation changes, yet the degree of irregularity does not. Even at a distance of several feet the water is observed to meet the sand in an irregular pattern. The irregularity of the coastline is statistically "self-similar"; the degree of irregularity is the same at any scale of observation (Wheatcraft and Tyler, 1988). This characteristic irregularity may be quantified by a parameter called the fractal dimension.

2. Fractal Dimension

Consider the following simple equation in one dimension:

$$F = N\xi^{\eta} \tag{1V-1}$$

Where: $\eta =$ dimension in euclidean space.

(η = 1)
N = number of measurements.
ξ = length of measurement.
F = length of a line.

For the one dimensional example if the length of F is fixed, then ξ is proportional to N. For simple geometrical shapes in η - dimensional euclidean space the above relationship is satisfied. eg.

$$\lim_{\xi \to 0} (N\xi) = F \tag{IV-2}$$

As the length of the measurement decreases the number of measurements (N) increases accordingly and F remains constant. Under conditions of self-similarity, however, this may not be the case. Returning to the example of the irregular coastline it may be observed that $F \rightarrow \infty$ as $\xi \rightarrow 0$ (Wheatcraft and Tyler, 1988). A three dimensional example of this may observed for the case of ordinary chalk. As the measurement scale decreases, the surface area rapidly increases. This implies that decreasing the scale at which an observation is made reveals increased irregularity. In general, the problem of the scale at the scale decreases is the scale decreases the scale decreases the scale decreases the scale decreases increased irregularity.

$$F = N\xi^{\eta-D} = \text{constant}$$
 (IV-3)
Where: $D = \text{fractal dimension and } D \le \eta$.

D is the dimension which provides a constant value for F as the scale ξ changes. If Equation IV-3 is re-arranged and solved for D the following equation is obtained:

$$D = \lim_{\xi \to 0} \left\langle \frac{\ln N}{\ln \frac{1}{\xi}} \right\rangle \quad \text{For a constant F.} \tag{IV-4}$$

This result is the basis for the box counting theorem which is sometimes used to calculate fractal dimension (Barnsley, 1988). In the box counting technique, the number of boxes (N) of side length (ξ) required to cover an irregular surface is determined for a range of ξ values. The slope of a plot of $\ln N$ vs $\ln \frac{1}{\xi}$ then approximately defines the fractal dimension of the surface. A variation of this technique applicable to irregular, two-dimensional curves is the so called "structured walk" method. Returning to the example of the coastline, we may define two points of reference A and B. Four measurements of the length of this coastline are taken, each time using a measuring stick of shorter length. The only rule in the measurement process is that the measuring stick cannot be bent. Figure IV-1 illustrates this process.

For the first measurement, a measurement scale of 6.3 miles was used. Under these conditions, only a single measurement was required and the length of the shoreline was 6.3 miles. During the second measurement, a scale of 3.0 miles was chosen and 2.36 measurements were required to reach point B. This procedure was repeated for scales of 1.5 and 0.75 miles, respectively. The results of these measurements are summarized in Figure IV-1. Figure IV-2 shows a plot of the natural log of the number of measurements vs the natural log of the reciprocal of the measurement scale employed. From Equation IV-4, the fractal dimension may be represented by the slope of the best-fit line through these points. From Figure IV-2, the fractal dimension describing the irregularity of our imaginary coastline is 1.2344. This means that the degree of irregularity or roughness of the curve may be defined as constant over the scales of measurement used in the analysis. It should be pointed out, however, that this does not necessarily mean that the same fractal dimension applies to measurement scales larger or smaller than those evaluated.





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irregular enough to nearly fill two-dimensional space. The object is considered to have a fractional dimension since it is neither a line nor a plane in the standard geometrical sense (Barnsley, 1988). The co-dimension (X) of this object is a measure of the void space on the plane and is defined by $H=\eta-D$ where $\eta=2$ (Goggin, 1988). The fractal co-dimension in Equation IV-3 is related to the semi-variogram in the following manner (Journel and Huijbregts, 1978):

$$\gamma(h) = \frac{1}{2} E\{(Z(x+h) - Z(x))^2\} = \frac{1}{2} \sigma^2 h^{2H}$$
(1V-5)
Where: σ^2 is the total variance.

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0 < H < 1
```

Equation IV-5 implies statistical self-similarity, because variance at any scale may be computed from the variance at any other measurement scale by the following relationship (Mishra, 1987):

$$\gamma(bh) = b^{2H} \gamma(h) \qquad (IV-6)$$

A visual realization of the effect of D on the shape of the semi-variogram curve is shown in Figure IV-5. The group of curves where 1.5 > D > 1.0 (0.5 < H < 1.0) represent data which are more regular than ordinary white noise. Ordinary white noise is represented by D=1.5 (H=0.5) and traces where 1.5 < D < 2.0 (0.0 < H < 0.5) are noisier and more erratic





Stated another way, if D<1.5, the data series may be considered to be heterogeneous in terms of long range effects eg. the data shows correlation at smaller scales of measurement. If D>1.5, a process is described which tends to fluctuate between high and low values more commonly than would be expected for a purely random variable.

The data series of interest in the current study consist of permeability, porosity and mean pore throat size values which are equally spaced in the spatial domain. These values could be plotted on Cartesian coordinates and the resulting curve analyzed for it's irregularity using the structured walk technique previously illustrated. The disadvantage of such an approach is that this method is very time consuming and tedious to implement. Geostatistical methods provide a more convenient approach to the measurement of fractal dimension. The following section will describe two such methods which were employed in this study.

B. Geostatistical Methods

A regionalized variable may be defined as a variable distributed in space (Journel and Huijbregts, 1978). Several methods for determining a spatial correlation structure for regionalized variables are available. After a review of the literature, the following methods were chosen as the basis for a fractal analysis of the available data:

- 1. Autocorrelation functions.
- 2. Semi-Variogram analysis.

Before providing a detailed description of these methods and their relationship to the fractal dimension D, some statistical terminology will be briefly defined. The remainder of this chapter will then deal with fractal theory and provide a mathematical definition of the fractal dimension D.

Lag Distance (h)

In the discussion to follow, reference will be made to the lag distance h as it applies to a data series. By way of example, consider a core of a fixed length.



At regular intervals along the length of the core, a permeability value is obtained by removing a small sample and analyzing it by mercury porosimetry methods. In our discussion the length of the constant length interval between samples is the value of the lag distance h. For example, lag 1 refers to one spacing interval between samples, lag 2 to two spacing intervals between samples, etc. For our example lag 1 spacing will provide 10 data values in the data series while lag 2 spacing will provide only 5 data points (Figure IV-6).

Mean, Variance and Autocovariance

A common measure of central tendency is given by the mean value defined in Equation IV-7.

$$\tilde{\mu}_{x} = E = \frac{1}{n} \sum_{i=1}^{n} Z(i)$$
 (IV-7)

Where: $\bar{\mu}_{x}$ = mean value

Z(i) = value at location i in the x direction n = number of observations

The operator E is known as the expectation operator and is equivalent to the mean value. The mathematical expectation E may be applied to measurements of a particular property but is not limited to measurements. The variance for a series of measurements may be defined by Equation IV-8 where the expectation operator is applied to the squared difference between the value at a given location and the mean value for all locations.

$$\sigma_x^2 = E(Z(x) - \overline{\mu}_x) = \frac{1}{n} \sum_{i=1}^n (Z(x) - \overline{\mu}_x)^2 \qquad (IV-8)$$

Where: $\sigma_x^2 = variance$

Variables separated by lag distance h may be related to one another though the use of the covariance function. In the case of measurements of the same property conducted of a regular lag spacing h, the term autocovariance is often employed. Equation IV-9 defines the autocovariance function:

$$C_{v}(x, x+h) = E((Z(x) - \overline{\mu}_{x})(Z(x+h) - \overline{\mu}_{x})) \qquad (IV-9)$$

Where: $C_{v}(x, x+h) =$ autocovariance function.
 $E =$ expectation operator.
 $\overline{\mu}_{x} = E(Z(x))$
 $h =$ lag distance.

The preceding discussion will provide the reader with a basis for the following discussion of two methods of determining the correlation structure of a property in space. The discussion of each method will conclude with the procedure required to use each method in the determination of the fractal dimension D.

1. Autocorrelation Functions

If two random variables separated in space by lag h are of the same type, eg. permeability, then a measure of their spatial correlation may be obtained through the use of the autocorrelation function. The autocorrelation function is defined by:

$$C(x,x+h) = \frac{C_{v}(x,x+h)}{\sigma_{v}^{2}} = \frac{E\{(Z(x)-\bar{\mu}_{x})(Z(x+h)-\bar{\mu}_{x})\}}{\sigma_{x}^{2}}$$
(IV-10)

Where:

C(x, x+h) = autocorrelation function.

 $C_{\nu}(x, x+h)$ = autocovariance function.

$$E = \frac{1}{n} \sum_{i=1}^{n} Z(i)$$

$$\overline{\mu}_{x} = E(Z(x))$$

$$\sigma_{x}^{2} = E(Z(x) - \overline{\mu}_{x})^{2}$$

h = lag distance

The autocorrelation function as defined in Equation IV-10 is the quotient of the autocovariance of the data series and the variance for the same series of data. If Z(x) is stationary in a statistical sense, eg. E(Z(x+h))and the autocovariance both exist and only depend on h, then C(x,x+h) may be interpreted as the population correlation coefficient for the pair of variables (Z(x), Z(x+h)) (Chung, 1984). A direct relationship between the autocorrelation function and the semi-variogram discussed in the next section exists under these conditions.

Semi-variance = Variance - Autosorrelation Function

The fractal dimension (D) may be estimated from the autocorrelation function from the following relationship (Mandelbrot, 1971).

$$D_{acf} = 2 - \left\{ \frac{\log(2C(x, x+h)+2)}{2\log 2} \right\}$$
(IV-11)

C(x,x+h) is the autocorrelation value at lag 1, eg. the smallest h of the data series. The autocorrelation method has the advantage of being computationally simple, but is essentially a blind method as it only looks at the autocorrelation at lag 1 without incorporating information on longer range correlation. This may be important if the length of lag 1 happens to coincide with a natural periodicity in the property under study. In this case the property may be spatially correlated at lag 1, but not at any other scale of measurement and the autocorrelation function may provide a misleading indicator of the degree of autocorrelation.

Several assumptions are implicit in the use of the autocorrelation function to calculate fractal dimension.

- 1. E(Z(x+h)) exists and is only dependent upon h.
- 2. The autocovariance exists and only depends upon h.
- 3. The data series is well represented by a standard Gaussian distribution (D is approximately 1.5).

2. Semi-Variogram Analysis

The semi-variogram is a cornerstone of geostatistical study and has found broad application in the mining industry. Consider two random variables Z(x) and Z(x+h)

separated by a lag vector h. The degree to which point measurements show correlation over some distance in space or time may be characterized by the variogram function (Journel and Huijbregts, 1978).

$$2\gamma(x,h) = E\{(Z(x) - Z(x+h))^2\}$$
 (IV-12)

Substituting for the expectation operator E and re-arranging provides an expression for the semi-variance estimator.

$$\gamma(x,h) = \frac{1}{2N(h)} \sum_{i=1}^{N(h)} \{Z(x_i) - Z(x_i+h)\}^2$$
 (IV-13)

In this case, $Z(x_i)$ is a variable at location x_i in one dimensional space, N(h) is the number of pairs of data points separated by lag value h and $\gamma(x,h)$ is the experimental semi-variance. In general, the semi-variance is \otimes function of the location x as well as the lag value h. In practice, $\gamma(x,h)$ is often assumed to be invariant with x. This assumption is verified by testing the value of the mean within a given core for statistical homogeneity using a standard one-way analysis of variance test (ANOVA). The results of these tests are noted in Appendix B and confirm the applicability of using the variogram approach for the data sets generated by subroutine AREA. The intrinsic hypothesis that $\gamma(h)$ semi-variogram from a series of equally spaced data. Figure IV-7 illustrates the calculation procedure for a one-dimensional semi-variance.



For data showing spatial correlation, y(h) increases from near zero as $h \rightarrow 0$ to a roughly constant value at some finite distance h. The value of this constant variance is known as the sill and is numerically close to the total variance of the data. The value of h at which the sill is reached is known as the range and represents the point at which spatial correlation ceases (Goggin, 1988) (Figure IV-8).



In some cases $\gamma(h)$ does not approach zero as $h \rightarrow 0$ and a finite discontinuity occurs at the origin. This discontinuity is known as the "nugget effect" and is caused by a combination of sampling errors and (or) small variations at spacings less than the lowest available h. Journel and Huijbregts, (1978) state that local variability on this scale may be likened to the random phenomena of white noise which cannot be resolved at lag h.

Also of interest is the limiting case where $\gamma(h)$ appears only as a discontinuity at the origin and may be modelled by a horizontal line. This case is known as a pure nugget effect and represents a total absence of autocorrelation at all available scales of measurement. For the purposes of this study the pure nugget effect represents the highest possible level of heterogeneity discernable at a given lag value h.

Structural analysis of a regionalized phenomena involves fitting a mathmatical model to the experimental semi-variogram (Figure IV-8). Typical functional forms for semi-variograms with finite sill values are shown in Table IV-1. Fitting a variogram model which characterizes the main features of the regionalization requires a good physical knowledge of the property being studied. The form chosen to model the data will have a significant influence upon the range of the semi-variogram and may significantly influence the variogram interpretation in some cases. Spherical and linear variogram models provided the best fit to the observed data in this study (Appendix B).

The determination of the fractal dimension (D) from semi-variogram analysis in this study will utilize the method of Burrough (1981). This approach uses a log-log plot of the variance of the data separated by lag h vs h. The limiting slope of the plot as $h \rightarrow 0$ is then equated to 4-2D. Alternatively, the fractal dimension D may be estimated by the equivalent formula for the case of the

Equation	Model	Comment			
Linear	$\gamma(h) = \sigma^2 \left[\frac{h}{a}\right]$ $= \sigma^2$	0 <h<a h≥a</h<a 			
Spherical	$\gamma(h) = \sigma^2 \left\langle \frac{3h}{2\alpha} - \frac{1}{2} \left[\frac{h}{\alpha} \right]^3 \right\rangle$ $= \sigma^2$	0 <h<a h≥a</h<a 			
Exponential	$\gamma(h) = \sigma^2 \left\{ 1 - \exp\left[-\frac{h}{a}\right] \right\}$				
Note: $h = lag$ distance, $a = range$					

Table IV-1 Semi-Variogram Functional Forms

semi-variogram.

$$\lim_{n \to 0} \left\langle \frac{\ln \gamma(h)}{\ln h} \right\rangle = 2 - D_{var} \qquad (1V - 14)$$

Previous studies indicate that the fractal dimension D is a useful indicator of the complexity of autocorrelation of natural phenomena over many scales of measurement (Burrough, 1981; Hewett, 1986). The use of Burrough's method of calculating D also greatly reduces the role of interpretation in structural analysis of variograms. For small distances $h \rightarrow 0$ a linear model of the form $\gamma(h) = \omega h$ with ω equal to the slope of the log-log plot at the y-axis may be fitted to any model with linear behavior near the y-axis (Table IV-1). This greatly reduces the importance of model selection, particularly in cases for which the semi-variance is unstable resulting in a large degree of variability in the plotted data. The insensitivity of the value of D to the variogram model has been confirmed in studies by various authors (Shaw, 1988; Crickmore, 1990).

The following assumptions are implicit in the use of the Burrough method.

1. E(Z(x+h)) exists and is only dependent upon h.

2. The covariance exists and depends only on h.

Both the autocorrelation method and the semi-variogram method are computationally simple. The semi-variogram method has the advantage of providing a visualization of the spatial correlation behavior of the variable over all lag values and provides a validation of the fractal dimension obtained from the autocorrelation method. This

is not to say that the absolute values of D from both methods will be the same. Indeed, the fact that the methods make differing assumptions as to the probability distribution function that fits the data suggests that they should not be the same. It has been found, however, that in the absence of an a priori indication of the probability distribution function of a given set of data a combination of both methods will give a more robust estimator of D than any single method (Crickmore, 1990). As a result, both methods of evaluating D will be used in this study. As Hewett (1986) points out, when the lag distance of the semi-variogram approaches half the total length of the domain being modelled, the reliability of the variogram in estimating correlation structure decreases markedly. Thus, the semi-variogram and, by association, the autocorrelation function will only be useful for determining correlation structure over distances small compared with the total domain. This is not seen as a serious limitation in this study. The Fortran source code used to generate the semi-variogram data as well as the fractal dimension using the autocorrelation method is listed in Appendix E.

V. Experimental Apparatus and Procedure

The experimental work undertaken in this study may be grouped into two broad categories. The initial coreflood studies were conducted to determine recovery factors and effective dispersion coefficients (K.) for ten different cores. Capillary pressure measurements were conducted subsequent to the corefloods in an attempt to quantify the spatial correlation structure of three different properties for each of the ten cores. The following discussion will deal with the experimental design, apparatus and procedures used in these studies.

A. Coreflood Studies

1. Coreflood Apparatus

The apparatus used in the miscible corefloods (Figure V-1) is composed of the following components:

- A solvent injection system permitting displacement at a constant rate.
- B. A sealed coreholder containing the consolidated porous media.
- C. A constant temperature cabinet containing a core rotation unit as well as upstream and downstream pressure gauges.
- D. A backpressure control unit.
- E. A refractometer.



➢ Ruska positive displacement pump was used as the solvent injection system. The pump was a double cylinder type capable of discharge rates of between 10 and 330 cm³ per hour per cylinder. Injection rates were varied by changing gears in the transmission of the pump. This pump was connected to cylinders containing *n*-hexane and cyclohexane by a four-way valve. This allowed for the injection of either solvent or resident oil as required.

The coreholder used was constructed of stainless steel and was 48 inches long with an internal diameter of 2.5 inches. Endcaps with o-ring type seals were then bolted to each end of the coreholder. The endcaps used a sintered metal screen to evenly distribute the injected and produced fluids across the inlet and outlet faces of each core. The core rotator shown in Figure V-1 allowed for the rotation of the core at a rate of four revolutions per hour. This was necessary to minimize the effect of gravity segregation of the miscible fluids within the porous medium. A Tescom 2500 backpressure regulator was used to regulate the pressure of the system. This unit was linked to a rotating sample collection tray containing a series of 50 cm³ graduated centrifuge tubes. The resulting coreflood effluent was analyzed for refractive index using an ABBE model A303 refractometer.

2. Coreflood Design

One of the primary objectives of this study was to quantify the effect of the porespace upon mixing efficiency at the floodfront. As a result, it was deemed important to design the corefloods in such a manner that the effect of the pore structure upon mixing could be isolated. It is important to note that in this study the absolute values obtained for the recovery factor and K, for the corefloods are less important than their relative values. The objective; therefore, was to hold constant or minimize all factors affecting convective mixing with the exception of the pore structure itself. By doing this, the effect of the pore structure on convective mixing is emphasized. (Brigham et al (1961), Perkins and Johnston (1963) and Stalkup (1983) found that the following criteria are important in the design of a laboratory coreflood:

- 1. Gravity segregation.
- 2. Viscous fingering.
- 3. Minimum miscibility pressure.
- 4. Solvent injection rate.
- 5. Residual water saturation.
- 6. Geometric effects.
- 7. Pore structure effects.

A two-component first-contact miscible system was used in the current study. The components chosen had to satisfy the following requirements:

- 1. First-contact miscibility at standard temperature and pressure conditions.
- 2. A large difference in refractive index between the components.
- 3. A small difference in density between the components.

The choice of *n*-hexane and cyclohexane as the two components was found to satisfy the first two requirements. Table V-1 illustrates the properties of these two chemicals (Reid *et al*, 1987).

Table V-1 Properties of Miscible Components

Component	μ@25°C (cp)	Density (kg/m ³)	Refractive Index
n-hexane (C ₆ H ₁₄)	0.30	662.7	1.3770
cyclohexane (C ₆ H ₁₂)	0.88	782.0	1.4235

Under conditions of miscibility, the mobility ratio of the flood may be obtained from Equation V-1.

$$M = \frac{\mu_o}{\mu_s} \qquad (V-1)$$

Where: $\mu_s = viscosity$ of the solvent. $\mu_s = viscosity$ of the resident oil.

By selecting *n*-hexane as the resident oil the effective mobility ratio for each coreflood is held constant at approximately 0.341. By maintaining a favorable mobility ratio, the effect of viscous fingering can be minimized. The density difference between the two components is significant and may result in gravity segregation and a resulting reduction in recovery efficiency. This possibility was addressed by rotating the core using the core rotation unit shown in Figure V-1. Comparison of floods conducted on rotated and unrotated cores shows that the effect of gravity segregation may be greatly reduced using this method. An recovery factor increase of 3.3% was noted for core A using a flood rate of 0.0222 cm/s when the core was rotated.

The floodrate criteria have been examined in detail by Perkins and Johnston (1963). Figure V-2 shows a plot of $\frac{\kappa_i}{D_o}$ vs $\frac{vod_p}{D_o}$. For values of $\frac{vod_p}{D_o} > 4$ convective dispersion dominates the mixing process.



This plot is based upon Equation II-2 which describes the growth of the longitudinal dispersion coefficient with velocity. In the absence of channeling of the solvent, K_i is equivalent to the effective dispersion coefficient K_i . The objective in setting injection rates is to minimize the effect of molecular diffusion on the process. Three injection rates were used in this study. Darcian velocities of 35.0 cm/hr, 55.0 cm/hr and 80.0 cm/hr were used for each of the corefloods. The following rough

calculation was done to examine the validity of using these injection rates. Values for D_o and d_p are average values for a homogeneous sandstone (Stalkup, 1983).

$$d_p = 0.01 \text{ cm.}$$

 $D_o \cong 1.0 \text{ x } 10^{-5} \text{ cm}^2/\text{s}$
 $\sigma_{\min} = 2.0$
 $v_{\min} = \frac{35.0 \text{ cm/hr}}{3600 \text{ s/hr}} = 9.722 \text{ x } 10^{-3} \text{ cm/s}$
 $\frac{v_{\min}\sigma_{\min}d_p}{D_o} = 19.44$

This result would seem to indicate that the effect of the molecular diffusion coefficient D_o is minimal, particularly at the highest velocity tested. In the region where D_o dominates the mixing process, K_i increases very slowly with the injection rate (Figure V-2). Three injection rates were used to ensure that the flood was conducted to the right of this region. By maintaining a constant Darcian velocity for each flood, the value of K_o will be dependent upon the heterogeneity coefficient σ , which; in turn, is a function of the structure of the porous medium.

The residual water saturation has also been found to affect the efficiency of the miscible displacement process (Stalkup, 1970). This factor was minimized by evacuating each of the cores for an extended period of time. This resulted in the vaporization and removal of most of the residual water from each core. In general, residual water saturation was not seen as a significant factor in these experiments as eight of the ten cores tested were outcrop samples with low residual water saturations.

Geometric effects were first investigated by Brigham et al, (1961) who found that boundary effects increased convective dispersion (K_{\bullet}) when core diameters were less than 3.2 cm. Core diameters in the current study varied between 4.445 cm and 5.556 cm. As a result, the diameter of the core was not seen as a controlling factor. The effect of core length can also have an influence upon the efficiency of the process. Brigham et al (1961) found that the amount of mixing was proportional to the square root of the distance travelled. This would imply that recovery is improved when the displacement path length is increased. For eight of the ten cores tested, the core length was held constant at 121.92 cm. The exceptions to this were two oilfield carbonate cores (core F and core G) which were 136.53 and 96.52 cm long, respectively (Table V-2).

Core	Lithology	Diameter (cm)	Length (cm)	Porosity %	k (md)
A	Indiana limestone	4.445	121.92	21.04	54.75
В	Brown sandstone	5.080	121.92	21.42	2035.48
С	Brown sandstone	5.080	121.92	21.90	1967.20
D	Indiana limestone	4.445	121.92	21.25	42.92
E	Berea sandstone	4.445	121.92	24.56	2231.72
F	Swan Hills limestone	5.080	136.53	12.26	138.36
G	Golden Spike limestone	4.445	96.52	11.58	51.21
Н	Berea sandstone	4.445	121.92	27.05	2155.91
I	Tyndle limestone	5.556	121.92	9.54	2.25
J	Tyndle limestone	5.556	121.92	8.99	5.76

Table V-2 Core Properties

If the second ption is made that the amount of mixing in a given products medium is proportional to K, then K, for the shorter core should be slightly lower than would be the case for a core 121.9 cm long. Conversely, the longer core should have a slightly higher effective dispersion coefficient than would be expected for a 121.9 cm core.

3. Coreflood Procedure

Eight of the ten cores shown in Table V-2 were obtained from outcrop samples. Cores A, B, C, D, E and H were purchased from Cleveland Quarries and were already cut to the appropriate length. Cores I and J were cored from a solid limestone block while cores F and G were oilfield cores. As can be seen from their characteristics, these cores represent a variety of pore structure styles with varying degrees of heterogeneity. The first step in the coreflood procedure involved installing the cores in the coreholder. A molten metal technique was used whereby each core was initially sealed with an epoxy paint to prevent any absorption of the molten metal into the porespace. The core was then centered in the core barrel using a three pin endcap which sealed one end of the coreholder. The core and coreholder were then placed vertically in an oven and heated to 150° F. A quantity of Cerrobend alloy was then melted and poured into the

Each core was saturated with *n*-hexane by first evacuating the core overnight using a portable vacuum pump. The quality of the vacuum was tested through the use of a vacuum gauge connected to the opposite end of the coreholder. This procedure allowed f the removal of the residual water which remained in the core. After evacuation, the coreholder was attached to the Ruska pump and *n*-hexane was injected into the core until the vacuum as measured by the gauge had dropped to zero and liquid was produced at the gauge end of the core. The amount of *n*-hexane injected was then used to calculate the porosity of the core using a bulk volume obtained from measurements of the core.

After saturation, the coreholder was mounted in the constant temperature cabinet shown in Figure V-1. Permeability was determined by injecting n-hexane at four

different rates and measuring the differential pressure across the core at each rate. From this data the liquid permeability was calculated from Darcy's law for linear fluid flow. From the pore volume measured during the core saturation, floodrates corresponding to the desired Darcian velocities were calculated. Because the pump transmission offered a finite choice of gear ratios it was not always possible to obtain the exact velocity required, however, it was usually possible to select an injection rate within 5% of the desired rate. The backpressure was arbitrarily set to 300 psi (2.068 MPA). Although not required for miscibility, this backpressure allowed for better control of the injection process as the Ruska pump would occasionally inject at too high a rate in the absence of backpressure. Once the pump speed was set, the four-way valve was switched over to cyclohexane and the core rotation unit was activated in preparation for the start of the flood. A volume of effluent corresponding to 0.05 pore volumes was calculated and this amount of fluid was collected in each centrifuge tube as the flood progressed. As the tubes were filled the refractive index of the effluent was measured using the refractometer. The flood was allowed to continue until 2.0 pore volumes of cyclohexane had been injected into the core. The data

collected consisted of a refractive index value for each 0.05 PV of cyclohexane injected. The computer analysis of this data will be described later in this chapter.

After each cyclohexane flood, the core was re-flooded with *n*-hexane until a refractometer test showed the effluent to be pure *n*-hexane. Due to the adverse mobility ratio of the re-flood, up to 4.0 pore volumes of *n*-hexane were required to achieve complete removal of the cyclohexane. Duplicate cyclohexane floods on core A separated by a *n*-hexane re-flood produced identical effluent concentration profiles. This indicated that the re-flooding procedure could be used to restore the core to it's initial state immediately after the initial saturation with *n*-hexane. Each core was flooded at the three rates described earlier, each cyclohexane flood being followed by a re-flood with *n*-hexane.

B. Mercury Porosimetry

The use of mercury porosimetry provides a means of evaluating the pore structure of discrete samples taken from each core. After the corefloods were complete, each core was dried by passing compressed air through it for several days. The core was then removed from the coreholder by melting the surrounding Cerrobend. Figure V-3 illustrates the sample locations chosen for this study. A

requirement for variogram or autocorrelation function calculation is that the data points be evenly spaced. As can be seen, a total of 50 samples per core were analyzed. This number was, by necessity, a compromise between 'ne data requirements of the statistical calculations and the time required to analyze each sample. Each sample had a volume of approximately 1 cm³ and was obtained by cutting the core into pieces using a diamond saw.



Once the samples were obtained they were first dried in an oven at $150^{\circ}F$ for several days and then coded as to their exact location within the core. The analysis of the samples was then undertaken.

The use of mercury porosimetry in the study of pore structure has been undertaken by several investigators. Wardlaw (1976) provides an excellent description of its use in an attempt to quantify pore structure behavior in. two-phase flow. In the current study mercury injection was used to determine values for the permeability, porosity and mean pore throat diameter for each of the 500 samples analyzed.

Purcell (1949) was the first to apply the capillary pressure curve determined from mercury porosimetry to the calculation of permeability. Equation V-2 illustrates Purcell's equation for permeability calculation. The integral in Equation V-2 is simply the area under a plot of $\frac{1}{p_c^2}$ vs the corresponding mercury saturation. Porosity is calculated by dividing the total volume of mercury injected into the sample at an arbitrarily high pressure by the bulk volume of the sample.

$$k = 10.24(\sigma_t \cos \theta)^2 \phi \lambda_L \int_{s=0}^{s=1} \frac{dS}{(P_t)^2}$$
 (V-2)

Where: k = permeability (md).

- $\phi = \text{porosity}.$
- S = fraction of perespace invaded by mercury.
- P_c = capillary pressure (psi).
- σ_t = interfacial tension (dynes/cm).
- θ = contact angle.
- λ_{L} = lithology factor.

The determination of the capillary pressure curve is relatively straightforward. The sample is placed in the sample cell containing mercury shown in Figure V-4.

The first step in the procedure involves the determination of a calibration curve for the apparatus used. The sample cell is first evatuated for several minutes using the vacuum pump. The cell is then flooded with mercury until the mercury reaches a mark in the top sightglass. Pressure is applied to the empty cylinder in increments from 0 to 1000 psi using the nitrogen source. At each pressure increment the piston crank is turned so as to bring the mercury level back up to the sightglass mark. After each movement of the piston crank a volume reading is taken and recordet from the scale on the piston crank.



In this way, a calibration curve of pressure vs volume of mercury injected is obtained. After the calibration curve is obtained the sample is placed in the cell and the cell is flooded with mercury. The bulk volume of the sample is then obtained from the difference between the measured volume of mercury and the known volume of the cell. After determination of the bulk volume a vacuum is then applied to the cell for several minutes. The same procedure of gradually increasing the pressure on the sample by increments while recording the volume of mercury forced into the porespace at each pressure was then followed. The same pressure increments were used as in the case of the calibration curve and care was taken to allow enough time between pressure increases for mercury intrusion within the sample to occur. Once the data was collected, the calibration converse was subtracted from the sample data leaving the capillary pressure curve. The volumes of mercury injected were converted into mercury saturations at each incremental pressure and the values for permeability, porosity and mean pore throat size were calculated by means of the computer program ANALYZE contained in Appendix E.

C. Data Analysis

Appendix E contains the source code for the Fortran programs EFFPLOT and ANALYZE which were used to analyze the raw data obtained from the coreflood and mercury porosimetry experiments, respectively. The datafile used by the EFFPLOT program consists of a set of calibration data as well as a refractive index value for each 0.05 P.V. of solvent injected. The calibration data was obtained by preparing a series of eleven mixtures of *n*-hexane and cyclohexane and measuring their refractive indices. The mixtures varied in composition from 100% *n*-hexane to 100% cyclohexane by volume and were prepared in increments of 10%. The first part of the EFFPLOT program uses a linear interpolation method to convert each of the refractive index values measured in the coreflood into a cyclohexane

concentration. The values for pore volumes injected are then converted to a lambda function by the following equation:

$$\lambda = \frac{\frac{v}{v_{a}} - 1}{\sqrt{\frac{v}{v_{a}}}}$$

Where: V = pore volumes injected.

 V_p = total pore volume.

The lambda values along with the corresponding cyclohexane concentrations are then output by the program and are listed in Appendix D. These values are then plotted on an arithemetic probability scale as shown in Figure V-5.



The straight line portion of the plot is used to calculate the effective dispersion coefficient (K_*) using the following equation:

$$K_{\varphi} = \nu L \left\{ \frac{\lambda_{\varphi 0} - \lambda_{10}}{3.625} \right\}^2$$

Where: $K_{\bullet} = \text{effective dispersion coefficient cm}^2/\text{s}.$

v = darcian velocity (cm/s).

L =length of the core (cm).

 $\lambda_{90}, \lambda_{10} = \lambda$ values at cyclohexane

concentrations of 90% and 10%.

This procedure was followed for each of the corefloods run. As a final step, the EFFPLOT program calculates the area underneath the cyclohexane concentration curve from 0 to 1.6 P.V. injected and uses this value to obtain a recovery factor for the flood. Since we are interested in the recovery factor for the *n*-hexane which is the resident oil, the recovery factor may be calculated from the following formula:

$$R.F. = 100 - \int_{0.0P.V.}^{1.0P.V.} C_* dC_* \qquad (V-3)$$

Where: R.F. = recovery factor for the resident oil. C_s = concentration of the solvent (%).

The integral in Equation V-3 was evaluated numerically using Simpson's 1/3 rule (Gerald and Wheatley, 1985). Due to errors in the calculation of the area beneath the curve
it was found that the recovery factor sometimes exceeded 100% after injection of 1.5 and 2.0 pore volumes of solvent. A maximum value of 105.5% recovery efficiency was noted for Core E after the injection of 2.0 pore volumes of solvent. The program automatically reduces any value over 100% back to 100% recovery before printout. This over prediction of recovery was not seen at the 1.0 pore volume injection level which was used for the comparisons in this study (Tables VII-1 - VII-3).

Program ANALYZE was used to process the data from the capillary pressure measurements into values for permeability, porosity and mean pore throat size for each sample. The main portion of the program reads in the data and subtracts the calibration curve values from the sample data. The resulting capillary pressure curve is then passed on to subroutine AREA where the porosity is calculated. The pressure at which 50% mercury saturation occurs is then calculated by linear interpolation. It has been noted that this pressure may be related to the mean pore throat size in the sample by Equation V-4 (Wardlaw, 1976). For the purposes of this calculation and the calculation of permeability, it was assumed that the interfacial tension was 480.0 dynes/cm and the contact angle between mercury and the non-wetting surface was 140°.

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$$d = \frac{4\sigma_{t}\cos\theta}{P_{c}} \qquad (V-4)$$

Where: $d = \text{mean pore throat diameter (10}^{-9} \text{ m.)}$ $\sigma_t = \text{interfacial tension (dynes/cm)}.$ $\theta = \text{contact angle}.$ $P_c = \text{capillary pressure (mpa)}.$

These are average values obtained from the original work by Purcell (1949). After the calculation of the mean pore throat diameter the program calculates permeability using Purcell's equation (Equation V-2). For this calculation, an average lithology factor (λ_i) of 0.216 was used (Amyx et al, 1960). The integral portion of Equation V-2 was calculated by fitting a natural spline curve to the calculated data points and numerically evaluating the area beneath the curve using the spline coefficients. This approach does not require equally spaced data points as is the case with Simpson's rule and is somewhat more accurate (Gerald and Wheatley, 1985). Once the permeability has been calculated a table is printed giving the sample location as well as the permeability, porosity and mean pore throat diameter for each sample. These tables are listed in Appendix A.

Subroutine VARIO uses the permeability data to calculate the semi-variogram and autocorrelation function

at lag 1 for the data series using the equations described in chapter IV. The data obtained was joined into one continuous series by taking each location series of ten data points (eg. location 2, levels A to J, Figure V-3) and adding the location series to the end of the previous location series (eg. location 1, levels A to J, Figure V-2). The values for the mean and variance were then calculated for the series and printed out. This method of calculation is similar to the calculation of the semi-variogram in two dimensions (Journel and Huijbregts, 1978 p 220.). The major difference is that the assumption is made that each of the location vectors (Figure V-3) may be represented by a column in a rectangular two-dimensional matrix. Each layer represents a row in the matrix and the semi-variance is calculated in the direction parallel to the axis of the core only. Due to the fact that the semi-variogram calculation is sensitive to the occurrence of extreme values it is common to calculate the log semi-variance of the property under study to minimize this effect (Chung, 1984; Mishra, 1987). This approach was followed in subroutine VARIO which calculates lag values and the natural log of the associated semi-variances. As a final step, the calculated autocorrelation function at lag 1 was transformed into a fractal dimension using Equation IV-11.

VI. Heterogeneity Description

Traditionally, heterogeneity within a porous medium has been defined in a variety of ways. In the case of immiscible displacement, by far the most commonly used estimator of heterogeneity is the Dykstra-Parsons permeability variation coefficient, V_{DP} . For the case of miscible displacement, the Koval heterogeneity factor (H) is commonly used. In this chapter the calculation of these heterogeneity indicators from the data obtained in the current study will be discussed. These calculations will form the basis for a comparison with a fractal heterogeneity estimator which will be discussed in the following chapters.

A. Dykstra - Parson's V Dr

Original work conducted by Dykstra-Parsons (1950) resulted in a method of calculating oil recovery from a field subject to immiscible displacement. It was found that recovery was a function of the mobility ratio (M), the initial oil and water saturations and the variation in absolute permeabilities within the porous medium. For our purposes, only the method of quantifying the permeability variation will be discussed in detail.

The calculation of the permeability variation

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coefficient V_{DP} assumes a log-normal distribution of permeability within the porous medium. The calculation procedure is as follows (Dykstra-Parsons, 1950):

- All permeabilities in a distribution are listed in decending order.
- 2. The percentage of the permeabilities exceeding each listed permeability are computed and tabulated as the portion of the total sample having a higher permeability.
- 3. The permeability values in step 1 are plotted on a log scale while the percentage of permeabilities greater than the permeability value are plotted on the probability axis of a log-probability scale.
- 4. A best-fit straight line is drawn through the points. If the points do not lie on a straight line, the terminal points are weighted less.
- 5. The permeability at 84.1 cumulative percent is subtracted from the median permeability. The result is then divided by the median permeability yielding the permeability variation V_{DP} .

Figure VI-1 illustrates the calculation procedure for a sample data set:



The graphical method of calculating V_{DF} implicitly ignores the effects of any extreme permeability values. These outliers are typically few in number. High values for permeability may occur due to localized fractures and will occur as points to the upper left on Figure VI-1. Conversely, low permeability outliers caused by impermeable streaks in the rock will occur to the lower right of Figure VI-1. The exclusion of these values is equivalent to a truncation of one or both tails of a frequency distribution curve for the data set in question. V_{DP} may also be calculated without the use of graphical techniques provided that the standard deviation of the natural log of the permeability distribution is known (Jensen and Lake, 1988). Equation VI-1 provides a means of calculating V_{DP} under these circumstances:

$$V_{DP} = 1 - e^{(-\sigma_{ln}(k))}$$
 (VI-1)

Where: V_{OP} = Dykstra-Parsons permeability variation.

 $\sigma_{in(k)}$ = standard deviation of the permeability distribution.

The use of Equation VI-1 will require the removal of any outliers in the data which may tend to cause bias in the results. The criteria for the removal of outliers is discussed further in Chapter VII. Appendix C contains a listing of the outliers removed as well as the calculated variance of the natural log of the permeability distribution for each core. After removal of the extreme values, $\sigma_{in(k)}$ was calculated through a slight modification of the source code of program ANALYZE in Appendix E. Table VI-1 lists the results of these calculations.

Core	Ø _{ln(k)}	V _{DP}
A	2.05	0.87
В	0.56	0.43
сс	0.99	0.63
D	1.83	0.84
E	1.09	0.66
F	2.44	0.91
G	2.42	0.91
Н	0.43	0.35
I	0.94	0.61
J	1.18	0.69

Table VI-1 V_{DT} Results

B. <u>Koval's H Parameter</u>

Koval (1963) adapted the Buckley-Leverett fractional flow equation for use in miscible displacement. In Koval's analysis, the flowing fraction of the displacing phase, F_* is assumed to be dependent upon solvent saturation, heterogeneity effects and viscosity differences. Equation VI-2 was developed by Koval for use in miscible displacement and is analogous to the familiar Buckley-Leverett fractional flow equation.

$$F_{s} = \frac{1}{1 + \left(\frac{1-s}{s}\right)\left(\frac{1}{H}\right)\left(\frac{1}{E}\right)}$$
(VI-2)

Where: F_s = flowing fraction of solvent.

S = solvent saturation.

H = heterogeneity factor.

E = effective viscosity ratio.

The effective viscosity ratio E is further defined by Equation VI-3 which is an empirical correlation based upon the fourth root mixing rule used in refinery calculations.

$$E = \left(0.78 + 0.22 \left(\frac{\mu_o}{\mu_*}\right)^{\frac{1}{4}}\right)^4$$
 (VI-3)

Where: *E* = effective viscosity ratio.

 μ_o = viscosity of the resident oil.

 μ_s = viscosity of the solvent.

Substituting K=HE into Equation VI-2 and re-arranging the terms yields:

$$V_{\rm pi} = \frac{K}{(K - F_{\rm e}(K - 1))^2}$$
(VI-4)

Where: V_{pi} = pore volumes of solvent injected.

Equation VI-4 suggests a simple means of calculating K and hence, the heterogeneity coefficient, H. At a point in time just prior to solvent breakthrough, $F_* = 0.0$. If we substitute $F_* = 0$ into Equation VI-4 we obtain the following expression for K based upon the volume of solvent injected at the time that breakthrough occurs (Koval, 1963):

$$K = \frac{1}{(V_{pi})_{b.t.}}$$
(VI-5)

Where: $(V_{pi})_{b.t.}$ = volume of solvent injected at breakthrough (P.V.) K = HE

Equation VI-5 is the basis for the calculation of the Koval heterogeneity factor H in the current study and is valid under the following conditions:

- 1. A linear, first-contact miscible system is used.
- 2. Solvent is injected continuously.
- 3. Gravity effects are eliminated.
- Flood rates are high in order to minimize diffusion effects.
- Permeability depends only upon the saturation of the oil or solvent.

The design of the coreflood experiments in this study satisfies conditions 1-4 while the 5th condition was assumed to be valid in the absence of interfacial tension.

Figures VI-2 to VI-7 show the effluent concentration profiles for the ten cores evaluated in this study. In each figure, the point at which solvent breakthrough occurred is indicated.







Based upon Equation VI-3 and the viscosity data contained in Table V-1, the effective viscosity ratio E for the floods shown in Figures VI-2 to VI-7 is 0.80803. For a completely homogeneous system. H=1 and breakthrough would occur at 1.237 P.V.. This calculation shows that Equation VI-3 cannot be applied to systems with favorable viscosity ratios without modification as material balance considerations limit the breakthrough pore volume to 1.0 P.V.. In light of this, we will simply assume that a completely homogeneous system will have an H value of 1.237. We can then normalize all of the H values by dividing them by 1.237 to be consistent with the scaling used by Koval. By doing this we can arrive at a relative scale where a completely homogeneous system is represented by a normalized H value of 1.0. This normalization is required to account for the fact that the mobility ratio of the displacements was less than 1.

Table VI-2 shows the results of these calculations.

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Core	P.V. @ Breakthrough	K	Н	Normalized H
A	0.56	1.785	2.209	1.785
В	0.75	1.333	1.650	1.333
С	0.91	1.098	1.359	1.098
D	0.56	1.785	2.209	1.785
E	0.92	1.086	1.345	1.086
F	0.56	1.785	2.209	1.785
G	0.62	1.612	1.996	1.612
H	0.83	1.204	1.491	1.204
I	0.27	3.703	4.583	3.703
J	0.27	3.703	4.583	3.703

Table VI-2 Koval's H Parameter

VII. Results and Discussion

A. Line of Investigation

A review of the literature reveals that the effect of spatial correlation of hydraulic properties (eg. permeability, porosity and pore throat characteristics) upon miscible recovery efficiency has received relatively little attention. In the current study an investigation of the following topics related to convective dispersion (Ke) and the efficiency of a first-contact miscible process was carried out:

- Relationship between floodrate, convective dispersion (K.) and recovery factor.
- Effect of lithology on convective dispersion
 (K.) and recovery efficiency.
- Effect of permeability, porosity and mean pore throat size variances on convective dispersion and recovery efficiency.
- Relationship between fractal dimension of permeability, porosity and mean pore throat size and dispersion and recovery efficiency behavior.

Unlike many coreflood experiments described in the literature, the current study was designed to isolate the effect of the pore struct re on convective dispersion and recovery factor. As a result, the effect of the lithology type upon K, and recovery efficiency may be qualitatively assessed. To the best of the author's knowledge, the current study is also unique in its evaluation of the relationship between fractal dimension and convective dispersion on the laboratory scale.

In the remainder of this chapter the four topics noted above will be discussed. In addition, a comparison of a fractal heterogeneity estimator with the well-known Dykstra-Parsons and Koval heterogeneity coefficients will be presented. As a final step, a possible method of predicting field scale dispersivities from laboratory measured dispersivities using fractal dimensions will be examined.

B. <u>Discussion</u> of <u>Results</u>

1. Effect of Floodrate and Convective Dispersion on Recovery Efficiency

Tables VII-1 through VII-3 summarize the results obtained for each of the corefloods undertaken in this study. 91

Core	Lithology	∨ (cm/s)	K. cm²/s	R.F. @ 1.0 P.V.	R.F. @ 1.5 P.V.	R.F. @ 2.0 P.V.
A	Indiana Lst.	0.0224	0.0948	87.6	95.1	96.1
В	Brown Sst.	0.0215	0.0270	92.6	98.3	100.0
С	Brown Sst.	0.0220	0.0105	94.6	100.0	100.0
D	Indiana Lst.	0.0223	0.0108	89.9	98.7	99.3
E	Berea Sst.	0.0216	0.0037	92.9	96.3	96.3
F	S.Hills Lst.	0.0229	0.3050	81.8	83.0	83.2
G	G.Spike Lst.	0.0213	0.1930	82.5	92.5	96.3
H	Berea Sst.	0.0222	0.0053	84.9	86.2	86.4
I	Tyndle Lst.	0.0229	0.2913	84.4	93.3	97.0
J	Tyndle Lst.	0.0230	0.5149	80.4	88.3	91.8

Table VII-1 Coreflood Results (Target rate=0.0222 cm/s)

Table VII-2 Coreflood Results (Target rate=0.0153 cm/s)

Core	Lithology	∨ (cm/s)	K. cm²/s	R.F. @ 1.0 P.V.	R.F. @ 1.5 P.V.	R.F. @ 2.0 P.V.
A	Indiana Lst.	0.0145	0.0437	88.0	94.0	94.7
В	Brown Sst.	0.0152	0.0165	94.2	100.0	100.0
С	Brown Sst.	0.0155	0.0062	96.9	100.0	100.0
D	Indiana Lst.	0.0167	0.0455	89.9	96.7	97.2
E	Berea Sst.	0.0161	0.0020	97.4	100.0	100.0
F	S.Hills Lst.	0.0185	0.0122	92.8	95.4	96.0
G	G.Spike Lst.	0.0164	0.1220	89.0	100.0	100.0
H	Berea Sst.	0.0142	0.0040	91.0	93.7	93.8
I	Tyndle Lst.	0.0146	0.1462	85.5	96.0	99.8
J	Tyndle Lst.	0.0148	0.2535	81.0	90.1	93.6

Core	Lithology	∨ (cm/s)	K cm²/s	R.F. @ 1.0 P.V.	R.F. @ 1.5 P.V.	R.F. @ 2.0 P.V.
A	Indiana Lst.	0.0110	0.0210	90.6	97.9	98.3
В	Brown Sst.	0.0088	0.0099	95.5	100.0	100.0
С	Brown Sst.	0.0086	0.0030	96.9	100.0	100.0
D	Indiana Lst.	0.0090	0.0246	90.4	98.8	99.3
E	Berea Sst.	0.0092	0.0011	98.0	100.0	100.0
F	S.Hills Lst.	0.0092	0.0294	93.7	96.7	96.8
G	G.Spike Lst.	0.0095	0.0709	89.5	100.0	100.0
н	Berea Sst.	0.0091	0.0017	94.7	97.0	97.2
I	Tyndle Lst.	0.0091	0.0768	86.2	98.1	100.0
J	Tyndle Lst.	0.0096	0.0924	83.6	93.9	97.4

Table VII-3 Coreflood Results (Target rate=0.0097 cm/s)

In general, it may be noted that recovery efficiency for a given rock type increases with a decrease in the flood velocity. Figure VII-1 illustrates the behavior of the ten cores studied. The first point on the left of each polygon represents a floodrate of 0.0222 cm/s (80 cm/hr), the second point, a rate of 0.0153 cm/s (55 cm/hr) and the point on the right a rate of 0.0097 cm/s (35 cm/hr). Note that the recovery factor shows an increase with decreasing rate in all cases.



This effect occurs in stable displacements as a result of the reduced effect of convective dispersion at lower flow rates (Perkins and Johnston, 1963). Shelton and Schneider (1975) also attributed this result to a more efficient stripping of the resident oil from dead-end porespace by diffusion processes.

Several studies have focused on the effect of convective dispersion upon the recovery efficiency of a first-contact miscible process. Spence and Watkins (1980) showed that recovery efficiency decreased with increasing convective dispersion. Stalkup (1983) observed that recovery efficiency is affected by the degree of dispersion in the miscible transition zone. Increasing amounts of convective dispersion tended to produce wider transition zones and less efficient displacements. Figure VII-2 shows a semi-log correlation of convective dispersion and the recovery factor. It may be observed that the recovery factor at 1 pore volume of solvent injected increases as K. decreases. Based upon studies conducted on Berea sandstones by other researchers, there is reason to suspect that the recovery factor of 84.87% obtained for Core H is unusually low for this type of sandstone (Shelton and Schneider, 1975). An inspection of the effluent concentration profile reveals that early breakthrough of the solvent occurred during flooding of Core H indicating that channeling of the solvent may have occurred, thus explaining the relatively low recovery factor. In any case, the value obtained for K, appears to be valid based upon the fact that K_{\star} is only calculated from the portion of the effluent concentration profile



which occurs after breakthrough as well as the fact that these portions of the profile are very similar for both Core E and Core H (Figure VI-3). A further validation of K. is found in the work of Spence and Watkins (1980) who reported K. values of approximately 0.005 cm²/s for Berea cores. If the Berea sandstone cores are grouped separately from all of the other rock types then the remaining data are well fit (r=0.9753) (Figure VII-2) by a logarithmic correlation of the following form:

$$K_{\bullet} = 199774015.9 (10)^{-0.107027(R.F.)}$$
(VII-1)

Where: $K_{\bullet} = \text{effective dispersion coefficient } (cm^2/s).$

R.F. = recovery factor @ 1 P.V. injected.

This equation is equivalent to plotting log K. vs R.F. and obtaining a linear "best-fit" line through the points. The data from the Berea cores also appear to show an inverse correlation between K, and the recovery factor although this inference is only based upon two data points. The results obtained agree with earlier findings regarding the inverse relationship between convective dispersion and recovery efficiency (Spence and Watkins, 1980).

2. Effect of Lithology on K, and Recovery Efficiency

Several observations were noted regarding the effect of lithology upon the value of K, and the associated recovery factor. Figure VII-3 shows the effect of grouping the core data points by lithology on a standard coordinate axis.



It is interesting to note that for the Berea cores, K, is as much as an order of magnitude less than the K, values for any of the other cores. This may be explained qualitatively by the fact that Berea sandstone is recognized as one of the most homogeneous rock types available. Hence, it would be expected that convective dispersion caused by heterogeneity in the pore structure would be relatively small.

In general, it was also noted that the cores having higher values of K_{\star} and lower recovery factors tended to be carbonates while those having higher recoveries tended to be sandstones (Figure VII-1). Also of interest is the fact that the recovery factor appears to be much more sensitive to changes in K, for the sandstones than is the case for the (presumably) more heterogeneous limestones. Figure VII-3 shows this effect. Note that the slope of the straight lines connecting cores of equivalent rock type tend to flatten out as one progresses from limestones to sandstones. The relationship between lithology and K_{\bullet} is a complex one and caution must be exercised in the interpretation of these results. The fact that only ten cores were tested under identical conditions suggests that these findings should be viewed as qualitative results only.

3. Effect of Permeability, Porosity and Mean Pore Throat Size Variance on Dispersion Behavior and Recovery Efficiency

Studies by Bretz et al (1988) related pore size variation to convective dispersion using thin-section techniques. It was found that wide pore size distributions could be correlated with larger amounts of convective dispersion. In the following discussion an examination will be carried out on the effect of variance of the three properties tested upon the values of K_{\star} , log K_{\star} and the recovery factor. The use of σ^2 to represent the width of the frequency distribution curve is valid provided that any outliers in the data are removed first. In general, outliers were recognized by comparing the data values obtained for each core and removing data which appeared to be unusually high. This process was fairly straightforward in that the outliers were few in number and were often at least an order of magnitude higher than values in the rest of the data set. Aside from making the estimation of variance more representative for the particular core being studied, the removal of outliers also has a physical basis. In the case of porosity, outlier values between 0.40 and 0.50 occurred occasionally and are probably the result of errors in the measurement of bulk volume. Permeability values greater than 1000 md were also excluded from the analysis on the basis that they appear to represent samples which were fractured. The case may be made that these fractures are an important part of the structure of the porous medium and should not be removed. As a test of this possibility, they were removed in calculating variance estimates but were used in the calculation of fractal dimension. None of the mean pore throat size data sets showed evidence of outliers using

the above criteria (Appendix C).

Table VII-4 illustrates the findings of the study regarding the relationship between variance estimates and K, and R.F.

Property Variance	vs	К.	log K.	R.F. @ 1 P.V.
Porosity		r=0.126	r=0.354	r=0.312
Permeability		r=0.100	r=0.284	r=0.289
Mean Pore Throat Diameter		r=0.349	r=0.036	r=0.214

Table VII-4 Correlation of Variance With K, and R.F.

A variety of correlation models were applied to the data. A linear "least-squares" model was found to yield the highest values for the correlation coefficient, r and was used in all cases.

In all cases, the degree of positive or negative correlation between the calculated parameters is quite low. This would indicate that convective dispersion and recovery factor are relatively insensitive to statistical variance estimates of the measured properties. Although this finding appears valid for the current study, it is by no means conclusive. It is quite possible that the number of samples studied and their location were inappropriate for the proper estimation of variance in each core. If the procedure is accepted as valid, however, then it appears that the use of property variance as a means of describing the heterogeneity of a porous medium and hence predicting its flow characteristics cannot be recommended.

4. Relationship Between Fractal Dimension of Permeability, porosity and Mean Pore Throat Size and Dispersion and Recovery Efficiency

Table VII-5 summarizes the results of the fractal dimensional analysis for each core. It may be noted that the fractal dimensions obtained using semi-variograms are 500ch higher and more erratic than those calculated from auxoscorrelation functions. This is due, in large part, to the fact that many of the semi-variograms were highly variable in nature thus making the determination of the slope near the origin a somewhat subjective process. Particular difficulty was experienced when the initial slope of the plot was negative. Figure VII-4 shows a sample calculation. Note that the determination of the slope as h approaches zero is subject to a fair amount of uncertainty. In addition, many of the semi-variograms exhibited a fairly low slope near the origin.

Core	Permeability		Porosity			n Pore at Size
	Daci	D _{var}	Daci	Ľ _{var}	Daci	D _{var}
A	1.546	1.071	1.437	1.937	1.437	1.821
В	1.521	1.889	1.496	1.998	1.620	1.985
С	1.510	1.621	1.489	1.996	1.357	1.975
D	1.513	1.917	1.577	1.992	1,688	1.948
Е	1.467	1.136	1.467	1.993	1.364	1.945
F	1.525	1.541	1.560	1.675	1.438	1.770
G	1.531	1.357	1.565	1.926	1.509	1.973
Н	1.479	1.972	1.403	1.996	1.546	1.953
I	1.518	1.862	1.532	1.989	1.636	1.996
J	1.546	1.437	1.391	1.987	1.469	1.968

Calcu

tal Dimensions

As a result of these factors, the fractal dimensions obtained from the two processes were treated separately (Appendix B).

The values shown in Table VII-5 were compared with K_{er} log K. and recovery factor (R.F.) using a linear "least-squares" approach. As was the case in the previous section, the effect of excluding data for the Berea cores was also evaluated. Correlation coefficients (r) were calculated for the case where all cores were included and the correlation was then recalculated excluding the Berea cores.



Tables VII-6 through VII-8 show the results of these calculations.

	vs	K.	log K.	R.F.
D _{acf} ⟨₩/O B. sst.)		r=0.800	r=0.634	r=0.697
D _{acj} (all cores)		r=0.740	r=0.858	r=0.515
D _{var} (w/o B. sst.)		r=0.208	r=0.274	r=0.374
D _{var} (all cores)		r=0.128	r=0.084	r=0.042

Table VII-6Permeability Fractal Correlations

	vs	K e	log 🔬	R.F.
D _{acf} (W/O B. sst.)		r=0.261	r=0.011	r=0.049
D _{acf} (all cores)		r=0.017	r=0.296	r=0.042
D _{var} (W/O B. sst.)		r=0.271	r=0.392	r=0.468
D _{var} (all cores)		r=0.343	r=0.431	r=0.454

Table VII-7Porosity Fractal Correlations

Table VII-8 Mean Pore Threat Size Fractal Correlations

	vs	K .	log K.	R.F.
$D_{acf}(w/o B.$ sst.)		r=0.189	r=0.004	r=0.060
D _{acj} (all cores)		r=0.008	r=0.251	r=0.133
D _{var} (w/o B. sst.)		r=0.079	r=0.243	r=0.270
D _{var} (all cores)		r=0.119	r=0.240	r=0.255

Calculations involving the fractal dimension of porosity and mean pore throat size did not show a significant degree of correlation with K. and R.F. This suggests that K. and R.F. are independent of the fractal dimension of porosity and mean pore throat size. This finding is not surprising in that flow characteristics in a porous medium have been found to be controlled by permeability variations. Dykstra and Parsons (1950) noted that although total recovery is a function of porosity, the recovery factor is influenced by permeability variations. Under conditions of miscible and immiscible displacements, we may expect the porosity distribution to influence only the total recovery and then only if all of the porosity were contacted with equal efficiency by the displacing fluid. This would require that _he permeability be the same at all locations.

The lack of significance of the fractal dimension for mean pore throat size may be attributed to the absence of capillary forces under miscible conditions. Under conditions of immiscible displacement, the size of the pore throats will have a direct influence upon relative permeability and hence will directly affect fluid flow in the porous medium. In the case of miscible displacement, mean pore throat size does not appear to be as important.

The most significant correlations were observed when the fractal dimension of permeability was examined. An examination of Table VII-6 shows that there is a large difference in the degree of correlation when using fractal dimensions calculated by the semi-variogram method (D_{var}) and using the method of autocorrelation functions

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 (D_{art}) . While the use of semi-variograms in the calculation of fractal dimension is valid on a theoretical basis, in practice it was found to be difficult to apply. In particular, it was difficult to determine the slope of the semi-variogram as $h \rightarrow 0$ due to the erratic behavior of some of the semi-variograms (Appendix B). The fitting of a model to a semi-variogram and the determination of the slope of the model as $h \rightarrow 0$ is a very subjective process for the most part and is subject to errors of interpretation on the part of the user. Because the calculation of the fractal dimension by the method of Burrough (Equation IV-14) is sensitive to these variations the use of D_{var} as a consistent indicator of fractal dimension cannot be recommended. A more promising approach involves the use of autocorrelation functions in the calculation of fractal dimension. This method has the advantage of eliminating subjectivity from the calculation procedure.

Figure VII-5 illustrates the degree of correlation between D_{ac} , and log K. for permeability. Note that the degree of correlation is generally very good and improves when the Berea cores are included in the analysis (r=0.858). Conversely, Table VII-6 shows that the degree of correlation between D_{acf} and R.F. decreased when



the Berea cores were included. It appears that this result may be attributed to a lower than expected recovery factor for core H. The calculated values for convective dispersion (K,) are considered to be accurate for all cores as discussed earlier. As a result, including the Berea cores improves the correlation in the case of the D_{acf} vs K, plot.

A correlation of the following form exists between D_{acf} and K. in Figure VII-5:

 $K_{\bullet} = 4.8698442 \times 10^{-60} (10)^{38.41279204(D_{acf})} (VII-3)$

Where: K_{e} = effective convective dispersion coefficient (cm^{2}/s).

D_{acf} = fractal dimension for permeability calculated from autocorrelation functions.

The fact that a correlation appears to exist between D_{aci} and K. (Equation VII-3) and also between K. and R.F. (Equation VII-1) suggests the development of an equation which directly relates D_{aci} to R.F.. Substituting K. from Equation VII-3 into Equation VII-1, we get:

$$R \cdot F \cdot = 631.73801 - 358.9075(D_{acf})$$
(VII-4)
$$1.4815 < D_{acf} < 1.7602$$

Where: R.F. = recovery factor (%).

 D_{acf} = fractal dimension for permeability determined from autocorrelation functions.

determined from autocorrelation functions

5. Comparison of D_{acf} with V_{DP} and Koval's H

Figure VII-6 provides a comparison of D_{acf} , the Dykstra-Parsons permeability variation, V_{DF} , and Koval's heterogeneity factor, H. Data from Tables VI-1, VI-2 and VII-5 were plotted vs the recovery factor at 1 P.V. injected.

While the spread of the data points around the best-fit line may seem otherwise, the best correlation with R.F. (r=0.806) is obtained for the Koval factor, H. The reason for this is that the calculation of H is very



sensitive to the degree of channeling which takes place within the core. This channeling effect is measured directly from effluent concentration profiles and has a direct influence upon the calculated R.F. (eg. early breakthrough implies that some of the oil is being bypassed and recovery will be less efficient). Also of interest is the fact that as the H factor increases, the data points appear to show a greater degree of scatter. This result appears to support the contention of McCaffery
et al. (1978) who questioned the assumption that effective permeability is dependent only upon the saturation of the oil or solvent in more heterogeneous systems.

It seems probable that the defining equation for the effective viscosity ratio (Equation VI-3) used by Koval is inadequate for describing the effective viscosity ratio in more heterogeneous systems. This would not be surprising in that Equation VI-3 was based upon experiments conducted in homogeneous cores only (Koval, 1963). This result invites further study into the applicability of the effective viscosity ratio equation in more heterogeneous systems.

Unlike Koval's H, D_{acf} and V_{DF} are calculated independently of the effluent concentration profile (eg. are independent variables) and show a lesser degree of correlation with recovery factor. Figure VII-7 shows the degree of correlation between the heterogeneity estimators and log K. Note that in this case the best correlation is provided by D_{acf} (r=0.840). This result was noted earlier however, Koval's H factor also shows a relatively good fit to log K. (r=0.785). This suggests that channeling is a strong contributor to the value of the effective dispersion coefficient K. The Dykstra-Parson's coefficient showed the lowest degree of correlation when



plotted vs R.F. and log K_{\bullet} . This result was expected in light of the fact that V_{DP} represents an estimate of the width of the permeability frequency distribution and should show similar results to attempts to correlate permeability variance with R.F. and log K_{\bullet} (Table VII-4).

Of importance in these comparisons between heterogeneity estimators are the treatment of outliers or extreme values. It is significant that V_{DP} and the permeability variance were calculated by excluding outliers from the analysis while D_{acf} included all values. It is possible that these extreme permeability values control the behavior of fluid flow within a porous medium by facilitating the channeling of solvent and bypassing of resident oil. Koval's H factor quantifies the effect of these extreme values in causing early breakthrough however, its efficiency in predicting R.F. and K. in more heterogeneous rock types is in doubt considering the spread of the data points about the best-fit line for more heterogeneous cores shown in Figures VII-6 and VII-7. The use of V_{pr} is unsuitable in that it ignores the effect of high permeability values which may predispose the porous medium to channeling of the solvent. Unlike waterflooding, channeling in miscible displacement is more critical to the process in that dilution of a solvent slug and loss of miscibility can occur very rapidly, thus rendering the process ineffective.

Figure VII-8 shows a comparison between D_{acf} and H. Although the correlation is only fair (r=0.523), the point of breakthrough on the effluent concentration profile is subject to an error of 0.05 P.V.. This translates into an error in H of \pm 0.22. Qualitatively, however, it appears that Dacf can give an indication of the degree of channeling provided that extreme values are included in the analysis.



6. Scale Dependence and D_{acf}

The question we must now ask is why go to the trouble of determining D_{acf} when Koval's H seems to correlate with log K, and R.F. nearly as well? The answer to this question lies in the fact that K, varies with the scale of the system under consideration. It is this scale dependence which creates difficulties in applying the results of laboratory investigations to field scale developments. The use of the fractal dimension (D) as an estimator of heterogeneity suggests a means of overcoming this difficulty. The basic advantage of this approach is that the fractal dimension is invarient over all scales of measurement that it is evaluated from. Wheatcraft and Tyler (1988) observed that dispersivity does not vary in a linear manner with scale but appears to be dependent upon the degree of "regularity" of the permeability variation at all available scales of measurement within a given system. Equation VII-5 illustrates an equation developed by Wheatcraft and Tyler (1988) describing the relationship between fractal dimension and dispersivity under conditions where the laboratory and field flood rates are the same.

$$Ke_{field} = Ke_{lab} \xi_c^{1-D} L^{D-1}$$
 (VII-5)

Where: Ke_{field} = field measured dispersivity (m).

Ke_{lab} = lab measured dispers'vity (m).

 ξ_c = smallest scale of measurement used

in determining D.

L = length of the system under study (m).

D = fractal dimension for permeability.

note that Ke_{field} and $Ke_{tab} = \frac{K_e}{v}$ where K sub e= the effective dispersion coefficient $\left(\frac{cm^2}{s}\right)$ and v = the darcian velocity of the displacement $\left(\frac{cm}{s}\right)$.

This equation has several interesting characteristics. For the limiting case where the porous medium is perfectly homogeneous $(D_{acj}=1.0)$ Ke_{jield} is equal to Ke_{lab} and is independent of scale. In addition, dispersivity increases proportionately to L^{D-1} . For the limiting case where D=2, dispersivity is proportional to the scale of measurement, L. A preliminary test of the applicability of this equation may be undertaken by substituting D_{acf} (permeability) from Table VII-5 for D, setting $\xi_c =$ 0.1219 m. and using Kelab values from Table VII-3 $(v \cong 0.0097 \text{ cm/s})$. Ke_{field} values for a variety of different scales (L) may then be generated. It was decided to group the ten cores by lithology with the value of D_{acf} (perm.) used being a mean value in cases where more than one core was available. The following is a sample calculation for the Indiana limestones.

> mean value of $K_{e} = 0.0228 \text{ cm}^{2}/\text{s}$ mean darcian velocity=0.01003 cm/s $Ke_{iab} = \frac{0.0228}{0.0100} = 2.2721 \text{ cm}.$ mean value of $D_{acf} = 1.5295.$ $L = 100 \text{ m}, Ke_{jield} = 79.31 \text{ m}.$ $L = 1000 \text{ m}, Ke_{jield} = 268.44 \text{ m}.$

A set of six curves were generated illustrating the growth of dispersivity with fractal dimension (Figure VII-9). As a test of the applicability of this approach, field data relating dispersivity to the scale of observation is also shown. This data, obtained by Gelhar *et al* (1985) represents the results of numerous tracer tests from around the world. Gelhar's data is based upon field displacement rates which are lower than the lowest rate evaluated in this study (Table VII-3). As a result each of the curves shown in Figure VII-9 may be treated as being upper boundaries on the expected value of Ke_{field} at a given scale (L) for each lithology type. In addition, the rock types represented in Gelhar's data were primarily sandstones with a small number of limestone aquifers being included.

Figure VII-9 shows that the curves generated for Indiana limestone, Berea and Brown sandstone as well as Swan Hills limestone show a realistic growth in dispersivity with scale as indicated by the clustering of the field data. The Golden Spike and Tyndle limestones tended to lie above the region occupied by the field data. These lithologies had the highest fractal dimension (D_{acf}) and tended to show a greater increase in dispersivity with scale. Although none of the field data obtained falls into



the region of these curves it does not necessarily follow that they are unrealistic. The first point to consider is that much of Gelhar *et al's* data described aquifers in which the source rock was significantly different than the Golden Spike and Tyndle limestones in the current study. It may be the case that all of the field data represents lithologies that are more homogeneous than Golden Spike or Tyndle limestones.

The second point to consider involves the behavior of dispersivity with displacement rate. In general, as the rate decreases, dispersivity also decreases for a given porous medium. For more heterogeneous rock types (eg. higher D_{acf}), it appears that this reduction of dispersivity is much more pronounced as the rate is decreased than is the case for more homogeneous rock types. If this is the case, a reduction to actual field rates may reduce Ke_{lab} for the limestones in question to the point where the upper two curves in Figure VII-9 lie within the upper boundary of the available field data.

Use of the fractal dimension as a means of determining dispersivities on a field scale will require a better validation of the preliminary comparison shown in Figure VII-9. It would be instructive to isolate the field tracer data by lithology and compare it with individual dispersivity curves shown in Figure VII-9. In addition, it is important that D_{acl} be determined using larger scales of measurement than were practical in the present study. Current research into this possibility is centering around the use of S.P. logs in the calculation of fractal dimension on a field wide basis. By defining the fractal dimension for a field or portion of a field it may be possible to develop a set of correlation curves relating dispersivity to scale using fractal dimension as a cross-plotting parameter.

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VIII. Conclusions and Recommendations

The relationship between several heterogeneity indicators and the mixing behavoir of a first-contact miscible process was investigated in the current study. Based upon the experiments conducted in this study, it may be concluded that:

- 1. he fractal dimension for permeability (D_{acf}) may be used to scale up dispersivity values $\left(\frac{\kappa_*}{\nu}\right)$ from laboratory scale to field scale.
- 2. The Dykstra-Parson's permeability variation V_{DP} does not correlate well with either log K. or R.F.. It appears that the calculation of V_{DP} ignores the effect of extreme permeability values which appear to influence channeling and convective dispersion.
- 3. A positive logarithmic correlation exists between K. and the calculated fractal dimension for permeability (D_{acl}) .
- 4. The recovery factor shows a linear correlation with D_{acf} .
- 5. The Koval heterogeneity factor (H) shows the best correlation with recovery factor but appears to be less reliable for more heterogeneous rock types.
- 6. Daci and Koval's H provide the best correlation with

 $\log K_{\bullet}$.

- Recovery efficiency increases with decreased flood velocity for a given rock type.
- Recovery efficiency (R.F.) shows an inverse logarithmic correlation with convective dispersion (K.) for all rock types.
- 9. Carbonates showed the highest values of K, and the lowest recovery factors while sandstones; in general, showed lower K, values and a higher recovery factor.
- 10. The more homogeneous the rock type, the greater the sensitivity of the recovery factor to changes in the degree of convective dispersion.
- 11. Permeability, porosity and mean pore throat size variances showed little correlation with K. or R.F. and cannot be recommended as a means of describing heterogeneity in porous media.
- 12. Calculation of fractal dimensions is best accomplished through the use of autocorrelation functions (D_{acl}) . The erratic behavoir of some of the semi-variograms near the origin makes their use onreliable.

Several additional areas of experimentation are suggested from the results of this study:

- The effect of length upon convective dispersion for different rock types needs to be quantified in greater detail.
- 2. A revised equation for the effective viscosity ratio (E) used in the calculation of Koval's H will need to be developed. The current equation does not appear to adequately describe the effective viscosity in more heterogeneous rock types.
- Methods of calculating fractal dimensions for permeability on a field wide basis will need to be developed. This work is ongoing.

In general, the use of the fractal dimension for permeability (D_{acf}) to describe heterogeneity shows promise. The chief advantage of this approach is that it opens up the possibility of scaling up laboratory measured dispersion to field levels in a theoretically valid manner. This study, although preliminary in nature, points out several avenues of additional research which may allow for the practical implementation of such a method in the future.

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Appendix A

The following is a listing of the raw data calculated for each core by the computer program ANALYZE which is listed in Appendix C. The layer and location headings refer to the location of the sample in the core (Figure V-3).

Layer	Location	Porosity	Permeability (md.)	Mean Pore Throat Size (nM)
A	1	0.45	170.19	6012.19
В	1	0.18	62.27	5650.28
с	1	0.11	30.93	1987.08
D	1	0.13	0.10	1631.11
Е	1	0.12	38.04	2884.06
F	1	0.11	3.69	1119.32
G	1	0.13	28.08	273.13
Н	1	0.13	396.65	5257.15
I	1	0.15	57.59	2546.02
J	1	0.16	74.48	6175.54
A	2	0.25	59.19	3173.74
В	2	0.41	187.34	6824.40
С	2	0.13	41.34	1503.45
D	2	0.13	47.99	1221.62
E	2	0.14	46.24	1494.82
F	2	0.14	38.88	2319.48
G	2	0.10	209.31	1141.39
H	2	0.15	53.52	3325.58
I	2	0.14	72.53	6219.95
J	2	0.15	0.1	2378.74
A	3	^ 44	243.18	7342.27
В	3	C.15	0.1	1105.09
С	3	0.15	37.37	2809.45
D	3	0.18	70.91	4140.57
E	3	0.15	0.1	5619.84

Table A-1 Data Summary - Core A

Layer	Location	Porosity	Permeability (md.)	Mean Pore Throat Size
				(nM)
F	3	0.14	44.41	2438.83
G	3	0.12	6885.13	4316.91
Н	3	0.15	207.48	1635.50
I	3	0.14	68.33	2278.93
J	3	0.14	170.41	1325.11
A	4	0.16	41.72	1514.36
В	4	0.42	234.77	7397.46
С	4	0.15	63.6	6208.80
D	4	0.14	41.99	2278.93
E	4	0.13	39.42	1699.48
F	4	0.14	0.1	1999.27
G	4	0.15	59.51	3089.14
H	4	0.16	55.27	3504.41
I	4	0.13	47.20	2546.03
J	4	0.13	38.11	1605.29
A	5	0.18	51.46	1234.01
В	5	0.23	54.38	2649.54
С	5	0.48	117.10	4232.77
D	5	0.13	53.06	2386.21
E	5	0.12	3.19	1141.38
F	5	0.14	26.73	1246.65
G	5	0.14	46.37	2201.92
H	5	0.12	55.18	2062.55
I	5	0.16	54.74	2509.26
J	5	0.14	36.70	1620.26

Table A-1 Data Summary - Core A (cont'd)

Layer	Location	Porosity	Permeability (md.)	Mean Pore Throat Size (nM)
A	1	0.19	51.09	4383.91
В	1	0.18	65.32	5731.44
С	1	0.18	72.59	6301.16
D	1	0.15	203.56	7718.46
Е	1	0.18	49.26	4474.66
F	1	0.13	27.34	3852.12
G	1	0.17	61.36	5591.47
Н	1	0.28	35.98	2334.58
I	1	0.22	53.09	4268.42
J	1	0.20	79.33	6178.19
A	2	0.27	121.99	6701.32
В	2	0.13	155.38	7892.10
С	2	0.15	44.9	4897.74
D	2	0.19	50.41	4163.66
E	2	0.22	269.89	7962.33
F	2	0.19	61.12	4195.31
G	2	0.17	62.03	5554.52
Н	2	0.21	94.12	6633.69
I	2	0.21	32.66	3437.19
J	2	0.18	49.07	4369.52
A	3	0.18	77.52	6664.83
В	3	0.14	34.69	3899.61
С	3	0.16	28.20	3615.50
D	3	0.18	47.24	4134.31
E	3	0.17	57.59	5237.65

Table A-2 Data Summary - Core B

T T T T T T T T T T	Location	Porosity	Permeability	Mean Pore Throat
Layer	LOCATION	Porosicy	(md.)	Size
				(nM)
F	3	0.12	33.13	4166.36
G	3	0.18	1610.78	7948.57
H	3	0.21	109.83	7106.77
I	3	0.21	55.97	4342.27
J	3	0.18	59.77	5155.08
A	4	0.20	67.28	5144.51
В	4	0.19	197.67	7975.11
с	4	0.28	54.60	2818.96
D	4	0.16	123.57	7911.63
E	4	0.19	91.69	6728.25
F	4	0.17	63.94	5549.46
G	4	0.17	48.38	4357.48
Н	4	0.21	59.56	4584.87
I	4	0.22	127.28	7264.23
J	4	0.19	77.21	7678.57
A	5	0.28	61.35	3022.98
В	5	0.11	231.01	7746,54
С	5	0.17	31.45	3683.43
D	5	0.16	44.12	4401.94
Е	5	0.20	70.04	5265.11
F	5	0.17	138.03	7273.44
G	5	0.18	164.76	7835.41
Н	5	0.22	56.48	4429.75
I	5	0.22	52.24	4112.57
J	5	0.22	159.84	7610.64

Table A-2 Data Summary - Core B (cont'd)

Layer	Location	Porosity	Permeability (md.)	Mean Pore Throat Size (nM)
A	1	0.19	84.59	6563.13
В	1	0.21	68.08	5219.72
С	2	0.22	75.11	5219.72
D	1	0.22	118.44	7104.54
E	1	0.22	72.75	5063.46
F	1	0.21	60.78	4629.70
G	1	0.21	45.04	3986.78
Н	1	0.21	72.51	5387.54
I	1	0.21	74.08	5330.06
J	1	0.18	125.71	5878.48
A	2	0.19	58.55	4970.03
В	2	0.20	72.78	5576.12
С	2	0.21	79.96	5957.44
D	2	0.21	62.36	4821.91
Е	2	0.22	65.94	4732.90
F	2	0.22	61.52	4492.25
G	2	0.18	36.25	3835.50
H	2	0.23	39.85	3573.02
I	2	0.23	53792.41	8107.00
J	2	0.24	82.05	5285.14
A	3	0.22	93.05	6817.42
В	3	0.22	107.88	7041.46
С	3	0.19	75.47	6213.79
D	3	0.20	70.52	5421.15
E	3	0.23	231.42	4793.16

Table A-3 Data Summary - Core C

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Layer	Location	Porosity	Permeability (md.)	Mean Pore Throat Size (nM)
F	3	0.22	85.97	6078.5
G	3	0.20	38.84	3601.23
Н	3	0.22	49.05	3803.93
I	3	0.24	58.30	4048.70
J	3	0.17	0.1	6256.92
A	4	0.2	100.13	6918.34
В	4	0.20	91.99	6698.60
С	4	0.20	127.94	6406.27
D	4	0.19	58.94	4877.23
E	4	0.20	56.87	4602.73
F	4	0.21	89.27	6519.36
G	4	0.23	88.18	5939.98
Н	4	0.23	82.69	5650.28
I	4	0.27	44.22	3688.59
J	4	0.19	117.51	7670.36
A	5	0.23	81.77	5585.43
В	¥,	0.21	161.86	7823.84
С	5	0.18	84.08	4793.15
D	5	0.21	56.39	4260.44
E	5	0.16	55.55	5479.36
F	5	0.21	63.31	4772.90
G	5	0.19	85.68	6694.06
Н	5	0.23	42.04	3749.22
I	5	0.24	46.54	3733.88
J	5	0.20	36.78	3587.71

Table A-3 Data Summary - Core C (cont'd)

Layer	Location	Porosity	Permeability (md.)	Mean Pore Throat Size (nM)
A	1	0.16	42.90	2278.93
В	1	0.13	29.51	1266.12
c	1	0.18	123.14	6591.68
D	1	0.20	100.01	4212.26
E	1	0.14	42.24	3242.84
F	1	0.19	85.71	5940.00
G	1	0.17	68.27	2791.40
Н	1	0.18	74.58	6282.96
I	1	0.11	51.79	4112.57
J	1	0.21	107.12	6476.17
A	2	0.22	49.05	2922.86
В	2	0.17	62.62	5484.70
С	2	0.23	135.89	6278.32
D	2	0.20	7910.38	2278.92
E	2	0.19	39.43	3926.72
F	2	0.16	48.20	2622.88
G	2	0.18	52.53	4949.16
Н	2	0.19	45.24	4007.20
I	2	0.16	67.82	5401.14
J	2	0.19	90.83	6288.49
A	3	0.14	38.48	1461.30
В	3	0.25	88.47	6384.11
С	3	0.18	34.41	2112.69
D	3	0.16	80.91	3926.72
E	3	0.15	48.12	3601.23

Table A-4 Data Summary - Core D

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Layer	Location	Porosity	Permeability (md.)	Mean Pore Throat Size (nM)
F	3	0.16	78.27	4426.86
G	3	0.16	54.09	3390.46
Н	3	0.16	0.1	6514.54
I	3	0.16	52.47	5020.39
J	3	0.21	149.66	6883.24
A	4	0.18	33.33	2062.54
В	4	0.14	53.35	2450.30
С	4	0.17	74.35	6245.27
D	4	0.15	48.29	2509.26
E	4	0.16	31.27	1180.13
F	4	0.16	120.07	4793.15
G	4	0.16	0.1	5873.07
Н	4	0.14	34.71	2062.55
I	4	0.15	0.1	2165.34
J	4	0.19	84.07	5387.56
A	5	0.19	32.87	1969.06
В	5	0.16	40.77	657.20
С	5	0.18	90.32	6311.02
D	5	0.20	74.64	375 6.94
E	5	0.18	34.09	2546.03
F	5	0.16	55.91	3325.58
G	5	0.13	691.36	2044.89
Н	5	0.19	91.46	4495.57
I	5	0.10	0.1	845.29
J	5	0.21	69.44	5000.02

Table A-4 Data Summary - Core D (cont'd)

Layer	Location	Porosity	Permeability (md.)	Mean Pore Throat Size (nM)
A	1	0.24	112.76	6767.60
В	1	0.21	90.77	6506.76
С	1	0.25	100.06	6233.25
D	1	0.27	58.08	3777.35
E	1	0.26	161.50	7462.50
F	1	0.24	92.31	5854.23
G	1	0.23	122.10	5310.07
Н	1	0.26	44.22	3383.12
I	1	0.24	89.82	6043.27
J	1	0.3	53.34	3417.12
A	2	0.22	87.18	5873.07
В	2	0.21	79.03	5763.72
С	2	0.28	56.25	4061.98
D	2	0.32	88.42	4518.14
Е	2	0.21	119.13	7678.10
F	2	0.22	86.38	6187.75
G	2	0.24	85.83	5278.44
Н	2	0.27	41.76	3371.66
I	2	0.27	47.12	3340.62
J	2	0.29	46.06	3189.92
A	3	0.24	170.43	7042.74
В	3	0.22	89.10	6275.62
С	3	0.24	84.83	5495.13
D	3	0.24	92.31	6068.39
E	3	0.22	146.38	7597.20

Table A-5 Data Summary - Core E

Layer	Location	Porosity	Permeability (md.)	Mean Pore Throat Size (nM)
F	3	0.20	95.38	6571.96
G	3	0.27	127.55	6747.16
H	3	0.25	188.21	7819.02
 	3	0.26	102.75	6298.82
 J	3	0.28	58.48	3523.35
A	4	0.22	83.87	6059.90
B	4	0.27	58.13	3677.44
c	4	0.25	80.38	4916.43
D	4	0.212	0.1	4639.59
E	4	0.21	85.99	6335.77
F	4	0.23	162.52	7799.96
G	4	0.24	558.16	7837.04
H	4	0.23	111.45	6808.85
I	4	0.28	47.89	3450.69
J	4	0.24	44.84	3601.23
A	5	0.22	62.39	4586.16
В	5	0.24	42.02	3504.41
С	5	0.22	65.92	4694.50
D	5	0.21	76.97	5608.84
Е	5	0.26	190.76	7793.72
F	5	0.22	84.64	6202.23
G	5	0.24	85.10	5496.60
Н	5	0.23	66.22	4351.48
I	5	0.25	423.64	7595.27
J	5	0.26	45.68	3601.23

Table A-5 Data Summary - Core E (cont'd)

Layer	Location	Porosity	Permeability (md.)	Mean Pore Throat Size (nM)
<u>A</u>	1	0.02	17.50	1266.12
В	1	0.10	42.34	6268.52
с	1	0.07	27.77	6607.49
D	1	0.09	50.93	6864.75
E	1	0.16	1310.61	7767.68
F	1	0.19	0.1	7931.10
G	1	0.06	27.19	6659.81
Н	1	0.07	30.50	7086.50
I	1	0.07	53.86	7191.98
J	1	0.11	78125.68	7432.90
A	2	0.03	15.38	7025.37
В	2	0.09	44.99	8149.99
С	2	0.06	0.82	1294.41
D	2	0.23	151.68	7741.50
E	2	0.05	43.09	7724.92
F	2	0.21	79.88	5932.47
G	2	0.04	13.02	3089.11
H	2	0.05	0.1	6773.46
I	2	0.09	19095.19	7164.41
J	2	0.15	69.32	7515.56
A	3	0.12	97.18	7651.40
В	3	0.16	88.99	7207.62
с	3	0.14	86.55	7177.13
D	3	J.06	0.1	2649.54
Е	3	0.05	19.25	3089.11

Table A-6 Data Summary - Core F

Layer	Location	Porosity	Permeability (md.)	Mean Pore Throat Size
				(nM)
F	3	0.20	0.1	7832.48
G	3	0.06	24.53	6349.95
Н	3	0.06	33.26	6409.44
I	3	0.03	33.06	2546.02
J	3	0.09	694.71	7467.20
A	4	0.04	1087.09	7692.99
В	4	0.05	0.1	2704.52
с	4	0.16	59.15	6104.72
D	4	0.16	86.53	7408.75
Е	4	0.15	66.04	7413.04
F	4	0.16	64.58	6570.25
G	4	0.03	513.92	7067.29
Н	4	0.11	0.1	6604.31
I	4	0.13	110.84	7515.14
J	4	0.10	61.32	6638.91
A	5	0.06	0.1	7513.88
В	5	0.09	48.33	7493.95
С	5	0.09	81.70	7363.09
D	5	0.08	36.62	6898.95
Е	5	0.10	47.14	6404.04
F	5	0.05	296.91	6991.46
G	5	0.10	47.14	6404.04
Н	5	0.08	2.10	523.04
I	5	0.10	25.06	3457.92
J	5	0.08	408.17	6283.63

Table A-6 Data Summary - Core F (cont'd)

Layer	Location	Porosity	Permeability (md.)	Mean Pore Throat Size (nM)
A	1	0.10	61.03	7428.62
В	1	0.05	33.11	6750.91
С	1	0.04	82913.06	6397.18
D	1	0.11	79196.68	7515.14
E	1	0.09	57.86	7686.32
F	1	0.04	287.60	7337.25
G	1	0.06	0.1	2546.02
Н	1	0.09	25.01	3986.77
I	1	0.08	22093.80	6970.35
J	1	0.09	35.53	6185.23
A	2	0.15	66.61	7581.13
В	2	0.02	53.40	4316.89
С	2	0.09	148.68	7537.29
D	2	0.11	44.72	6791.11
E	2	0.14	67.04	7275.73
F	2	0.06	0.1	6568.65
G	2	0.05	0.1	6206.89
H	2	0.07	33.88	7216.14
I	2	0.07	22.84	4464.77
J	2	0.07	400.67	7035.49
A	3	0.13	63.22	7058.58
В	3	0.03	31724.70	3926.72
С	3	0.08	0.1	7276.91
D	3	0.12	100.38	7802.32
E	3	0.19	0.1	7911.17

Table A-7 Data Summary - Core G

Layer	Location	Porositý	Permeability (md.)	Mean Pore Throat Size (nM)
F	3	0.09	57.29	7408.75
G	3	0.06	24.90	6563.12
н	3	0.07	581.13	7700.57
I	3	0.08	44.09	6862.62
J	3	0.08	0.1	7244.05
A	4	0.13	388.88	7232.95
В	4	0.03	14.29	5387.54
С	4	0.07	313.14	6669.48
D	4	0.05	14.99	7353.02
E	4	0.14	79.54	7252.53
F	4	0.07	55.70	7329.22
G	4	0.04	10.37	2405.08
Н	4	0.05	283643.6	7193.34
I	4	0 .09	50.9	6961.81
J	4	0.06	2.81	6818.44
A	5	0.15	70.05	6710.14
В	5	0.06	27.85	7363.09
С	5	0.08	57.96	6740.03
D	5	0.12	93.09	6807.81
Е	5	0.09	0.1	7267.12
F	5	0.06	388891.3	7358.56
G	5	0.04	242.51	7105.81
H	5	0.09	5.24	6935.66
I	5	0.08	71.12	7408.75
J	5	0.1	45.65	6442.02

Tabile A-7 Data Summary - Core G (cont'd)

Layer	Location	Porosity	Permeability (md.)	Mean Pore Throat Size (nM)
A	1	0.27	90.52	4882.94
В	1	0.26	99.01	5849.11
С	1	0.25	52.02	3765.51
D	1	0.23	44.76	3697.00
E	1	0.23	101.58	6451.03
F	1	0.25	46.39	3629.89
G	1	0.25	45.91	3479.78
H	1	0.23	85.65	5786.16
I	1	0.36	69.15	3512.62
J	1	0.33	33.40	2365.23
A	2	0.21	71.55	5296.94
В	2	0.22	63.53	4516.34
С	2	0.25	46.41	3508.18
D	2	0.23	35.30	3427.12
E	2	0.24	90.18	5492.32
F	2	0.21	79.34	5854.23
G	2	0.25	66.47	4221.56
H	2	0.25	46.56	3661.94
I	2	0.24	40.88	3463.91
J	2	0.23	38.66	3534.64
A	3	0.26	105.98	6298.82
В	3	0.25	76.58	4988.44
С	3	0.26	54.12	3697.00
D	3	0.26	43.23	3447.36
E	3	0.29	53.71	3519.99

Table A-8 Data Summary - Core H

Layer	Location	Porosity	Permeability (md.)	Mean Pore Throat Size (nM)
F	3	0.25	72.03	4404,42
G	3	0.25	46.68	3436.33
Н	3	0.22	143.92	7654.46
I	3	0.24	41.67	3476.36
J	3	0.22	67.26	4522.37
A	4	0.26	133.59	7033.29
В	4	0.24	37.84	3336.68
С	4	0.26	48.20	3542.50
D	4	0.21	56.54	4365.08
E	4	0.24	70.68	4499.80
F	4	0.24	39.12	3476.36
G	4	0.22	92.57	6454.78
Н	4	0.23	46.59	3613.21
I	4	0.24	40.06	3636.70
J	4	0.34	230.49	6161.84
A	5	0.23	142.96	7717.34
В	5	0.23	56.01	4053.04
С	5	0.25	43.79	3504.41
D	5	0.25	45.48	3483.12
E	5	0.21	66.05	4755.68
F	5	0.24	40.50	3483.59
G	5	0.26	39.05	3238.80
н	5	0.24	40.98	3536.09
I	5	0.23	37.49	3512.62
J	5	0.23	35.46	3420.79

Table A-8 Data Summary - Core H (cont'd)

Layer	Location	Porosity	Permeability (md.)	Mean Pore Throat Size (nM)
A	1	0.18	21.09	2066.90
В	1	0.13	26.59	3145.02
С	1	0.24	50.60	2916.32
D	1	0.16	17.92	1661.97
E	1	0.14	28.58	3300.31
F	1	0.16	24.19	2315.73
G	1	0.17	22.64	2187.84
н	1	0.12	36.73	4048.70
I	1	0.12	6.60	1978.34
J	1	0.18	29.64	2720.91
A	2	0.15	30.98	2834.53
В	2	0.13	19.77	2484.08
c	2	0.12	21.02	3259.05
D	2	0.13	25.08	3311.49
Е	2	0.15	28.91	2903.32
F	2	0.14	2.79	1536.18
G	2	0.16	28,96	2975.76
Н	2	0.13	14.06	1931.54
I	2	0.15	25.09	2896.87
J	2	0.19	66.56	5581.84
A	3	0.18	24.16	2846.26
В	3	0.19	38.59	2630.09
С	3	0.18	31.95	2473.54
D	3	0.16	39.56	3045.80
Е	3	0.12	24.51	2833.88

Table A-9 Data Summary - Core I
Layer	Location	Porosity	Permeability (md.)	Mean Pore Throat Size (nM)
F	3	0.14	21.11	2622.88
G	3	0.12	16.81	2062.54
Н	3	0.15	44.81	2008.07
I	3	0.14	73.45	7713.36
J	3	0.16	26.74	2325.84
A	4	0.14	17.11	3255.79
В	4	0.15	29.52	1999.26
с	4	0.20	35.11	1742.07
D	4	0.17	16.64	2062.54
Е	4	0.15	21.37	2056.24
F	4	0.19	0.1	1925.43
G	4	0.13	29.52	3844.00
Н	4	0.16	36.62	2129.95
I	4	0.08	6.72	1947.00
J	4	0.17	26.34	2835.78
A	5	0.18	22.81	2566.42
В	5	0.16	29.45	2062.54
С	5	0.15	29.10	2751.22
D	5	0.19	10.58	2132.13
Е	5	0.14	28.36	2728.78
F	5	0.16	28.01	2556.01
G	5	0.12	23.08	3648.99
н	5	0.18	22.42	1864.74
I	5	0.14	35.99	4197.72
J	5	0.14	25.86	3003.69

Table A-9 Data Summary - Core I (cont'd)

Layer	Location	Porosity	Permeability (md.)	Mean Pore Throat Size (nM)
A	1	0.11	31.94	3863,24
В	1	0.16	21.76	2323.24
С	1	0.14	30.60	2936.02
D	1	0.15	27.00	2922.86
Ē	1	0.17	20.45	2284.49
F	1	0.20	44.36	2495.16
G	1	0.18	24.35	2195.98
H	1	0.18	0.1	2419.96
I	1	0.17	25.16	2598.01
J	1	0.12	27.66	3968.92
A	2	0.14	22.21	2410.42
В	2	0.16	17.60	1999.27
С	2	0.16	15.24	1934.00
D	2	0.15	31.95	2691.62
Е	2	0.17	22.48	1918.34
F	2	0.16	36.49	2564.81
G	2	0.18	32.72	2995.92
Н	2	0.16	18.55	2479.96
I	2	0.15	28.54	3050.83
J	2	0.14	29.58	3417.12
A	3	0.15	0.1	1893.95
В	3	0.19	16.88	2546.02
С	3	0.19	22.29	2139.94
D	3	0.15	32.15	2955.39
E	3	0.12	24.93	3852.11

Table A-10 Data Summary - Core J

Layer	Location	Porosity	Permeability (md.)	Me∴n Pore Throat Size (nM)
F	3	0.22	55.18	3399.93
G	3	0.15	27.30	2717.53
Н	3	0.21	15.16	1426.12
I	3	0.16	30.01	2445.70
J	3	0.16	30.19	3048.47
A	4	0.14	22.69	2682.73
В	4	0.15	19.81	3167.95
С	4	0.15	297.83	2573.44
D	4	0.11	18.83	2051.72
Е	4	0.14	21.64	1987.07
F	4	0.12	20.45	3109.21
G	4	0.17	21.51	2159.01
Н	4	0.19	36.08	2775.66
1	4	0.14	37.45	4056.11
J	4	0.13	19.29	2713.40
A	5	0.17	15.56	2475.98
B	5	0.14	15.76	2515.52
с	5	0.11	31.42	4371.18
D	5	0.18	31.05	2546.02
E	5	0.18	39.63	3202.98
F	5	0.21	33.41	2454.91
G	5	0.18	19.24	2319.48
Н	5	0.18	21.49	2627.05
I	5	0.14	44.12	4178.50
J	5	0.14	22.68	3060.73

Table A-10 Data Summary - Core J (cont'd)

Appendix B

1. ANOVA Calculations

An important requirement for the use of semi-variograms and autocorrelation functions in the calculation of fractal dimension is that the mean be stationary in a statistical sense. Recall from Chapter IV the assumption that E(Z(x+h)) exists and is only dependent upon h. In practical terms, this assumption may be tested by a comparison of the variance of properties within each layer of the core to the variance between each layer of core (Figure V-3). If the variance within the layers and between the layers are similar then the mean is stationary and the application of semi-variograms and autocorrelation functions to the data set is valid (W. H. Griffin, personal communication).

The comparison of variances between the two groups is readily achieved through the use of a standard one-way analysis of variance (ANOVA) test. Till (1974) provides a good description of this test which calculates the ratio of the variance between layers in the core and variance within layers of the core. This variance ratio is expressed as an F statistic which may be tested by comparing it to the value of F at the 1% confidence level. If $F_{calc} < F_{0.01}$ then the assumption that the mean is stationary is valid. The following calculation is an example of the ANOVA procedure as applied to Core A permeability data contained in Appendix A.

Data Groups	Group A	Group B	Group C	Group D	Group E
(md)	170.2	62.3	30.9	0.1	38.0
	59.2	187.3	41.3	48.0	46.2
	243.2	0.1	37.4	71.0	0.1
	41.7	234.8	63.6	42.0	39.4
	51.5	54.4	118.0	53.1	3.2
Mean	113.1	107.8	58.2	42.8	25.4
	Group F	Group G	Group H	Group I	Group J
	3.7	28.1	396.6	57.5	74.5
	38.9	209.3	53.5	72.5	0.1
	44.4	6885.1	207.5	68.3	170.4
	0.1	59.5	55.3	47.2	38.1
	26.7	46.3	55.2	54.7	36.7
Mean	22.8	1807.1	85.8	60.1	63.9

$$\overline{x} = \frac{1}{10} \sum_{i=A}^{J} \overline{x}_{i} = 238.7$$

$$BSS = n \sum_{i=A}^{J} (\overline{x}_{i} - \overline{x})^{2} = 5(1342011.12) = 6710055.6$$

$$WSS = \sum_{i=A}^{J} \sum_{j=1}^{5} (x_{j} - \overline{x}_{i})^{2} = 12063086.0$$

Variation	Sum of Squares	Degrees of Freedom	Mean Sum of Squares	Ratio (F)
Between Groups	6710055.6	9	745561.73	
Within Groups	12063086.0	40	301577.15	2.472
Total	18773141.6	49		

ANOVA Analysis

 $F_{(0.01;9,40)} = 2.89$

Since 2.472 < 2.89 we accept the hypothesis that the mean is stationary.

This test was repeated for each property in each of the ten cores. In each case the test of the stationary mean was satisfied.

2. Semi-Variogram Plots

The following pages contain the semi-variogram plots used to calculate the D_{var} values shown in Table VII-5. The procedure for the calculation of the slope as h approached zero followed several rules:

- The model fit to the semi-variogram weighted the first two points of the semi-variogram more highly than the rest of the points because the first two lag spacings are based upon the greatest number of data pairs.
- 2. The slope was arbitrarily defined as the slope of the tangent line to the semi-variogram model between h=9.6 inches and h=4.8 inches.
- 3. The slope was determined between h=17.28 and h=4.8 in the following manner:

 $slope = \frac{\Delta y}{\ln 17.28 - \ln 4.8} = \frac{\Delta y}{\ln 3.6}$

4. The slope was always positive.



































Appendix C

1. Calculation of Property Mean and Variance

Table 0-2 Clustrates the calculation procedure for property variance using the lata in Appendix A. This information was correlated with K, and log K. in Table VII-4. The waring ce of the natural log of permeability $\sigma_{in(k)}^2$ is also listed. This parameter was calculated by program ANAL-2E listed in Appendix E after the exclusion of outliers and was used in the calculation of the Dykstra-Parsons permeability variation in Table VI-1.

Note that the mean values of porosity and permeability $\bar{\mu}$ calculated here do not correspond to porosity and permeability values for the entire core shown in Table V-2. There are three reasons for this:

- A maximum of fifty samples are averaged in this calculation procedure while the data in Table V-2 were calculated on a whole core basis.
- 2. Exclusion of the outliers reduces the mean value.
- 3. Different procedures were used in the calculation. Table V-2 porosity values were calculated from a core saturation procedure using *n*-hexane and permeability values were determined by flooding the core with *n*-hexane at four different rates and plotting differential pressures. The permeability

was then calculated from Darcy's law for linear flow. In the case of the data in Appendix A porosity and permeability were both determined by mercury porosimetry methods.

		ū	σ²	$\sigma_{\ln(k)}^2$	DATA
CORE A	OUTLIERS		-	$O_{\ln(k)}$	
POROSITY	A-1, A-3, B-4, C-5	0.138	0.00223		46
PERMEABILITY	G-3	73.11	5917.24	4.207	49
MEAN PORE SIZE		2985.6	3462742		50
CORE B	OUTLIERS	μ	σ²	$\sigma^2_{\ln(k)}$	DATA
POROSITY		0.189	0.00138		50
PERMEABILITY	G-3	82.96	3030.66	0.309	49
MEAN PORE SIZE		5486.15	2607612		50
CORE C	OUTLIERS	μ	σ²	$\sigma^2_{\ln(k)}$	DATA
POROSITY		0.21	0.000386		50
PERMEABILITY	I-2	76.3	1276.1	0.988	49
MEAN PORE SIZE		5385.3	1421689		50
CORE D	OUTLIERS	μ	٥²	$\sigma_{\ln(k)}^2$	DATA
POROSITY		0.172	0.00081		50
PERMEABILITY	D-2	73.52	9057.84	3.343	49
MEAN PORE SIZE		3955.55	3290024		50

Table C-1 Property Mean and Variance Calculations

Table C-1							
Property	Nean	and	Variance	Calculations			
		(C	cont'd)				

			3		
CORE E	OUTLIERS	μ	٥²	$\sigma_{\ln(k)}^2$	DATA
POROSITY		0.243	0.00069		50
PERMEABILITY		103.88	7951.4	1.181	50
MEAN PORE SIZE		5520.4	2176609		50
CORE F	OUTLIERS	μ	σ²	$\sigma_{\ln(k)}^2$	DATA
POROSITY		0.097	0.00256		50
PERMEABILITY	E-1, J-1, I-2, A-4	80.5	18064	5.946	46
MEAN PORE SIZE		6218.44	3800035		50
CORE G	OUTLIERS	ħ	σ²	$\sigma_{\ln(k)}^2$	DATA
POROSITY		0.082	0.00124		50
PERMEABILITY	C-1, D-1, J-1 B-3, H-4, F-5	85.31	15313.22	5.854	44
MEAN PORE SIZE		6645.9	1543871		50
CORE H	OUTLIERS	٦Ļ	σ²	$\sigma_{\ln(k)}^2$	DATA
POROSITY		0.247	0.00086		50
PERMEABILITY		65.92	1306.74	0.185	50
MEAN PORE SIZE		4375.4	1557655		50
CORE I	OUTLIERS	μ	σ²	$\sigma_{\ln(k)}^2$	DATA
POROSITY		0.154	0.00076		50
PERMEABIL, ITY		26.88	171.305	0.879	50
MEAN PORE SIZE		2758.6	1045225		50

Table C-1Property Mean and Variance Calculations(cont'd)

CORE J	OUTLIERS	μ	σ²	$\sigma^2_{\ln(k)}$	DATA
POROSITY		0.158	0.00064		50
PERMEABILITY		31.06	1547.68	1.405	50
MEAN PORE SIZE		2738.5	401720		50

Appendix D

This appendix contains the data used in the calculation of K. using Brigham's lambda factor method for each of the corefloods. The values for v and L used in the calculations come from Tables VII-1 through VII-3. The results of these calculations are also shown in Tables VII-1 through VII-3. Equation II-4 was used in the calculations with K. being substituted for K_i .



1. Sample Calculation

Core J

Rate = 0.0222 cm/s $\lambda_{90} - \lambda_{10} = 1.554$ K. = $(0.02298 \ cm/s)^*(121.92 \ cm)^*\left(\frac{1.554}{3.625}\right)^2 = 0.5149 \ cm^2/s$ Rate = 0.01528 cm/s $\lambda_{90} - \lambda_{10} = 1.359$ K. = $(0.0148 \ cm/s)^*(121.92 \ cm)^*\left(\frac{1.359}{3.625}\right)^2 = 0.2535 \ cm^2/s$ Rate = 0.009722 cm/s $\lambda_{90} - \lambda_{10} = 1.021$ K. = $(0.00955 \ cm/s)^*(121.92 \ cm)^*\left(\frac{1.021}{3.625}\right)^2 = 0.09237 \ cm^2/s$

2. Effluent Concentration Profile and λ Values

Tables D-1 through D-10 contain the effluent concentration profiles as well as the λ values calculated by program EFFPLOT for each coreflood. These λ values were plotted as shown on the previous page and K, was calculated using the same procedure as the sample calculation.

	Rate=0.02222 cm; 5		Rate=0. cm		Rate=0 cm	1
P.V.	% C6H12	λ	% C ₆ H ₁₂	λ	% C ₆ H ₁₂	λ
$\begin{array}{c} 0.050\\ 0.100\\ 0.150\\ 0.200\\ 0.250\\ 0.300\\ 0.350\\ 0.400\\ 0.450\\ 0.550\\ 0.600\\ 0.550\\ 0.600\\ 0.650\\ 0.700\\ 0.650\\ 0.700\\ 0.850\\ 0.900\\ 0.950\\ 1.000\\ 1.000\\ 1.050\\ 1.000\\ 1.550\\ 1.300\\ 1.350\\ 1.400\\ 1.550\\ 1.600\\ 1.650\\ 1.700\\ 1.650\\ 1.750\\ 1.600\\ 1.650\\ 1.750\\ 1.900\\ 1.950\\ 2.000\\ \end{array}$	$\begin{array}{c} 1.786\\ 1.965\\ 1.786\\ 1.965\\ 1.786\\ 1.$	$\begin{array}{c} -4.249\\ -2.846\\ -2.195\\ -1.789\\ -1.500\\ -1.278\\ -1.099\\ -0.949\\ -0.949\\ -0.820\\ -0.707\\ -0.607\\ -0.516\\ -0.434\\ -0.359\\ -0.224\\ -0.163\\ -0.289\\ -0.224\\ -0.163\\ -0.051\\ -0.051\\ 0.000\\ 0.049\\ 0.095\\ 0.140\\ 0.049\\ 0.095\\ 0.140\\ 0.338\\ 0.374\\ 0.442\\ 0.474\\ 0.506\\ 0.537\\ 0.596\\ 0.653\\ 0.680\\ 0.707\end{array}$	$\begin{array}{c} 1.608\\ 2.500\\ 2.500\\ 2.500\\ 2.679\\ 1.786\\ 2.500\\ 2.$	$\begin{array}{c} -4.249\\ -2.846\\ -2.195\\ -1.789\\ -1.500\\ -1.278\\ -1.099\\ -0.949\\ -0.949\\ -0.820\\ -0.707\\ -0.607\\ -0.516\\ -0.434\\ -0.359\\ -0.289\\ -0.224\\ -0.163\\ -0.105\\ -0.051\\ 0.000\\ 0.049\\ 0.095\\ 0.140\\ 0.095\\ 0.140\\ 0.095\\ 0.140\\ 0.095\\ 0.140\\ 0.095\\ 0.140\\ 0.095\\ 0.140\\ 0.095\\ 0.140\\ 0.095\\ 0.140\\ 0.095\\ 0.140\\ 0.095\\ 0.140\\ 0.095\\ 0.165\\ 0.653\\ 0.680\\ 0.707\end{array}$	0.892 1.429 1.429 1.429 1.429 1.250 0.715 0.892 1.608 0.892 2.321 3.571 6.786 10.626 10.000 99.756 100.000	$\begin{array}{c} -4.249\\ -2.846\\ -2.195\\ -1.789\\ -1.500\\ -1.278\\ -1.099\\ -0.949\\ -0.949\\ -0.949\\ -0.949\\ -0.516\\ -0.434\\ -0.359\\ -0.224\\ -0.163\\ -0.229\\ -0.224\\ -0.163\\ -0.289\\ -0.224\\ -0.163\\ -0.263\\ -0.263\\ -0.301\\ 0.095\\ 0.140\\ 0.95\\ 0.140\\ 0.95\\ 0.567\\ 0.596\\ 0.625\\ 0.653\\ 0.653\\ 0.680\\ 0.707\end{array}$

Table D-1 Effluent Concentration Profiles (Core A)

	Rate=0 cm		Rate=0 cm		Rate=0. cm	
P.V.	% C ₆ H ₁₂	λ	% C ₆ H ₁₂	λ	% C ₆ H ₁₂	λ
0.050 0.100 0.150 0.200 0.250 0.300 0.350 0.400 0.550 0.600 0.550 0.650 0.700 0.750 0.800 0.900 1.000 1.000 1.100 1.250 1.300 1.350 1.400 1.550 1.600 1.550 1.600 1.550 1.900 1.950 2.000	$\begin{array}{c} 1.429\\ 3.571\\ 4.465\\ 4.821\\ 4.465\\ 4.821\\ 4.465\\ 4.286\\ 3.579\\ 3.036\\ 1.9659\\ 2.859\\ 7.142\\ 10.781\\ 18.906\\ 2.8589\\ 7.142\\ 10.780\\ 31.965\\ 2.8589\\ 7.142\\ 10.780\\ 31.965\\ 2.8589\\ 7.142\\ 10.780\\ 3.036\\ 5.532\\ 50.000\\ 61.667\\ 71.905\\ 80.294\\ 90.244\\ 92.926\\ 94.878\\ 96.341\\ 96.341\\ 96.341\\ 96.341\\ 99.551\\ $	$\begin{array}{c} -4.249\\ -2.846\\ -2.195\\ -1.789\\ -1.500\\ -1.278\\ -1.099\\ -0.820\\ -0.949\\ -0.820\\ -0.516\\ -0.359\\ -0.224\\ -0.165\\ -0.224\\ -0.165\\ -0.224\\ -0.165\\ -0.224\\ -0.165\\ -0.224\\ -0.359\\ -0.224\\ -0.165\\ -0.224\\ -0.516\\ -0.359\\ -0.226\\ -0.226\\ -0.556\\ -0.653\\ -0.567\\$	2.679 2.679 2.500 1.786 2.142 1.608 1.786 2.500 1.244 3.331 890.366 97.360 98.561 99.512 99.267 99.267 99.267	$\begin{array}{c} -4.249\\ -2.846\\ -2.195\\ -1.789\\ -1.500\\ -1.278\\ -1.099\\ -0.820\\ -0.949\\ -0.820\\ -0.707\\ -0.607\\ -0.516\\ -0.289\\ -0.224\\ -0.163\\ -0.289\\ -0.224\\ -0.163\\ -0.289\\ -0.224\\ -0.163\\ -0.289\\ -0.224\\ -0.516\\ -0.516\\ -0.516\\ -0.516\\ -0.516\\ -0.516\\ -0.516\\ -0.556\\ -0.556\\ -0.567\\ -0.596\\ -0.567\\ -0.596\\$	$\begin{array}{c} 1.250\\ 2.500\\ 1.786\\ 1.965\\ 1.786\\ 2.321\\ 1.786\\ 2.321\\ 1.786\\ 2.509\\ 2.509\\ 2.679\\ 2.679\\ 2.679\\ 2.679\\ 2.679\\ 2.679\\ 2.679\\ 2.679\\ 2.679\\ 2.679\\ 2.679\\ 2.679\\ 2.679\\ 2.682\\ 92.682\\ 92.682\\ 92.926\\ $	$\begin{array}{c} -4.249\\ -2.846\\ -2.195\\ -1.789\\ -1.500\\ -1.278\\ -1.099\\ -0.949\\ -0.949\\ -0.949\\ -0.949\\ -0.516\\ -0.434\\ -0.359\\ -0.224\\ -0.163\\ -0.289\\ -0.224\\ -0.163\\ -0.289\\ -0.224\\ -0.163\\ -0.359\\ -0.224\\ 0.434\\ -0.359\\ -0.224\\ 0.434\\ -0.359\\ -0.224\\ 0.434\\ -0.516\\ -0.051\\ 0.000\\ 0.049\\ 0.095\\ 0.140\\ 0.338\\ 0.374\\ 0.408\\ 0.442\\ 0.474\\ 0.506\\ 0.537\\ 0.567\\ 0.596\\ 0.653\\ 0.680\\ 0.707\end{array}$

Table D-2 Effluent Concentration Profiles (Core B)

	Rate=0 cm		Rate=0 cm		Rate=0. cm/	
P.V.	% C ₆ H ₁₂	λ	% C ₆ H ₁₂	λ	% C ₆ H ₁₂	λ
$\begin{array}{c} 0.050\\ 0.100\\ 0.150\\ 0.200\\ 0.250\\ 0.300\\ 0.350\\ 0.400\\ 0.550\\ 0.650\\ 0.550\\ 0.650\\ 0.550\\ 0.650\\ 0.700\\ 0.850\\ 0.950\\ 0.950\\ 1.000\\ 1.950\\ 1.350\\ 1.350\\ 1.350\\ 1.600\\ 1.550\\ 1.600\\ 1.550\\ 1.600\\ 1.550\\ 1.600\\ 1.550\\ 1.900\\ 1.950\\ 2.000\\ \end{array}$	0.892 1.071 1.965 1.786 2.500 1.786 2.679 4.465 7.142 7.858 7.500 7.142 6.071 5.358 6.786 13.438 33.829 65.208 79.523 87.567 92.926 96.341 97.804 99.756 99.756 99.756 99.756 99.756 99.756 100.000 100.000 100.000 100.000 100.000	$\begin{array}{c} -4.249\\ -2.846\\ -2.195\\ -1.789\\ -1.500\\ -1.278\\ -1.099\\ -0.949\\ -0.949\\ -0.949\\ -0.516\\ -0.434\\ -0.359\\ -0.289\\ -0.224\\ -0.163\\ -0.289\\ -0.224\\ -0.163\\ -0.051\\ -0.051\\ -0.051\\ -0.051\\ 0.095\\ -0.263\\ -0.263\\ -0.263\\ -0.263\\ -0.263\\ -0.263\\ -0.567\\ -0.596\\ -0.537\\ -0.596\\ -0.596\\ -0.537\\ -0.596\\ -0.596\\ -0.537\\ -0.596\\ -0.596\\ -0.537\\ -0.596\\ -0.596\\ -0.537\\ -0.596\\ -0.577\\ -0.596\\ -0.577\\ -0.596\\ -0.577\\ -0.596\\ -0.577\\ -0.596\\ -0.577\\ -0.596\\ -0.577\\ -0.596\\ -0.577\\ -0.596\\ -0.577\\ -0.596\\ -0.577\\ -0.596\\ -0.577\\ -0.596\\ -0.577\\ -0.596\\ -0.577\\ -0.596\\ -0.577\\ -0.596\\ -0.577\\ -0.596\\ -0.577\\ -0.577\\ -0.576\\ -0.577\\ -0.576\\ -0.577\\ -0.576\\ -0.577\\ -0.576\\ -0.577\\ -0.576\\ -0.577\\ -0.576\\ -0.577\\ -0.576\\ -0.577\\ -0.576\\ -0.577\\ -0.576\\ -0.577\\ -0.576\\ -0.577\\ -0.576\\ -0.577\\ -0.576\\ -0.577\\ -0.576\\ -0.577\\ -0.576\\ -0.577\\ -0.576\\ -0.577\\ -0.576\\ -0.577\\ $	$\begin{array}{c} 1.965\\ 1.786\\ 1.786\\ 2.321\\ 1.786\\ 2.500\\ 1.250\\ 1.250\\ 1.250\\ 1.429\\ 1.250\\ 1.786\\ 1.608\\ 2.142\\ 1.608\\ 2.142\\ 1.608\\ 2.142\\ 1.608\\ 2.55\\ 59.750\\ 78.572\\ 90.244\\ 92.926\\ 96.097\\ 97.804\\ 98.781\\ 99.512\\ 99.267\\ 99.756\\ 100.000\\ 99.756\\ 100.000\\ 100$	$\begin{array}{c} -4.249\\ -2.846\\ -2.195\\ -1.789\\ -1.500\\ -1.278\\ -1.099\\ -0.949\\ -0.949\\ -0.820\\ -0.707\\ -0.607\\ -0.516\\ -0.434\\ -0.359\\ -0.289\\ -0.224\\ -0.163\\ -0.105\\ -0.051\\ -0.051\\ -0.051\\ -0.051\\ -0.051\\ -0.051\\ -0.338\\ -0.289\\ -0.224\\ -0.163\\ -0.289\\ -0.224\\ -0.516\\ -0.516\\ -0.557\\ -0.596\\ -0.597\\ -0.596\\ -0.596\\ -0.596\\ -0.597\\ -0.596\\ -0.596\\ -0.596\\ -0.596\\ -0.596\\ -0.597\\ -0.596\\ -0.596\\ -0.596\\ -0.597\\ -0.596\\ -0.596\\ -0.596\\ -0.597\\ -0.596\\ -0.596\\ -0.596\\ -0.597\\ -0.596\\ -0.596\\ -0.597\\ -0.596\\ -0.597\\ -0.596\\ -0.597\\ -0.596\\ -0.597\\ -0.596\\ -0.597\\ -0.596\\ -0.597\\ -0.596\\ -0.597\\ -0.596\\ -0.597\\ -0.596\\ -0.597\\ -0.596\\ -0.597\\ -0.596\\ -0.596\\ -0.597\\ -0.596\\ -0.596\\ -0.597\\ -0.596\\ -0.596\\ -0.597\\ -0.596\\ -0.596\\ -0.597\\ -0.596\\ -0.596\\ -0.597\\ -0.596\\ -0.596\\ -0.597\\ -0.596\\ -0.597\\ -0.596\\ -0.597\\ -0.596\\ -0.597\\ -0.596\\ -0.597\\ -0.596\\ -0.597\\ -0.596\\ -0.597\\ -0.596\\ -0.597\\ -0.596\\ -0.597\\ -0.596\\ -0.597\\ -0.596\\$	1.786 1.429 1.608 1.071 1.071 1.786 0.715 1.786 1	$\begin{array}{c} -4.249\\ -2.846\\ -2.195\\ -1.789\\ -1.500\\ -1.278\\ -1.099\\ -0.949\\ -0.949\\ -0.820\\ -0.707\\ -0.607\\ -0.516\\ -0.434\\ -0.359\\ -0.289\\ -0.224\\ -0.163\\ -0.105\\ -0.051\\ 0.000\\ 0.095\\ 0.140\\ 0.095\\ 0.140\\ 0.095\\ 0.140\\ 0.095\\ 0.301\\ 0.338\\ 0.374\\ 0.408\\ 0.442\\ 0.474\\ 0.506\\ 0.537\\ 0.567\\ 0.596\\ 0.653\\ 0.680\\ 0.707\end{array}$

Table D-3 Effluent Concentration Profiles (Core C)

	Rate=0.02222 cm/s		Rate=0.01528 cm/s		Rate=0.00972 cm/s	
P.V.	% C ₆ H ₁₂	λ	% C ₆ H ₁₂	λ	% C6H12	λ
0.050 0.100 0.200 0.250 0.300 0.350 0.400 0.550 0.600 0.550 0.600 0.650 0.700 0.850 0.900 0.950 1.000 1.250 1.200 1.350 1.400 1.550 1.600 1.550 1.600 1.550 1.600 1.650 1.650 1.900 1.950 2.000	2.679 3.929 4.465 4.465 3.750 2.858 2.679 2.500 1.965 3.358 2.679 2.500 1.398 8.036 11.563 16.094 23.333 192 41.778 68.124 76.667 82.4863 90.4863 93.901 95.853 97.560 93.901 95.853 97.560 97.560 98.293 98.293 99.267	$\begin{array}{c} -4.249\\ -2.846\\ -2.195\\ -1.789\\ -1.500\\ -1.278\\ -1.099\\ -0.949\\ -0.949\\ -0.516\\ -0.434\\ -0.359\\ -0.224\\ -0.163\\ -0.289\\ -0.224\\ -0.163\\ -0.105\\ -0.051\\ 0.000\\ 0.049\\ -0.263\\ -0.263\\ -0.263\\ -0.263\\ -0.557\\ -0.557\\ -0.596\\ -0.537\\ -0.596\\ -0.596\\ -0.596\\ -0.537\\ -0.596\\ -0.596\\ -0.553\\ -0.596\\ -0.596\\ -0.553\\ -0.596\\ -0.553\\ -0.596\\ -0.553\\ -0.596\\ -0.553\\ -0.596\\ -0.553\\ -0.596\\ -0.553\\ -0.596\\ -0.553\\ -0.596\\ -0.553\\ -0.596\\ -0.553\\ -0.596\\ -0.553\\ -0.596\\ -0.553\\ -0.596\\ -0.553\\ -0.555\\ -0.555\\ -$	0.892 3.036 2.142 1.608 1.786 1.965 1.608 2.142 1.786 0.892 1.608 2.142 1.786 0.892 1.608 2.142 1.786 0.892 1.608 2.571 7.142 10.626 23.333 32.127 39.574 50.749 61.458 67.292 74.049 78.810 85.946 88.918 93.415 93.459 94.8781 95.745 93.659 94.8781 95.7561 99.756 100.000 100.000 100.000	$\begin{array}{c} -4.249\\ -2.846\\ -2.195\\ -1.789\\ -1.500\\ -1.278\\ -1.099\\ -0.949\\ -0.949\\ -0.820\\ -0.707\\ -0.607\\ -0.516\\ -0.434\\ -0.359\\ -0.224\\ -0.163\\ -0.289\\ -0.224\\ -0.163\\ -0.051\\ -0.051\\ -0.051\\ -0.051\\ -0.051\\ -0.051\\ -0.359\\ -0.224\\ -0.163\\ -0.516\\ -0.516\\ -0.516\\ -0.556\\ -0.556\\ -0.557\\ -0.567\\ -0.567\\ -0.567\\ -0.567\\ -0.596\\ -0.597\\ -0.596\\ -0.596\\ -0.596\\ -0.597\\ -0.596\\$	0.536 1.786 1.965 1.786 1.965 1.786 2.321 1.608 1.608 1.965 1.786 3.571 4.821 7.858 1.407 16.094 24.889 34.042 43.999 55.000 64.166 69.791 76.667 83.513 89.108 91.707 94.878 96.341 96.341 97.560 98.293 99.025 98.781 99.025 99.756 99.756	$\begin{array}{c} -4.249\\ -2.846\\ -2.195\\ -1.789\\ -1.500\\ -1.278\\ -1.099\\ -0.949\\ -0.949\\ -0.820\\ -0.707\\ -0.607\\ -0.516\\ -0.434\\ -0.359\\ -0.224\\ -0.163\\ -0.289\\ -0.224\\ -0.163\\ -0.289\\ -0.224\\ -0.163\\ -0.359\\ -0.224\\ -0.163\\ -0.556\\ -0.557\\ 0.567\\ 0.596\\ 0.625\\ 0.653\\ 0.680\\ 0.707\end{array}$

Table D-4 Effluent Concentration Profiles (Core D)

	Rate=0.02222 cm/s		Rate=0.01528 cm/s		Rate=0.00972 cm/s	
P.V.	% C ₆ H ₁₂		% C ₆ H ₁₂	λ	% C ₆ H ₁₂	λ
0.050 0.100 0.150 0.200 0.250 0.300 0.350 0.400 0.450 0.5500 0.600 0.650 0.650 0.700 0.800 0.900 0.950 1.000 1.050 1.000 1.250 1.300 1.350 1.400 1.550 1.650 1.650 1.650 1.650 1.900 1.900 1.950 1.900 1.900 1.950 1.900	$\begin{array}{c} 1.786\\ 2.142\\ 2.679\\ 2.679\\ 2.679\\ 2.500\\ 1.786\\ 2.500\\ 1.786\\ 2.500\\ 1.786\\ 2.679\\ 6.429\\ 10.313\\ 9.821\\ 8.036\\ 8.215\\ 27.777\\ 69.167\\ 78.095\\ 89.459\\ 94.634\\ 97.318\\ 97.804\\ 96.341\\ 97.318\\ 97.804\\ 98.781\\ 99.025\\ 99.756\\ 100.000\\ 100.000\\ 100.000\\ 100.000\\ 100.000\\ 100.000\\ 99.756\\ 100.000\\ 99.756\end{array}$	$\begin{array}{c} -4.249\\ -2.846\\ -2.195\\ -1.789\\ -1.500\\ -1.278\\ -1.099\\ -0.949\\ -0.820\\ -0.707\\ -0.607\\ -0.516\\ -0.434\\ -0.359\\ -0.289\\ -0.224\\ -0.163\\ -0.105\\ -0.051\\ 0.000\\ 0.049\\ 0.095\\ 0.140\\ 0.224\\ 0.263\\ 0.301\\ 0.338\\ 0.374\\ 0.408\\ 0.474\\ 0.506\\ 0.537\\ 0.567\\ 0.596\\ 0.653\\ 0.653\\ 0.653\\ 0.653\\ 0.707\end{array}$	0.000 0.892 1.071 1.250 1.250 1.250 1.250 1.786 1.071 0.892 1.429 2.679 3.571 2.679 2.858 8.036 40.888 74.049 93.901 94.8781 98.781 99.267 99.267 99.267 99.267 99.267 99.267 99.267 99.267 99.756 100.000 100.000 100.000 99.756 100.000 99.756	$\begin{array}{c} -4.249\\ -2.846\\ -2.195\\ -1.789\\ -1.500\\ -1.278\\ -1.099\\ -0.949\\ -0.949\\ -0.516\\ -0.434\\ -0.359\\ -0.224\\ -0.163\\ -0.229\\ -0.224\\ -0.163\\ -0.051\\ 0.000\\ 0.049\\ -0.051\\ 0.000\\ 0.049\\ 0.095\\ 0.140\\ 0.338\\ 0.374\\ 0.408\\ 0.408\\ 0.442\\ 0.506\\ 0.537\\ 0.596\\ 0.653\\ 0.653\\ 0.680\\ 0.707\end{array}$	100.000 100.000 100.000	

Table D-5 Effluent Concentration Profiles (Core E)

Rate=0.02222 cm/s		Rate=0.01528 cm/s		Rate=0.00972 cm/s		
P.V.	% C ₆ H ₁₂	λ	% C ₆ H ₁₂	λ	% C ₆ H ₁₂	λ
0.050 0.100 0.200 0.250 0.300 0.350 0.400 0.550 0.600 0.550 0.650 0.700 0.850 0.900 1.000 1.050 1.250 1.300 1.400 1.550 1.550 1.600 1.550 1.600 1.550 1.600 1.550 1.600 1.900 1.950 2.000	0.000 0.358 0.000 0.892 0.892 0.892 0.715 1.429 1.608 3.392 5.715 9.642 15.625 35.319 65.208 83.513 96.097 96.341 97.560 98.781 98.781 99.267 98.781 99.267 98.781 99.512 99.756 100.000 98.781 99.512 99.756	$\begin{array}{c} -4.249\\ -2.846\\ -2.195\\ -1.789\\ -1.500\\ -1.278\\ -1.099\\ -0.949\\ -0.820\\ -0.707\\ -0.607\\ -0.516\\ -0.434\\ -0.359\\ -0.224\\ -0.163\\ -0.2289\\ -0.224\\ -0.163\\ -0.000\\ 0.049\\ -0.224\\ -0.163\\ -0.224\\ -0.163\\ -0.224\\ -0.516\\ -0.516\\ -0.537\\ -0.506\\ 0.537\\ 0.596\\ -0.596\\ -0.537\\ 0.596\\ -0.596\\ -0.537\\ -0.596\\ -0.596\\ -0.596\\ -0.537\\ -0.596\\ -0.596\\ -0.537\\ -0.596\\ -0.596\\ -0.537\\ -0.596\\ -0.596\\ -0.537\\ -0.596\\ -0.596\\ -0.537\\ -0.596\\ -0.596\\ -0.537\\ -0.596\\ -0.596\\ -0.537\\ -0.596\\ -0.596\\ -0.597\\ -0.596\\ -0.596\\ -0.597\\ -0.596\\ -0.597\\ -0.596\\ -0.597\\ -0.596\\ -0.597\\ -0.596\\ -0.597\\ -0.596\\ -0.597\\ -0.596\\ -0.597\\ -0.596\\ -0.597\\ -0.596\\ -0.597\\ -0.596\\ -0.596\\ -0.597\\ -0.596\\ -0.596\\ -0.596\\ -0.597\\ -0.596\\ -0$	0.715 1.071 1.608 1.250 0.892 1.429 0.358 1.071 0.179 0.892 1.429 0.892 1.429 0.892 1.429 1.429 0.892 1.429 1.429 0.892 1.429 1.429 0.892 1.429 0.8537 98.537 98.293 99.025 99.267	$\begin{array}{c} -4.249\\ -2.846\\ -2.195\\ -1.789\\ -1.500\\ -1.278\\ -1.099\\ -0.820\\ -0.707\\ -0.607\\ -0.516\\ -0.434\\ -0.359\\ -0.289\\ -0.224\\ -0.163\\ -0.289\\ -0.224\\ -0.163\\ -0.051\\ 0.000\\ 0.095\\ 0.140\\ 0.095\\ 0.140\\ 0.095\\ 0.140\\ 0.338\\ 0.224\\ 0.474\\ 0.506\\ 0.537\\ 0.596\\ 0.596\\ 0.653\\ 0.596\\ 0.653\\ 0.596\\ 0.653\\ 0.596\\ 0.653\\ 0.596\\ 0.655\\ 0.653\\ 0.596\\ 0.655\\ 0.596\\ 0.707\\ 0.596\\ 0.707\\ 0.596\\ 0.596\\ 0.707\\ 0.596\\ 0.707\\ 0.596\\ 0.596\\ 0.707\\ 0.596\\ 0.655\\ 0.55\\ 0.$	2.500 1.786 2.321 1.429 1.786 1.071 1.071 1.250 1.429 1.250 1.429 1.250 1.608 1.786 1.786 2.500 2.679 5.358 15.782 42.888 68.959 82.972 91.463 94.145 96.097 96.585 97.560 98.781 99.025 99.025 99.267 99.267 99.267 99.756 100.000 99.756 100.000 100.000 100.000	$\begin{array}{c} -4.249\\ -2.846\\ -2.195\\ -1.789\\ -1.500\\ -1.278\\ -1.099\\ -0.820\\ -0.949\\ -0.820\\ -0.707\\ -0.607\\ -0.516\\ -0.434\\ -0.359\\ -0.289\\ -0.224\\ -0.163\\ -0.289\\ -0.224\\ -0.163\\ -0.051\\ -0.051\\ -0.051\\ -0.051\\ -0.051\\ -0.051\\ -0.289\\ -0.224\\ -0.163\\ -0.289\\ -0.224\\ -0.506\\ -0.596\\$

Table D-6 Effluent Concentration Profiles (Core F)

	Rate=0.02222 cm/s		Rate=0.01528 cm/s		Rate=0.00972 cm/s	
P.V.	% C ₆ H ₁₂	λ	% C ₆ H ₁₂	λ	% C ₆ H ₁₂	λ
0.050 0.100 0.150 0.200 0.250 0.300 0.400 0.550 0.600 0.650 0.650 0.600 0.700 0.850 0.900 1.000 1.000 1.100 1.250 1.250 1.300 1.400 1.550 1.400 1.550 1.600 1.550 1.600 1.550 1.600 1.900 1.900 1.900 1.950 1.900 1.950 1.900 1.950 1.900 1.950 1.900 1.950 1.900 1.950 1.900 1.950 1.900 1.950 1.900 1.950 1.900 1.950 1.900 1.950 1.900 1.950 1.950 1.900 1.950 1.950 1.900 1.950 1.950 1.900 1.950 1.950 1.900 1.950 1	0.715 0.358 1.071 1.250 0.358 0.536 0.536 0.179 2.500 3.215 4.108 5.179 26.444 37.447 49.777 53.999 64.166 69.167 71.905 81.892 83.513 84.595 86.755 88.918 91.463 91.952 91.952 92.438 92.682 93.171 93.415 93.415	$\begin{array}{c} -4.249\\ -2.846\\ -2.195\\ -1.789\\ -1.500\\ -1.278\\ -1.099\\ -0.949\\ -0.820\\ -0.707\\ -0.607\\ -0.516\\ -0.434\\ -0.359\\ -0.289\\ -0.224\\ -0.163\\ -0.289\\ -0.224\\ -0.163\\ -0.051\\ -0.051\\ 0.000\\ 0.049\\ 0.095\\ 0.140\\ 0.2263\\ 0.301\\ 0.338\\ 0.374\\ 0.408\\ 0.474\\ 0.506\\ 0.537\\ 0.567\\ 0.596\\ 0.653\\ 0.680\\ 0.707\end{array}$	0.358 0.358 0.536 0.715 1.071 0.715 0.536 0.715 1.2858 3.392 4.108 11.407 17.969 29.332 32.979 38.297 38.297 38.297 38.297 52.749 52.749 52.749 58.500 67.292 69.374 74.523 77.382 80.541 82.162 83.783 86.216 88.108 89.459 90.244 90.975 90.730 90.975 91.463 91.952 92.194 92.438	$\begin{array}{c} -4.249\\ -2.846\\ -2.195\\ -1.789\\ -1.500\\ -1.278\\ -1.099\\ -0.949\\ -0.820\\ -0.707\\ -0.607\\ -0.607\\ -0.516\\ -0.434\\ -0.359\\ -0.229\\ -0.224\\ -0.163\\ -0.105\\ -0.051\\ -0.051\\ -0.051\\ -0.051\\ -0.051\\ 0.000\\ 0.049\\ 0.095\\ 0.140\\ 0.263\\ 0.301\\ 0.338\\ 0.374\\ 0.408\\ 0.408\\ 0.442\\ 0.474\\ 0.506\\ 0.537\\ 0.567\\ 0.596\\ 0.625\\ 0.653\\ 0.680\\ 0.707\end{array}$	0.358 0.536 0.715 1.071 0.536 0.358 1.071 1.608 1.786 2.500 2.679 3.215 3.750 7.142 18.437 39.787 27.112 37.660 41.333 50.749 53.250 62.499 64.792 70.238 73.333 77.856 79.523 81.080 83.242 85.134 87.026 88.918 90.244 90.244 90.486 90.730 90.975	$\begin{array}{c} -4.249\\ -2.846\\ -2.195\\ -1.789\\ -1.500\\ -1.278\\ -1.099\\ -0.949\\ -0.949\\ -0.820\\ -0.707\\ -0.516\\ -0.434\\ -0.359\\ -0.289\\ -0.224\\ -0.163\\ -0.289\\ -0.224\\ -0.163\\ -0.051\\ 0.000\\ 0.095\\ 0.140\\ 0.095\\ 0.140\\ 0.338\\ 0.374\\ 0.223\\ 0.301\\ 0.338\\ 0.374\\ 0.408\\ 0.442\\ 0.474\\ 0.506\\ 0.537\\ 0.596\\ 0.653\\ 0.680\\ 0.707\end{array}$

Table D-7 Effluent Concentration Profiles (Core G)
	Rate=0.02222 cm/s		Rate=0.01528 cm/s		Rate=0.00972 cm/s	
P.V.	% C ₆ H ₁₂	λ	% C ₆ H ₁₂	λ	% C ₆ H ₁₂	λ
0.050 0.100 0.200 0.250 0.300 0.350 0.400 0.450 0.550 0.600 0.550 0.600 0.700 0.800 0.900 0.900 1.000 1.000 1.250 1.300 1.400 1.550 1.400 1.550 1.600 1.550 1.600 1.550 1.600 1.550 1.900 2.000	0.000 0.892 1.071 0.892 1.786 0.892 1.071 1.608 0.358 0.358 0.892 1.429 2.142 1.786 1.965 1.429 2.142 1.965 1.608 92.682 94.878 97.560 97.804 97.560 97.804 97.560 97.804 97.560 97.560 97.804 98.537 98.537 98.537 98.537 99.267 98.537 99.267 99.267 99.267 99.267 99.267 99.512 99.267 100.000 99.756 100.000 100.000 99.756 100.000	$\begin{array}{c} -4.249\\ -2.846\\ -2.195\\ -1.789\\ -1.500\\ -1.278\\ -1.099\\ -0.949\\ -0.949\\ -0.820\\ -0.707\\ -0.607\\ -0.516\\ -0.4359\\ -0.224\\ -0.163\\ -0.224\\ -0.163\\ -0.051\\ -0.051\\ -0.051\\ -0.051\\ -0.051\\ -0.051\\ -0.516\\ -0.516\\ -0.516\\ -0.556\\ -0.557\\ -0.567\\ -0.537\\ -0.567\\ -0.557\\ -0.557\\ -0.556\\ -0.557\\ -0.556\\ -0.557\\ -0.556\\ -0.557\\ -0.556\\ -0.557\\ -0.556\\ -0.625\\ -0.653\\ -0.680\\ -0.707\\ -0.577\\ -0.568\\ -0.557\\ -0.568\\ -0.557\\ -0.568\\ -0.557\\ -0.568\\ -0.557\\ -0.568\\ -0.557\\ -0.568\\ -0.557\\ -0.568\\ -0.557\\ -0.568\\ -0.557\\ -0.568\\ -0.557\\ -0.556\\ -0.688\\ -0.707\\ -0.557\\ -0.568\\ -0.557\\ -0.568\\ -0.557\\ -0.568\\ -0.557\\ -0.568\\ -0.557\\ -0.568\\ -0.557\\ -0.556\\ -0.688\\ -0.707\\ -0.557\\ -0.568\\ -0.557\\ -0.568\\ -0.557\\ -0.557\\ -0.556\\ -0.557\\ -0.557\\ -0.556\\ -0.557\\ -0.557\\ -0.556\\ -0.557\\ -0.557\\ -0.556\\ -0.557\\ -0.556\\ -0.557\\ -0.557\\ -0.556\\ -0.557\\ -0.557\\ -0.556\\ -0.557\\ -0.556\\ -0.557\\ -0.557\\ -0.557\\ -0.556\\ -0.557$	0.892 0.715 1.071 0.892 0.715 1.071 0.892 0.358 0.358 0.715 0.536 1.071 5.358 7.500 8.036 6.965 10.781 30.213 61.667 87.830 91.707 93.901 95.366	$\begin{array}{c} -4.249\\ -2.846\\ -2.195\\ -1.789\\ -1.500\\ -1.278\\ -1.099\\ -0.949\\ -0.949\\ -0.820\\ -0.707\\ -0.607\\ -0.516\\ -0.434\\ -0.359\\ -0.224\\ -0.163\\ -0.289\\ -0.224\\ -0.163\\ -0.051\\ -0.051\\ -0.051\\ -0.051\\ -0.051\\ -0.051\\ -0.599\\ -0.224\\ -0.163\\ -0.599\\ -0.224\\ -0.163\\ -0.599\\ -0.224\\ -0.567\\ -0.596\\$	0.892 1.429 1.250 0.892 0.892 0.715 0.715 0.715 0.92 0.715 0.892 0.715 0.892 0.715 0.892 0.715 0.892 0.715 0.892 0.715 0.992 1.429 10.000 43.999 75.951 88.918 92.438 95.122 96.8293 98.048 98.781 99.025 99.512 100.000 100.000 99.756	$\begin{array}{c} -4.249\\ -2.846\\ -2.195\\ -1.789\\ -1.500\\ -1.278\\ -1.099\\ -0.949\\ -0.820\\ -0.707\\ -0.607\\ -0.516\\ -0.434\\ -0.359\\ -0.289\\ -0.224\\ -0.163\\ -0.289\\ -0.224\\ -0.163\\ -0.051\\ 0.000\\ 0.049\\ -0.051\\ 0.009\\ 0.095\\ 0.140\\ 0.049\\ 0.095\\ 0.140\\ 0.338\\ 0.374\\ 0.408\\ 0.442\\ 0.506\\ 0.537\\ 0.567\\ 0.567\\ 0.567\\ 0.596\\ 0.625\\ 0.653\\ 0.680\\ 0.707\end{array}$

Table D-8 Effluent Concentration Profiles (Core H)

	Rate=0.02222 cm/s		Rate=0.01528 cm/s		Rate=0.00972 CE/S	
P.V.	% C ₆ H ₁₂	λ	% C ₆ H ₁₂	λ	% C ₆ H ₁₂	λ
0.050 0.100 0.200 0.250 0.300 0.400 0.450 0.550 0.600 0.650 0.650 0.700 0.850 0.900 0.950 1.000 1.000 1.250 1.300 1.350 1.400 1.550 1.600 1.550 1.600 1.750 1.600 1.900 1.950 2.000	0.358 0.715 0.715 0.892 1.608 2.858 3.036 3.215 5.000 6.250 7.321 15.001 19.844 23.778 34.681 36.382 56.750 66.875 73.572 77.143 79.523 83.242 85.405 86.216 87.026 87.3659 94.389 94.878 95.366 95.853	$\begin{array}{c} -4.249\\ -2.846\\ -2.195\\ -1.789\\ -1.500\\ -1.278\\ -1.099\\ -0.949\\ -0.820\\ -0.707\\ -0.516\\ -0.359\\ -0.224\\ -0.163\\ -0.289\\ -0.224\\ -0.163\\ -0.289\\ -0.224\\ -0.163\\ -0.359\\ -0.224\\ -0.163\\ -0.359\\ -0.224\\ -0.163\\ -0.516\\ -0.516\\ -0.516\\ -0.551\\ -0.556\\ -0.625\\ -0.596\\$	$0.358 \\ 0.358 \\ 0.892 \\ 1.608 \\ 1.786 \\ 1.608 \\ 1.608 \\ 2.679 \\ 3.571 \\ 5.358 \\ 7.858 \\ 15.786 \\ 15.782 \\ 20.223 \\ 15.782 \\ 23.112 \\ 23.112 \\ 24.499 \\ 66.875 \\ 71.190 \\ 79.523 \\ 83.513 \\ 84.863 \\ 85.675 \\ 89.459 \\ 90.000 \\ 91.219 \\ 92.926 \\ 93.659 \\ 94.389 \\ 95.366 \\ 95.$	$\begin{array}{c} -4.249\\ -2.846\\ -2.195\\ -1.789\\ -1.500\\ -1.278\\ -1.099\\ -0.949\\ -0.820\\ -0.707\\ -0.607\\ -0.516\\ -0.434\\ -0.359\\ -0.289\\ -0.224\\ -0.163\\ -0.289\\ -0.224\\ -0.163\\ -0.051\\ -0.051\\ 0.000\\ 0.095\\ 0.140\\ 0.095\\ 0.140\\ 0.338\\ 0.374\\ 0.442\\ 0.474\\ 0.506\\ 0.537\\ 0.567\\ 0.596\\ 0.625\\ 0.680\\ 0.707\end{array}$	0.358 0.358 0.536 1.608 1.786 2.142 2.679 3.215 4.286 5.358 9.465 13.907 16.562 19.844 21.555 24.001 31.703 40.223 48.000 56.500 61.041 65.625 71.190 77.143 79.523 82.162 83.242 84.3255 88.918 90.000 90.975 91.463 92.194 92.438 93.659 93.901 94.389 94.878	$\begin{array}{c} -4.249\\ -2.846\\ -2.195\\ -1.789\\ -1.500\\ -1.278\\ -1.099\\ -0.949\\ -0.949\\ -0.820\\ -0.707\\ -0.607\\ -0.516\\ -0.434\\ -0.359\\ -0.289\\ -0.556\\ -0.557\\ -0.596\\ 0.625\\ 0.625\\ 0.625\\ 0.625\\ 0.680\\ 0.707\\ \end{array}$

Table D-9 Effluent Concentration Profiles (Core I)

	Rate=0.02222 cm/s		Rate=0.01528 cm/s		Rate=0.00972 cm/s	
P.V.	% C ₆ H ₁₂	λ	% C ₆ H ₁₂	λ	% C ₆ H ₁₂	λ
$\begin{array}{c} 0.050\\ 0.100\\ 0.100\\ 0.200\\ 0.200\\ 0.250\\ 0.300\\ 0.400\\ 0.450\\ 0.550\\ 0.550\\ 0.600\\ 0.550\\ 0.600\\ 0.550\\ 0.600\\ 0.650\\ 0.700\\ 0.850\\ 0.900\\ 0.950\\ 1.000\\ 1.000\\ 1.050\\ 1.000\\ 1.050\\ 1.000\\ 1.550\\ 1.500\\ 1.550\\ 1.550\\ 1.600\\ 1.550\\ 1.600\\ 1.550\\ 1.600\\ 1.950\\ 2.000\\ \end{array}$	0.536 0.715 1.071 0.892 1.965 3.215 5.179 6.608 6.429 7.500 15.312 19.687 23.778 36.3892 36.3892 36.3892 56.500 67.292 79.049 83.242 86.216 86.755 86.292 79.049 83.242 86.216 86.755 88.379 90.486 90.730 90.730 90.730 91.463 94.155 95.608 96.097 96.585	$\begin{array}{c} -4.249\\ -2.846\\ -2.846\\ -2.895\\ -1.99\\ -1.999\\ -1.999\\ -0.820\\ -0.707\\ -0.607\\ -0.516\\ -0.359\\ -0.289\\ $	0.358 0.358 0.358 0.358 0.892 1.071 1.786 1.786 2.858 5.358 8.036 10.626 15.782 41.778 23.332 33.616 53.250 62.251 71.190 75.951 79.523 81.080 83.783 84.595 85.946 87.296 87.296 89.188 90.730 90.9752 91.9524 171 93.171 93.901 94.389 94.145 95.122 96.341	$\begin{array}{c} -4.249\\ -2.846\\ -2.195\\ -1.789\\ -1.500\\ -1.278\\ -1.099\\ -0.949\\ -0.949\\ -0.949\\ -0.516\\ -0.434\\ -0.359\\ -0.224\\ -0.163\\ -0.224\\ -0.163\\ -0.224\\ -0.163\\ -0.224\\ -0.163\\ -0.224\\ -0.163\\ -0.224\\ -0.359\\ -0.224\\ -0.516\\ -0.359\\ -0.224\\ -0.516\\ -0.516\\ -0.557\\ -0.556\\ -0.653\\ -0.567\\$	0.179 0.358 0.536 1.429 1.608 1.429 1.786 3.215 5.358 9.286 14.531 16.406 20.000 21.778 23.778 31.489 39.787 47.777 56.249 61.041 65.416 70.477 76.667 79.762 81.621 83.242 84.863 86.755 88.918 91.952 92.194 91.463 91.952 92.194 92.194 92.194 92.194 93.415 93.901 94.389 94.145 94.878	$\begin{array}{c} -4.249\\ -2.846\\ -2.195\\ -1.789\\ -1.500\\ -1.278\\ -1.099\\ -0.949\\ -0.820\\ -0.707\\ -0.607\\ -0.516\\ -0.434\\ -0.359\\ -0.289\\ -0.224\\ -0.163\\ -0.289\\ -0.224\\ -0.163\\ -0.289\\ -0.224\\ -0.163\\ -0.359\\ -0.224\\ -0.163\\ -0.359\\ -0.224\\ -0.516\\ -0.516\\ -0.551\\ 0.000\\ 0.049\\ 0.095\\ 0.140\\ 0.338\\ 0.374\\ 0.408\\ 0.442\\ 0.506\\ 0.537\\ 0.567\\ 0.596\\ 0.625\\ 0.680\\ 0.707\\ \end{array}$

Table D-10 Effluent Concentration Profiles (Core J)

Appendix E

1. Program EFFPLOT

Sample Data File:

The following is a complete data file used for program EFFPLOT.



```
*
      PROGRAM EFFPLOT
                                         *
*
* THIS PROGRAM APPLIES THE EXPERIMENTAL
                                         *
* CALIBRATION CURVE TO THE RAW EFFLUENT
                                         *
* CONCENTRATION DATA IN ORDER TO OBTAIN
                                         *
* A PROFILE OF CYCLOHEXANE CONCENTRATION
                                         *
* VS PORE VOLUMES OF CYCLOHEXANE INJECTED *
*
    THIS PROFILE IS THEN USED TO
                                         *
* CALCULATE VALUES FOR HEXANE RECOVERY
* AFTER INJECTION OF 1.0, 1.5 AND 2.0 PV
                                         *
                                          4
* OF CYCLOHEXANE.
REAL A(41), B(11), C(41), D(41), PV(41), U(41), NUMB
     REAL AAREA1, AAREA2, AAREA3
     INTEGER I, J, K
     READ(5, *) (B(J), J=1, 11)
     READ(5, *) (PV(I), I=1, 41)
     READ(5,*) (A(I), I=1,41)
     DO 10 I=1,41
       IF (PV(I).GT.0.0) THEN
       U(I) = ((PV(I) - 1.0) / SQRT(PV(I)))
       ENDIF
       DO 20 J=1,10
        IF (A(I).LE.B(J)) THEN
        C(I) = 0.0
        D(I) = C(I) / 100.0
        GO TO 10
       ELSEIF (A(I).GT.B(J).AND.A(I).LE.B(J+1)) THEN
       NUMB=A(I)-B(J)
       RATIO=NUMB/(B(J+1)-B(J))
       C(I) = RATIO * 10.0 + (10.0 * (J-1))
       D(I) = C(I) / 100.0
       GO TO 10
         ELSEIF (A(I).GT.B(11)) THEN
          C(I) = 100.0
          D(I) = C(I) / 100.0
          GO TO 10
         ELSE
         GO TO 20
         ENDIF
 20
        CONTINUE
 10
      CONTIN
```

```
WRITE(6,4)
     FORMAT ('1', 'EFFLUENT CONCENTRATION PROFILE')
 4
     WRITE(6,5)
     FORMAT ('CORE J - RATE=116.20 CM3/HR (RUN #3)'/)
5
     WRITE(6,6)
     FORMAT('PV',8X,'PERCENT C6H12',8X,'LAMBDA'/)
 6
     DO 30 K=1,41
      PRINT 25, PV(K), C(K), U(K)
     CONTINUE
 30
     FORMAT(' ', F6.3, 6X, F8.3, 7X, F6.3)
 25
*********************
* CALCULATE AREAS UNDER
                           *
                           *
* EFFLUENT CONCENTRATION
                           *
* PROFILES USING SIMPSON'S
                           +
* 1/3 RULE
********
*****
* AREA TO 1.0 PV *
*****
AAREA1=C(1)+4.0*C(2)+2.0*C(3)+4.0*C(4)+2.0*C(5)+4.0*C(6)
+2.0*C(7)+4.0*C(8)+2.0*C(9)+4.0*C(10)+2.0*C(11)+4.0*C(12)
+2.0*C(13)+4.0*C(14)+2.0*C(15)+4.0*C(16)+2.0*C(17)
         +4.0*C(18)+2.0*C(19)+4.0*C(20)
     +
      AFREA1=AAREA1+C(21)
************
* AREA TO 1.5 PV *
******
     AAREA2=AAREA1+2.0*C(21)+4.0*C(22)+2.0*C(23)+4.0*C(24)
         +2.0*C(25)+4.0*C(26)+2.0*C(27)+4.0*C(28)
     +
         +2.0*C(29)+4.0*C(30)
      AFREA2=AAREA2+C(31)
*****
* AREA TO 2.0 PV *
*********
```

```
AAREA3 = AAREA2 + 2.0 \times C(31) + 4.0 \times C(32) + 2.0 \times C(33) + 4.0 \times C(34)
          +2.0*C(35)+4.0*C(36)+2.0*C(37)+4.0*C(38)
     +
          +2.0*C(39)+4.0*C(40)
     +
      AFREA3=AAREA3+C(41)
************************
* CALCULATE % HEXANE RECOVERY *
*******
      HEX1=100.0-(0.05/3.0*AFREA1)
      HEX2 = 150.0 - (0.05/3.0 \times AFREA2)
      HEX3=200.0-(0.05/3.^*AFREA3)
      IF (HEX2 .GT. 100.000) HEX2=100.00
      IF (HEX3 .GT. 100.000) HEX3=100.00
      WRITE(6,7) HEX1
      FORMAT(/'HEXANE RECOVERY @ 1.0 PV = ',F8.3,'%')
 7
      WRITE(6,8) HEX2
      FORMAT ('HEXANE RECOVERY @ 1.5 PV = ', F8.3, '%')
 8
      WRITE(6,9) HEX3
      FORMAT ('HEXANE RECOVERY @ 2.0 PV = ', F8.3, '%')
 9
      STOP
      END
```

```
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```

2. Program ANALYZE

Sample Data File:

The following is a sample of the data input format required for program ANALYZE. The data shown is for the first 3 samples only. Additional data may be added up to a maximum of 50 samples per computer run. Processing of greater than 50 samples per computer run will require modification of the source code. The data which starts with 1.000 in each line corresponds to HG injection readings at the following pressures: 20, 30, 40, 50, 60, 80, 100, 125, 150, 175, 200, 300, 400, 500, 600, 700, 800, 900 and 1000 psi.



PROGRAM ANALYZ ANALYZE × * THIS PROGRAM PERFORMS DATA ANALYSIS UPON * * * + CAPILLARY PRESSURE DATA OBTAINED ON A CORE BY CORE BASIS. THE MAIN PORTION OF THE PROGRAM ÷ READS IN THE MERCURY INJECTION DATA AND ÷ + SUBTRACTS THE APPROPRIATE CALIBRATION CURVE. * * THE CORRECTED DATA IS THEN PASSED ON TO OTHER * + ÷ SUBROUTINES WHERE FURTHER DATA PROCESSING ÷ OCCURS. * * SUBROUTINES: * - CALCULATES THE AREA UNDER THE * AREA * CAPILLARY PRESSURE CURVE. ALSO * CALCULATES POROSITY, PERMEABILITY AND * THE MEAN PORE THROAT DIAMETER FOR EACH * SAMPLE. * - CALCULATES EXPERIMENTAL SEMI-VARIOGRAM * VARIO FOR THE CORE BASED UPON THE LOGARITHM * LOGARITHM OF THE DATA VALUES + * CALCULATED BY SUBROUTINE AREA. ALSO * CALCULATES THE AUTOCORRELATION FUNCTION * * AND FRACTAL DIMENSION. REAL CALA(50), CALB(50), CAL3(50), CAL4(50), BV(50), CU(50), CV(50) REAL MI(50,20),MIC(50,20),PRES(50),POR(50),PERM(50),D50(50) REAL PORS(50), PERMS(50), D50S(50), VRR(50), MVRR(50), XVAR(50) REAL CMU, CMV CHARACTER LEVEL(50)*1,LEVELS(50)*1,PTYPE(50)*1,ALPHA(50)*1 INTEGER I, J, LOC (50), LOCS (50), COUNT, IX, VARBLE ********************************** * DATA FOR THE CALIBRATION CURVES * AND CORRESPONDING PRESSURES ARE -* * READ INTO 1-D ARRAYS USING DATA * STATEMENTS. *************************

```
DATA CALA
/1.000,0.965,0.964,0.963,0.962,0.960,0.959,0.954,
     + 0.949,0.945,0.942,0.939,0.925,0.910,0.895,0.881,
     + 0.866,0.852,0.836,31*0.821/
      DATA CALB
/1.000,0.910,0.904,0.900,0.896,0.893,0.886,0.882,
     + 0.876,0.871,0.865,0.861,0.839,0.811,0.788,0.760,
     + 0.732,0.704,0.678,31*0.654/
      DATA PRES
/0.0,20.0,30.0,40.0,50.0,60.0,80.0,100.0,125.0,
     + 150.0,175.0,200.0,300.0,400.0,500.0,600.0,700.0,
     + 800.0,900.0,31*1000.0/
      DATA
ALPHA/'A','B','C','D','E','F','G','H','I',41*'J'/
      DO 10 I=1,50
      READ(5, *)
LEVEL(I), PTYPE(I), BV(I), LOC(I), (MI(I,J), J=1, 20)
   10 CONTINUE
      CAL3(1) = 0.000
      CAL4(1) = 0.000
      DO 20 J=1,19
       CAL3(J+1) = CALA(J) - CALA(J+1)
       CAL4 (J+1) = CALB (J) - CALB (J+1)
   20 CONTINUE
      DO 30 I=1,50
       MIC(I,1) = 0.000
       DO 40 J=1,19
    MIC(I,J+1) = MI(I,J) - MI(I,J+1)
   40 CONTINUE
   30 CONTINUE
      DO 50 I=1,50
        IF(PTYPE(I).EQ.'L') THEN
      DO 60 J=1,20
       MI(I,J) = MIC(I,J) - CAL3(J)
   60 CONTINUE
        ELSE
      DO 70 J=1,20
       MI(I,J) = MIC(I,J) - CAL4(J)
```

```
50 CONTINUE
     DO 80 I=1,50
      DO 90 J=1,20
      MIC(I,J) = MI(I,J)
      IF(MIC(I,J).LE.0.000) THEN
       MIC(I, J) = 0.000
      ENDIF
  90 CONTINUE
  80 CONTINUE
***********************************
* CALL SUBROUTINE AREA FOR THE
* CALCULATION OF POROSITY, PERMEABILITY *
* AND MEAN PORE THROAT SIZE FOR EACH
                                     ×
* SAMPLE.
                                      *
*******
     CALL AREA (BV, PRES, MIC, POR, PERM, D50)
************************
* SORT ARRAYS BY LOCATION *
*********
     COUNT=0
     DO 250 I=1,5
      DO 260 J=1,50
       IF(LOC(J).EQ.I) THEN
        COUNT=COUNT+1
        LOCS(COUNT) = LOC(J)
        LEVELS (COUNT) =LEVEL (J)
        PORS(COUNT) = POR(J)
        PERMS (COUNT) = PERM (J)
        D50S(COUNT) = D50(J)
       ENDIF
  260 CONTINUE
  250 CONTINUE
*****
* SORT ARRAYS BY LEVEL *
********
     DO 270 I=1,10
      COUNT=I
      DO 280 J=1,50
       IF(LEVELS(J).EQ.ALPHA(I)) THEN
        LEVEL (COUNT) =LEVELS (J)
```

70 CONTINUE ENDIF 193

```
LOC(COUNT)=LOCS(J)
        POR(COUNT) = PORS(J)
        PERM(COUNT) = PERMS(J)
        D50(COUNT) = D50S(J)
        COUNT=COUNT+10
       ENDIF
 280 CONTINUE
  270 CONTINUE
**********
* PRINT OUT RESULTS FROM
                            *
                            *
* SUBROUTINE AREA.
*********
     WRITE(6,200)
  200 FORMAT (T20, 'DATA SUMMARY-CORE
J'//,T10,'LEVEL',T17,'LOCATION',
          T27, 'POROSITY', T37, 'PERMEABILITY', T51, 'MEAN PORE
     +
THROAT'/,
          T42, 'md', T57, 'nM'/, 60('-'))
    +
     DO 210 I=1,50
      WRITE(6,220) LEVEL(I),LOC(I),POR(I),PERM(I),D50(I)
 220 FORMAT (T12, A2, T20, I2, T30, F6.3, T42, F11.3, T55, F8.3)
  210 CONTINUE
******************
* CALL SUBROUTINE VARIO THREE TIMES USING
                                          *
* THE DATA CALCULATED IN SUBROUTINE AREA.
VARBLE=1
      CALL VARIO (PERM, VARBLE)
      VARBLE=2
      CALL VARIO (POR, VARBLE)
      VARBLE=3
      CALL VARIO(D50, VARBLE)
      STOP
      END
      SUBROUTINE AREA (BVOL, PRESS, MICOR, P, PM, DIA)
```

```
REAL
BVOL(50), PRESS(50), PC(50), MICOR(50,20), HG(50,20), Y(50)
      REAL
P(50), PM(50), DIA(50), TOTAL, PC50(50), NUMB, RATIO, RES
      REAL
BPAR(4), INT, S1, S2, S3, S4, IFT, ANGLE, LFAC, C(19, 3), X(50)
      INTEGER I, J, IC, NX, IER
      TOTAL=0.000
***************************
* CONVERT PRESSURE VECTOR TO
* UNTOS OF ATM E-2 AND CALCULATE *
* HG ___ URATION VS PRESSURE
****************************
      DO 100 J=1,20
       PC(J) = (1.0/((PRESS(J)/14.7)+1.0422)) **2
  100 CONTINUE
      DO 110 I=1,50
       DO 120 J=1,20
        TOTAL=MICOR(I, J) +TOTAL
        HG(I,J) = TOTAL
  120
       CONTINUE
       P(I)=TOTAL/BVOL(I)
       TOTAL=0.000
  110 CONTINUE
      DO 130 I=1,50
       DO 140 J=1,20
        MICOR(I,J)=HG(I,J)/(P(I)*BVOL(I))
  140
       CONTINUE
  130 CONTINUE
*******************************
* FIND CAPILLARY PRESSURE
                                   *
* CURRESPONDING TO 50% HG
                                   *
                                   *
* SATURATION WITHIN THE SAMPLE.
* USE THIS VALUE TO CALCULATE
                                   *
* THE MEAN PORE THROAT DIAMETER. *
************************
      DO 150 I=1,50
       DO 160 J=1,19
         IF(0.50.GT.MICOR(I,J).AND.0.50.LE.MICOR(I,J+1))
THEN
          NUMB=0.50-MICOR(I,J)
          RATIO=NUMB/(MICOR(I,J+1)-MICOR(I,3))
```

PC50(I)=(PRESS(J)+RATIO*(PRESS(J+1)-PRESS(J)))+15.3204 GO TO 150 ENDIF 160 CONTINUE 150 CONTINUE ************** * CALCULATION OF MEAN PORE THROAT * * SIZE USING PRESSURE VALUES FROM * * * ARRAY PC50. *************** ************************************* * THE MEAN PORE DIAMETER IN nM IS CALCULATED. * * * THE CONTACT ANGLE IS CONVERTED TO RADIANS * * AND THE MEAN PRESSURE TO MPA. ************** DO 200 I=1,50 DIA(I)=-4.0*480.0*COS(140.0*0.01745329)/(PC50(I)*0.0067708) 200 CONTINUE IC=19 NX=20 ********* * INITIALIZE END CONDITIONS * * * PARAMETERS FOR NATURAL * SPLINE FUNCTION. ÷ ******** BPAR(1) = 0.0BPAR(2) = 0.0BPAR(3) = 0.0BPAR(4) = 0.0*********************** * CALL IMSL SUBROUTINE ÷ * ICSICU WHICH CALCULATES * THE SPLINE COEFFICIENTS * ********* DO 170 I=1,50 X(1) = MICOR(I, 1)Y(1) = PC(1)DO 180 J=1,19 IF(MICOR(I,J).GE.MICOR(I,J+1)) THEN RES=(MICOR(I,J)-MICOR(I,J+1))+0.001

```
MICOR(I, J+1) = MICOR(I, J+1) + RES
        X(J+1) = MICOR(I, J+1)
       ELSE
        X(J+1) = MICOR(I, J+1)
        ENDIF
         Y(J+1) = PC(J+1)
 180 CONTINUE
      CALL ICSICU(X,Y,NX,BPAR,C,IC,IER)
************************
* USING THE CALCULATED SPLINE *
* COEFFICIENTS THE AREA UNDER *
* THE CAPILLARY PRESSURE
* CURVE MAY BE FOUND BY
                             ×
* NUMERICAL INTEGRATION.
                              *
************************
      INT=0.0
      DO 190 J=1,19
       S1=(X(J+1)-X(J))
       S2=S1**2
       S3=S1**3
       S4=S1**4
INT=INT+C(J,3)/4.0*S4+C(J,2)/3.0*S3+C(J,1)/2.0*S2+Y(J)*S1
  190 CONTINUE
*******************************
                                    *
* CALCULATE PERMEABILITY FROM
                                    *
* PURCELL'S EQUATION. FOR OUR
* CASE:
*
* INTERFACIAL TENSION=480 DYNES/CM *
* CONTACT ANGLE=140 DEGREES
                                    ×
* LITHOLOGY FACTOR=0.216
                                    ÷
+
* WHEN K IS LOW, THE SPLINE
* FUNCTION MAY OCCASIONALLY RESULT *
* IN A NEGATIVE AREA CALCULATION.
                                    ÷
                                    +
* SINCE THIS IS PHYSICALLY
* IMPOSSIBLE, ANY OCCURENCES
                                    *
                                    ×
* RESULTING IN A NEGATIVE K VALUE
                                    ¥
* ARE ARBITRARILY SET TO 0.10 MD.
*******************
```

```
ANGLE=140.0
     IFT=480.0
     LFAC=0.216
     RAD=ANGLE*0.01745329
PM(I)=10.24*((IFT*COS(RAD))**2)*LFAC*P(I)*(INT/14.7**2)
     IF(PM(I).LE.0.000) THEN
      PM(I) = 0.10
     ENDIF
  170 CONTINUE
     RETURN
     END
_
______
     SUBROUTINE VARIO (VAR, VARBLE)
*****************
* THIS SUBROUTINE CALCULATES THE VARIOGRAM
                                            ÷
* FOR THE DATA IN QUESTION BASED UPON THE
                                            *
* LOGARITHM OF THE DATA VALUES CALCULATED
                                            *
* IN SUBROUTINE AREA. THE AUTOCORRELATION
                                           *
* FUNCTION IS THEN CALCULATED AND CONVERTED
                                           *
* TO A FRACTAL DIMENSION.
***************
      REAL VAR(50), VR(50), LAGV(50), ARRAY(50, 50)
      REAL LAG, AC, ACM, ACV, FDIM
      INTEGER I, J, K, NPR, LG, NX, VARBLE
      PARAMETER (LAG=4.8, NX=10.0)
      IF (VARBLE .EQ. 1) THEN
       WRITE(6,398)
       ELSEIF (VARBLE . EQ. 2) THEN
        WRITE(6,399)
        ELSE
        WRITE(6,400)
      ENDIF
  398 FORMAT('1', 'VARIOGRAM SUMMARY ',
          '(PERMEABILITY)',/)
     +
  399 FORMAT('1', 'VARIOGRAM SUMMARY ',
           '(POROSITY)',/)
     +
  400 FORMAT ('1', 'VARIOGRAM SUMMARY ',
     +
           '(PORE THROAT DIAMETER)',/)
      WRITE(6,401)
```

```
401 FORMAT ('X VALUE', 6X, 'Y VALUE', /)
```

```
402 CONTINUE
     LG=NX-1
* THE VECTOR VAR IS CONVERTED INTO A
                                         ×
* TWO DIMENSIONAL ARRAY TO FACILITATE
                                         *
* THE CALCULATION OF THE SEMI-VARIANCES. *
DO 403 J=1,5
       DO 404 I=1,NX
        \operatorname{ARRAY}(I,J) = \operatorname{VAR}((J-1) * 10 + I)
  404
       CONTINUE
  403 CONTINUE
      DO 405 K=1,LG
       VR(K) = 0_{\circ} 0
       DO 406 J=1,5
        DO 407 I=1,NX-K
         VR(K) = VR(K) + (ARRAY(I + K, J) - ARRAY(I, J)) **2
  407
        CONTINUE
      CONTINUE
  406
       NPR=5*(NX-K)
       VR(K) = 0.5 * VR(K) / FLOAT(NPR)
       LAGV(K) = FLOAT(K) * LAG
       WRITE(6,408) LAGV(K),VR(K)
  408 FORMAT(F8.2,5X,F12.3)
  405 CONTINUE
************************************
                                      *
* CALCULATE AUTOCORRELATION FUNCTION
* AND ASSOCIATED FRACTAL DIMENSION.
*****************************
      ACM=0.0
      ACV=0.0
      AC=0.0
      DO 409 I=1,50
       VAR(I) = EXP(VAR(I))
       ACM = ACM + VAR(I)
  409 CONTINUE
      ACM=ACM/50
      DO 410 I=1,50
       ACV=ACV+(VAR(I)-ACM)**2
  410 CONTINUE
```

DO 402 I=1,50

VAR(I)=LOG(VAR(I))

```
199
```

```
ACV=ACV/50
      DO 411 I=: .49
       AC=AC+((VAR(I)-ACM)*(VAR(I+1)-ACM))
 411 CON 🤟
               T 10)
      AC=.
                    <sup>•</sup>2.0*AC+2.0)/(2.0*LOG(2.0)))
      FDIM-
                     CM, ACV
      WRITE
                   VALUE = ', F9.3, /'MEAN VARIANCE =
  412 FORMA
               .
',F14.3)
      WRITE(6, .... AC, FDIM
  413 FORMAT ('AUTOCORRELATION FUNCTION =', F7.3,
          /'FRACTAL DIMENSION =', F7.3)
     +
      RETURN
      END
```

200