Solute transport in variable-aperture fractures:
An investigation of the relative importance
of Taylor dispersion and macrodispersion

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Abstract. Dispersion of solutes in a variable aperture fracture results from a combination of molecular diffusion and velocity variations in both the plane of the fracture (macrodispersion) and across the fracture aperture (Taylor dispersion). We use a combination of physical experiments and computational simulations to test a theoretical model in which the effective longitudinal dispersion coefficient \( D_L \) is expressed as a sum of the contributions of these three dispersive mechanisms. The combined influence of Taylor dispersion and macrodispersion results in a nonlinear dependence of \( D_L \) on the Peclet number \( (Pe = Vb/D_m) \), where \( V \) is the mean solute velocity, \( b \) is the mean aperture, and \( D_m \) is the molecular diffusion coefficient. Three distinct dispersion regimes become evident: For small \( Pe \) (\( Pe \ll 1 \)), molecular diffusion dominates resulting in \( D_L \propto Pe^0 \); for intermediate \( Pe \), macrodispersion dominates \( (D_L \propto Pe) \); and for large \( Pe \) (Taylor dispersion dominates \( (D_L \propto Pe^2) \). The \( Pe \) range corresponding to these different regimes is controlled by the statistics of the aperture field. In particular, the upper limit of \( Pe \) corresponding to the macrodispersion regime increases as the macrodispersivity increases. Physical experiments in an analog, rough-walled fracture confirm the nonlinear \( Pe \) dependence of \( D_L \) predicted by the theoretical model. However, the theoretical model underestimates the magnitude of \( D_L \). Computational simulations, using a particle-tracking algorithm that incorporates all three dispersive mechanisms, agree very closely with the theoretical model predictions. The close agreement between the theoretical model and computational simulations is largely because, in both cases, the Reynolds equation describes the flow field in the fracture. The discrepancy between theoretical model predictions and \( D_L \) estimated from the physical experiments appears to be largely due to deviations from the local cubic law assumed by the Reynolds equation.

1. Introduction

Solute transport in a rough-walled fracture is controlled by diffusive and advective processes. The Peclet number \( Pe = Vb/D_m \), where \( V \) is the mean solute velocity, \( b \) is a characteristic length scale (e.g., fracture aperture), and \( D_m \) is the molecular diffusion coefficient, defines the relative importance of each transport process. Molecular diffusion dominates for \( Pe \ll 1 \). Within the advection-dominated regime (larger \( Pe \) values), two different mechanisms lead to dispersion because of variable velocity within the rough-walled geometry: Taylor dispersion and macrodispersion. Taylor dispersion results from mixing induced by velocity variations across the fracture aperture. Macrodispersion is caused by velocity variations in the plane of the fracture that result from aperture variability. Because of their different origins, Taylor dispersion and macrodispersion exhibit different fundamental dependence on \( Pe \), with Taylor dispersion proportional to \( Pe^2 \) and macrodispersion proportional to \( Pe \).

Computational simulations of transport in fractures have incorporated the influence of either Taylor dispersion (i.e., parallel plate fractures [e.g., Hull et al., 1987; Ippolito et al., 1994]) or macrodispersion (i.e., constant velocity across the fracture aperture [e.g., Moreno et al., 1988; Thompson, 1991; Thompson and Brown, 1991]) but not both. Recent experimental evidence [Ippolito et al., 1994] and scaling analyses [Roux et al., 1998] suggest that dispersion in variable-aperture fractures can be described as a sum of molecular diffusion, Taylor dispersion, and macrodispersion. Roux et al. [1998] also presented scaling arguments suggesting that the \( Pe \) range within which each dispersion process dominates is controlled by the mean, variance, and correlation scale of the aperture field.

To date, there are no experimental or computational studies that fully delineate the various regimes of solute dispersion in variable-aperture fractures. Ippolito et al. [1994] experimentally demonstrated the influence of two distinct dispersion regimes (i.e., Taylor dispersion and macrodispersion). However, they did not quantify the statistics of the aperture field in their experimental fracture, making it difficult to generalize their results. Keller et al. [1995, 1999] measured the longitudinal dispersion coefficient \( D_L \) over a range of \( Pe \) in two different natural fractures in granite. They compared these results to \( D_L \) predicted using stochastic theory and the measured statistics of...
their aperture fields. However, large-scale aperture variability (clearly evident in images of the aperture fields in both of their fractures) likely dominated the dispersion process, partially invalidating comparisons of experimental results to stochastic theory. Dronfield and Silliman [1993] demonstrated a nonlinear relationship between $D_L$ and $Pe$ ($D_L \propto Pe^{-1.4}$), based on transport experiments in a sand-roughened analog fracture. This result suggests that their experiments (run over a narrow range of $Pe$) were in the transition zone between Taylor dispersion and macrodispersion.

In this paper we use a combination of physical experiments and computational simulations to explore the $Pe$ ranges of the different dispersion regimes, as controlled by the mean, variance, and correlation scale of the aperture field. We also present a theoretical expression for $D_L$ that combines Taylor dispersion and macrodispersion. Our experiments use a light transmission technique that yields high-resolution, accurate measurements of both aperture fields and solute concentration fields in transparent analog fractures. This approach offers the advantages that the aperture field is measured at the time of the experiment and dispersion of a dye pulse within the fracture is followed directly, avoiding assumptions about mixing in the inflow and outflow manifolds that are required when estimating $D_L$ from breakthrough curves measured at the outflow. Our computational model tracks particles through a variable aperture fracture. The velocity field within the fracture is specified using a parabolic velocity profile across the aperture, where the local, aperture-averaged velocity is obtained from a numerical solution of the Reynolds equation. In addition to advection within this three-dimensional velocity field, particles undergo three-dimensional molecular diffusion. Thus the mechanisms that cause both Taylor dispersion and macrodispersion are incorporated into the model.

We first performed experiments in a Hele-Shaw cell (flat, parallel-plate fracture) and used the results to verify our computational model in the absence of aperture variation and to measure $D_m$ for our solute. We then experimentally investigated the range of $Pe$ where the transition between macrodispersion and Taylor dispersion occurs in a rough-walled fracture. To transcend experimental limitations and explore Taylor dispersion and macrodispersion regimes over a wide range of $Pe$ for aperture fields with different statistics, we designed a sequence of computational simulations. First, we compared our modeling approach to the experiments to evaluate model error and then simulated a much wider $Pe$ range within a synthetic aperture field with similar statistics to our experimental fracture. Finally, we simulated transport through two additional synthetic aperture fields to consider the influence of aperture variance and correlation length on the $Pe$ range over which the different dispersion mechanisms dominate.

Our simulations support the Roux et al. [1998] scaling estimates of the $Pe$ range corresponding to the relative dominance of Taylor dispersion and macrodispersion. The theoretical expression for $D_L$, which is a sum of the macrodispersion coefficient [Gelhar, 1987, 1993] and Taylor dispersion coefficient and shares the fundamental assumptions of the Reynolds equation, agrees closely with our computational results. However, for our rough-walled experimental fracture, theoretical estimates of $D_L$ are significantly less than the experimentally measured values (e.g., by 51% at $Pe = 300$). We believe that this discrepancy between experiment and theory is primarily due to the inability of the Reynolds equation, upon which the stochastic theory is based, to fully describe the velocity field within a rough-walled fracture [e.g., Yeo et al., 1998; Nicholl et al., 1999].

2. Theoretical Description of Dispersion in Variable Aperture Fractures

In a parallel-plate fracture the primary mechanism causing dispersion is the well-known phenomenon of Taylor dispersion [Taylor, 1953; Aris, 1956]. The Taylor dispersion coefficient for a parallel-plane fracture is [e.g., Fischer et al., 1979]

$$D_{L,Taylor} = \frac{V^2 b^2}{210 D_m},$$

where $V$ is the average velocity in the fracture, $b$ is the fracture aperture, and $D_m$ is the molecular diffusion coefficient.

For transport in a variable-aperture fracture, Gelhar [1987, 1993] developed a stochastic analysis of flow and solute transport. Gelhar’s analysis assumes that the logarithm of the aperture ($\beta = \ln b$) is a statistically stationary, Gaussian random field and that the flow within a variable-aperture can be described by the Reynolds equation. The Reynolds equation is based on the assumptions that aperture variations are relatively smooth and that the velocity profile across the aperture is parabolic, corresponding to local, plane Poiseuille flow [e.g., Zimmerman and Bodvarsson, 1996]. The stochastic analysis of flow reveals that the effective hydraulic aperture is equal to the geometric mean aperture. The stochastic transport analysis neglects the influence of local dispersion and results in the following expression for the macrodispersion coefficient:

$$D_{L,macro} = \sigma_b^2 \lambda \left[ 3 + I(\sigma_b^2/\sigma_{\beta}^2) \right] V = \sigma_b^2 \lambda B V,$$

In (2), $\sigma_{\beta}^2$ and $\lambda$ are the variance and integral scale of $\beta$, respectively,

$$B = 3 + I(\sigma_b^2/\sigma_{\beta}^2)$$

$$I(\sigma_b^2) = \int_0^\infty \left[ \exp (R_{\beta b}(u)) - 1 \right] du,$$

where $R_{\beta b}(u)$ is the covariance function of $\beta$ and $u$ is the nondimensional spatial separation variable, which equals the spatial separation divided by $\lambda$. Note that for an exponential covariance, $\lambda$ is equivalent to the correlation length or the length scale over which correlation in $\beta$ persists. Also note that $V$ in (2) is the mean solute velocity, equal to the mean flux through the fracture divided by the mean aperture $(<b>)$ [Gelhar, 1993]. Expression (2) incorporates the influence of variations in the mean flow velocity within the fracture plane but not the influence of Taylor dispersion. Equations (1) and (2) demonstrate that $D_{L,macro}$ is proportional to $V$ and that $D_{L,Taylor}$ is proportional to $V^2$, which suggests that at high flow rates, Taylor dispersion may dominate over macrodispersion, even in a variable-aperture fracture. Expressions (1) and (2) may be rewritten in a nondimensional form, in terms of $Pe = V/b/D_m$, where $(b)$ is the mean aperture and $V$ is the mean solute velocity:}

$$D_{L,Taylor}/D_m = Pe^2/210 = \alpha_{Taylor} Pe^2$$

$$D_{L,macro}/D_m = \sigma_b^2 \lambda B (b) Pe = \alpha_{macro} Pe.$$
In (5) and (6), $\sigma_{\text{Taylor}}$ and $\sigma_{\text{Macro}}$ are nondimensional coefficients for the contributions of Taylor dispersion and macrodispersion, respectively. Note that (5) represents an "effective" Taylor dispersion coefficient in a rough-walled fracture based on the mean aperture and mean solute velocity. A stochastic analysis of Taylor dispersion in a variable-aperture fracture is required to establish the validity of such a representation of the effective Taylor dispersion coefficient.

Roux et al. [1998] used scaling arguments to suggest three primary dispersion regimes in variable-aperture fractures: molecular diffusion, "geometric" dispersion, and Taylor dispersion. The "geometric" dispersion regime corresponds to the range of $Pe$ where velocity variations in the plane of the fracture dominate the mixing process and $D_L \approx Pe$. We note that this is equivalent to macrodispersion as described by (2) and (6), and we use the latter term to refer to this regime in the remainder of the paper. Roux et al. [1998] used scaling relationships for $D_L$ in the Taylor dispersion and macrodispersion regimes to define the approximate $Pe$ range within which macrodispersion is the dominant dispersion mechanism. However, they did not use the precise relationships (5) and (6) to quantify the macrodispersion and Taylor dispersion coefficients. Roux et al. [1998] also suggested that $D_L$ can be expressed as a sum of the three different components. This results in a first-order approximation of the total nondimensional longitudinal dispersion coefficient of the form:

$$\frac{D_L}{D_m} = \tau + \alpha_{\text{Macro}} Pe + \alpha_{\text{Taylor}} Pe^2,$$

where $\tau$ is the tortuosity for diffusion within the fracture, typically $<1.0$, reflecting the reduced rate of molecular diffusion in a geometrically complex void space. For typical $Pe$ ranges, $\tau$ is an insignificant contribution to $D_L/D_m$ and may be dropped from (7). Equation (7) suggests that $\alpha_{\text{Macro}}$ will be influenced by the statistics of the aperture field and will increase with $\sigma_{\text{b}}^2$ and $\lambda/(b)$. Additionally, the $Pe$ range over which macrodispersion can be expected to dominate is

$$1/\alpha_{\text{Macro}} < Pe < \alpha_{\text{Macro}}/\alpha_{\text{Taylor}},$$

where $\tau$ is assumed to be approximately 1.

Figure 1 shows $D_L/D_m$, from (7), plotted against $Pe$ for two hypothetical values of $\alpha_{\text{Macro}}$ (0.2 and 20) with $\tau$ and $\alpha_{\text{Taylor}}$ equal to 1 and 1/210, respectively. The curve for $\alpha_{\text{Macro}} = 0.2$ transitions directly from a molecular diffusion regime (slope equal to 0) to the Taylor dispersion regime (slope equal to 2), whereas the curve for $\alpha_{\text{Macro}} = 20$ exhibits a large region, $O(10^{-1}) < Pe < O(10^4)$, where macrodispersion dominates (slope equal to 1). To highlight these different dispersion regimes, it is useful to plot $D_L$ in the nondimensional form $D_L/(V(b))$ against $Pe$ (Figure 2). Figures 1 and 2 demonstrate the importance of quantifying $\alpha_{\text{Macro}}$ to determine the $Pe$ range associated with the different dispersion regimes in a given fracture.

To fully study the $Pe$ range corresponding to different dispersion regimes requires a fracture that is long compared to $\lambda$. Theoretical results for a two-dimensional, isotropic, random field with exponential covariance suggest that $D_L$ should reach 99% of its asymptotic value after the solute has traveled a distance of $\sim 20\lambda$ [e.g., Dagan, 1984]. Similarly, for transport between parallel plates, Taylor dispersion should become fully developed at a distance of $\sim 0.4(b)/Pe$ [e.g., Fischer et al., 1979]. Thus, to make a meaningful comparison of physical and computational experiments to theory, we require a stationary field with small $\lambda$ compared to the dimensions of the field. This ensures that $D_L$ will become relatively constant within the scale of the experiment. We also require a source whose transverse dimensions extend over $\sim 20\lambda$ to eliminate nonergodic effects. However, to avoid an increase in Taylor dispersion due to the velocity variations at the lateral boundaries of the fracture [e.g., Doshi et al., 1978], the source should also be narrow.
Table 1. Fracture Dimensions and Measured Aperture Statistics

<table>
<thead>
<tr>
<th>Fracture Type</th>
<th>Hele-Shaw</th>
<th>Rough-Walled</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dimensions (cm × cm)</td>
<td>15.3 × 30.5</td>
<td>14.8 × 30.2</td>
</tr>
<tr>
<td>Dimensions (pixels × pixels)</td>
<td>972 × 1940</td>
<td>958 × 1958</td>
</tr>
<tr>
<td>Dimensions (λ × λ)</td>
<td>...</td>
<td>336 × 686</td>
</tr>
<tr>
<td>Pixel size, cm</td>
<td>1.57 × 10⁻²</td>
<td>1.54 × 10⁻²</td>
</tr>
<tr>
<td>Minimum aperture, cm</td>
<td>1.72 × 10⁻²</td>
<td>1.30 × 10⁻³</td>
</tr>
<tr>
<td>Maximum aperture, cm</td>
<td>2.09 × 10⁻²</td>
<td>3.85 × 10⁻²</td>
</tr>
<tr>
<td>(b), cm</td>
<td>1.93 × 10⁻²</td>
<td>2.21 × 10⁻²</td>
</tr>
<tr>
<td>σ₂(b)/b</td>
<td>2.73 × 10⁻²</td>
<td>2.72 × 10⁻¹</td>
</tr>
<tr>
<td>λ long axis, cm</td>
<td>...</td>
<td>4.4 × 10⁻²</td>
</tr>
<tr>
<td>λ short axis, cm</td>
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<tr>
<td>RMS Error, % of mean</td>
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<td>1.2</td>
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*In this study, λ represents the integral scale calculated by numerically integrating \( \rho(r) dr \), where \( \rho(r) = 1 - \gamma(r)/\sigma_σ^2 \) is the correlation function and \( \gamma(r) \) is the semivariogram. Previous studies using similar fractures [e.g., Glass et al., 1998; Nicholl et al., 1999] measured λ as the separation length at which the semivariogram reached the level of the sill (≈0.08 cm).

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3. Experimental Investigations

We carried out experiments in two analog fractures: a Hele-Shaw cell fabricated from two pieces of flat glass and a rough-walled fracture fabricated by mating two pieces of textured glass (fabrication details for both fractures are described by Nicholl et al., 1999). Table 1 presents the dimensions of each of the fractures. Experiments in the Hele-Shaw cell provided a test for our computational model as well as a measurement of \( D_m \). The rough-walled fracture was designed to have a stationary, isotropic aperture field with a correlation scale much smaller than the dimensions of the fracture and values of \( \sigma_β^2 \) and (b) in the range of values reported for natural fractures. These transparent analog fractures offered the additional advantage of allowing direct, full-field aperture and solute concentration measurements, with no disturbance of the fracture between the aperture measurements and transport experiments. Thus our experiments can be directly compared to computational simulations and theoretical results.

A test cell frame supported the fractures and allowed light from a feedback-controlled source to be transmitted through the entire fracture (Figure 3a). An electronically cooled, 12-bit, shuttered, charge-coupled-device (CCD) camera (2045 × 2033 pixels and 4096 gray levels) supported above the test cell measured the intensities of transmitted light. We measured aperture fields and solute concentration fields using a light transmission technique first proposed by Glass et al., 1991. The details of the experimental system, the aperture measurement technique, and a method for quantifying aperture errors are presented by Detwiler et al., 1999.

3.1. Measurement Techniques

A light-absorbing dye (Warner Jenkins FD&C Blue #1 dye) was used both as a tracer during transport experiments and as a light-absorbing solute for aperture measurement. The Beer-Lambert law describes the absorbance of monochromatic light by a dyed solution as a function of the distance through the solution (b) and the dye concentration (C). According to the Beer-Lambert law the absorbance at each pixel (identified using a double subscript “ij”, where i and j refer to the row and column index of the pixel) within a two-dimensional field is given by

\[
A_{ij} = \ln \left( \frac{I_{cl,ij}}{I_{dye,ij}} \right) = \mu C_{ij}b_{ij},
\]

where \( A_{ij} \) is the absorbance, \( I_{cl,ij} \) and \( I_{dye,ij} \) are the intensities transmitted through a clear and a dyed solution, respectively, \( \mu \)

Figure 3. (a) Schematic plan view of fracture cell and plumbing layout. (b) Cross section X-X’ of rough-walled fracture. (c) Cross section X-X’ of Hele-Shaw cell.
is the absorptivity of the solute, $C_{ij}$ is the dye concentration, and $b_{ij}$ is the thickness of the absorbing layer or the local aperture. Images of a fracture filled entirely with clear and dyed solutions yield arrays of intensity measurements, $I_{cij}$ and $I_{b wij}$ for use in (9). Normalizing (9) by its mean then gives an expression for the aperture:

$$b_{ij} = A_{ij}(b)/\langle A \rangle,$$

where $\langle b \rangle$ is the independently measured mean aperture and $\langle A \rangle$ is the mean (over all $ij$) of $A_{ij}$. We determined $\langle b \rangle$ by injecting a known volume of fluid into the fracture and measuring the area occupied by the fluid. Once $b_{ij}$ is calculated for the entire fracture, we can measure concentrations ($C_{ij}$) using images taken during transport experiments by directly applying (9).

If the light source is polychromatic, as is the case with our measurement system, the linear relationship between absorbance and concentration (9) is only approximate. Despite efforts to remove the influence of nonlinear absorbance on our measurements by reducing the measured wavelengths with a band-pass filter (Andover Corporation, 630 nm ± 5 nm) on the camera lens, some nonlinearity remained. Slight nonlinearity in dye absorbance results in aperture measurement errors that increase with dye concentration, but reducing dye concentration results in an increase in random errors due to signal noise caused by the smaller signal range (i.e., difference between $I_{cij}$ and $I_{b wij}$). We reduced the influence of random errors by averaging 80 images of each field and used the procedure described by Detwiler et al. [1999] to determine the dye concentration for measuring $I_{b wij}$ (0.05 g/L) that resulted in the minimum total error. We estimated root-mean-square (RMS) aperture measurement errors of 0.8 and 1.2% of the mean aperture for the Hele-Shaw cell and the rough-walled fracture, respectively.

For transport experiments we could not average multiple images, so we used a higher dye concentration that utilized the full dynamic range of the CCD camera. The use of a higher dye concentration reduces the influence of noise on measurements of $C_{ij}$ thus increasing the sensitivity of our measurements, especially in regions of low concentration (i.e., solute plume tails). However, a higher dye concentration also results in nonlinear absorbance in regions of high concentration. We accounted for the influence of nonlinear absorbance on concentration measurements by fitting the following function to a series of measurements made at different dye concentrations (0.025, 0.05, 0.10, 0.15, 0.20, and 0.25 g/L):

$$A_{ij} = e_{ij}C_{ij}/(f_{ij} + C_{ij}).$$

In (11), $e_{ij}$ and $f_{ij}$ are fitting parameters determined at each pixel in the array. Note that for low concentrations (i.e., $C_{ij} \ll f_{ij}$), (11) reduces to a linear relationship similar to (9). To obtain the concentration at each location, we used the fitted parameters $e_{ij}$ and $f_{ij}$ and solve (7) for $C_{ij}$:

$$C_{ij} = (f_{ij}/e_{ij})/(1/A_{ij} - 1/e_{ij}).$$

The maximum RMS concentration error at any location is $\sim 0.023 C_{ij} C_{ij} = 0.25$ g/L, based on an analysis discussed in detail in the appendix.

3.2. Measured Aperture Fields

Table 1 provides a summary of the dimensions and statistics of the two aperture fields. The aperture field for the Hele-Shaw cell exhibits a narrow distribution (variance, $\sigma_i^2 = 2.78 \times 10^{-7}$ cm$^2$) about its mean ($\langle b \rangle = 0.0193$ cm). The $i$ and $j$ semivariograms (long and short principal axes, respectively) for the Hele-Shaw cell (Figure 4) show a relatively strong trend along the short axis and a milder trend along the long axis. Although the glass plates were flat, clamping pressure resulted in smaller apertures along the edges of the cell ($\sim 0.018$ cm), larger apertures along the centerline of the cell (0.020 cm), and uneven compression of the plastic shims along the length of the fracture (Figure 3b). These features led to apparent large-scale trends in the aperture field as evident in the variograms shown in Figure 4.

The aperture field of the rough-walled fracture has a wider, negatively skewed distribution (see Figure 7), with $\sigma_i^2 = 3.62 \times 10^{-5}$ cm$^2$ and $\langle b \rangle = 0.0221$ cm. Nicholl et al. [1999] present an image of a portion of the rough-walled aperture field. The semivariograms for the rough-walled fracture (Figure 4) indicate that the field is isotropic and reaches the level of the sill at a separation of $\sim 0.08$ cm. The semivariograms also indicate a slight negative correlation at separations between $\sim 0.08$ cm and $\sim 0.18$ cm caused by the repetitive nature of the individual fracture surfaces. Numerically integrating the correlation functions of $b_{ij}$ yields estimates of $\lambda$ of $\sim 0.044$ cm along both the long and short axes.

3.3. Transport Experiments

We conducted a series of transport experiments in each fracture over a range of flow rates. To facilitate reproducibility, a computer-controlled flow through the fracture by activating/deactivating solenoid valves measured flow rates by recording outflow mass at equal intervals and triggered the CCD camera to acquire images at specified times. Inflow and outflow man-
folds provided uniform pressure across the width of the two ends of each fracture, and no-flow boundaries were applied to the sides of each fracture (Figure 3). A constant head reservoir at the inflow and a stabilized drip point at the outflow created steady, reproducible flow rates through the fracture. Table 2 summarizes the flow rates for each experiment. The CCD camera required ~11 s to write each image to disk, so we chose the maximum flow rate for each fracture so that we could acquire at least 10 images (i.e., concentration fields) during an individual transport experiment. The minimum flow rate represents the reproducible limit achievable using the constant head reservoirs of our system.

The initial condition for each experiment is shown in the first frame of Plates 1a and 1b. This initial condition provides two advantages: a known initial concentration distribution and negligible edge effects caused by the no-flow boundaries of the fracture. We pumped a 0.375 g/L slug of solute through the injection port in the center of the inflow manifold (Figure 3a) into the center of the inflow end of the fracture. We then flushed the inflow manifold by pumping clear water into the two ends (i.e., inflow and waste line in Figure 3a) and out through the injection port. After flushing the inflow manifold with clear water, we stopped the pump, closed the valve at the waste side of the inflow manifold, and opened the outflow manifold, initiating flow through the fracture under constant gradient. In designing the transport experiments we considered the possible influence of gravitational effects on dispersion in the fractures. Relatively small density gradients can lead to enhanced dispersion at early times (i.e., before density gradients are reduced by dispersion) [e.g., Reejhsinghani et al., 1966]. Our results (discussed in section 3.5) confirm that density effects were negligible.

Plates 1a and 1b show sequences of three solute concentration fields from experiments in the Hele-Shaw cell and the rough-walled fracture. The concentration fields demonstrate the initial condition in the first frame and the effect of dispersion on the plumes in the two successive frames. The role of aperture variability in enhancing dispersion is clearly evident in Plate 1b. The progressive reduction in the peak concentration is also evident in Plates 1a and 1b.

### 3.4. Analysis of Concentration Fields

The product of a measured solute concentration field and the measured aperture field yields the solute mass at each pixel in the field, and summation over the entire field yields the total mass within the system. The total mass measured in each of the fields obtained during any of the experiments varied by no more than ±3% from the total mass averaged over all fields from a single experiment. Good mass conservation further confirms the accuracy of our measurement techniques. The rates of change of the first and second spatial moments ($M_{1x}$ and $M_{2x}$, respectively) of solute mass in the flow direction with time are equivalent to the mean solute velocity ($V$) and are twice the longitudinal dispersion coefficient ($2D_L$), respectively [e.g., Aris, 1956]. We calculated $M_{1x}$ for each solute mass field using

$$M_{1x} = \sum_{i=1}^{nx} \sum_{j=1}^{ny} C_{ij}b_{ij}x_{ij} \left/ \sum_{i=1}^{nx} \sum_{j=1}^{ny} C_{ij}b_{ij} \right.$$ (13)

where $C_{ij}$, $b_{ij}$, and $x_{ij}$ are the concentration, aperture, and $x$ coordinate at pixel $ij$, respectively, and $nx$ and $ny$ are the number of measurements in the $x$ and $y$ directions. We then calculated $M_{2x}$ for each field using

$$M_{2x} = \sum_{i=1}^{nx} \sum_{j=1}^{ny} C_{ij}b_{ij}(x_{ij} - M_{1x})^2 \left/ \sum_{i=1}^{nx} \sum_{j=1}^{ny} C_{ij}b_{ij} \right.$$ (14)

We then estimated $D_L$ and $V$ by plotting $M_{1x}$ and $M_{2x}$ against time. Note that this method of measuring $D_L$ requires no assumptions about the initial condition for the experiment. The initial concentration distribution is directly measured. This approach overcomes the difficulties involved in using breakthrough curves and an analytical solution to the one-dimensional advection-diffusion equation for estimating trans-
Plate 1. A portion of three concentration fields measured during (a) an experiment in the Hele-Shaw cell, (b) an experiment in the rough-walled fracture, and (c) a simulation in the rough-walled fracture. Each field represents a 3 cm × 9 cm region of the fracture centered on the first spatial moment of solute mass in the x direction and the center of the fracture in the y direction. In the Hele-Shaw cell, $M_{1x} = 1.1, 6.6, \text{ and } 12.0 \text{ cm}$ in the three frames and in the rough-walled fracture, $M_{1x} = 1.6, 5.4, \text{ and } 9.4 \text{ cm}$ in the three frames. In the Hele-Shaw cell, the effect of Taylor dispersion is evident as growth of the plume in the flow direction with little spreading perpendicular to the flow direction. In the rough-walled fracture, the enhanced dispersion caused by aperture variability is evident.
and macrodispersion (Pe = 800. We determined
required for development of Taylor dispersion (Pe)
On the basis of theory presented in section 2, the distances
in section 2 were met at very early times in our experiments.
the initial concentration field until solute began to exit the
faster is, no enhanced dispersion was observed at early time (i.e.,
respectively. Linear plots of M
parameters for the rough-walled fracture. The error bars represent
the possible range of D_L estimates resulting from concentra-
tion measurement errors (see the appendix for a discussion of
errors).
port parameters. The latter approach assumes initial
conditions that are typically difficult to verify, and mixing in the
inflow and outflow systems leads to inaccuracies in estimates of
transport parameters.
3.5. Experimental Results
Experiments in both the Hele-Shaw cell and the rough-
walled fracture resulted in linear plots of M_{22}, versus time from
the initial concentration field until solute began to exit the
fracture. This indicates that the conditions for full development
of both macrodispersion and Taylor dispersion described
in section 2 were met at very early times in our experiments.
On the basis of theory presented in section 2, the distances
required for development of Taylor dispersion (-0.4(b)Pe)
and macrodispersion (20a) are ~0.9~7.0 cm and ~0.9 cm,
respectively. Linear plots of M_{22}, versus time over the full range
of experimental flow rates also confirm that the influence of
natural convection due to density gradients was negligible; that
is, no enhanced dispersion was observed at early time (i.e.,
faster M_{22} growth), as would be expected with significant nat-
ural convection [e.g., Stockman, 1997].
We used the procedures described in section 3.4 to calculate
the mean solute velocity and dispersion coefficient for each
experiment (Table 2) (the appendix provides a detailed discus-
sion of errors in our estimates of D_L). For both fractures we
demonstrated the reproducibility of our experiments by re-
peating a number of experiments at one flow rate (see exper-
iments 5a, 5b, and 5c in Figure 5 and Table 2). Figure 5 is a plot
of D_L/D_m versus Pe for each set of experiments. The range of
Pe values covered by our experiments extends from ~100 to
800. We determined D_m to be 5.67 \times 10^{-6} cm^2/s by fitting (1),
the theoretical expression for Taylor dispersion between par-
allel plates, to the experimental results from the Hele-Shaw
cell. Although, as discussed in section 3.3, the aperture field in
the Hele-Shaw cell is not perfectly uniform (because of bowing of
the glass plates), the plates are essentially parallel along the
central portion of the cell traversed by the solute plume. Note
that this method is analogous to an accepted method for mea-
suring D_m that involves measuring the dispersion of a solute in
laminar flow through a tube and using the theoretical expres-
sion for Taylor dispersion in a tube to calculate D_m [e.g.,
Cussler, 1984].
In the rough-walled fracture the relationship between
D_L/D_m and Pe is also nonlinear (Figure 5), indicating that
Taylor dispersion contributes to dispersion over the Pe range
of our experiments. Dropping \tau and fitting (7) to our experi-
mental data results in estimates of \sigma_{\text{macro}}, \alpha, and D_L that
agree more closely with experimental results than several other
variations) to define the transmissivities between adjacent grid
blocks. For comparison with our experimental results we used
a grid that corresponded to the measured aperture field (e.g.,
958 \times 1958 with dimensions of each grid block equal to
0.0154 \times 0.0154 cm); this is approximately the same discreti-
ation used by Nicholl et al. [1999] to fabricate additional aperture fields for experimental investigation.

4. Computational Investigations
The theory presented in Section 2 suggests that the nature of
the Pe dependence of D_L will depend on aperture statistics
((b), \sigma_{\text{ap}}, and \lambda). To investigate the influence of Pe and aperture
variability on D_L, over a wider range of parameters (Pe,
\sigma_{\text{ap}}, and \lambda) than is possible in a single experimental frac-
ture, we simulate flow and transport through computer-
generated aperture fields. Measured aperture and concentra-
tion fields from our physical transport experiments allow us to
first test the computational model through direct comparison
before proceeding to additional computational studies.

4.1. Flow and Transport Solvers
To simulate the velocity field within the fracture, we used the
flow solver developed by Nicholl et al. [1999], which uses a
finite difference discretization of the Reynolds equation. We
used the harmonic average (found by Nicholl et al. [1999] to
agree more closely with experimental results than several other
variations) to define the transmissivities between adjacent grid
blocks. For comparison with our experimental results we used
a grid that corresponded to the measured aperture field (e.g.,
958 \times 1958 with dimensions of each grid block equal to
0.0154 \times 0.0154 cm); this is approximately the same discreti-
ization used by Nicholl et al. [1999] for this fracture.
Nicholl et al. [1999] compared flow computations made with
this flow solver to saturated flow experiments in the two frac-
tures used in this study, as well as a third fracture fabricated by
mating a single piece of rough glass with a single piece of
smooth glass. Their comparison indicated that the Reynolds
equation (and other two-dimensional variants that account for
convergence/divergence of flow and tortuosity of the center-
line) overestimates flow through the rough-walled fracture
used in the current study by ~26%. High-resolution simula-
tions on a subset of the entire field indicated that increasing the
resolution of aperture measurements, and subsequently the
finite difference grid, had a negligible influence (~2%) on the
results. Thus they concluded that it may be necessary to solve
the three-dimensional Stokes equations in situations where
improved accuracy is required. Because the Reynolds equation

Figure 5. Nondimensional dispersion coefficient (D_L/D_m) plotted against the Peclet number (Pe) for experiments in both fractures. The curves through the data points are the result of fitting (7) to the data. Table 3 lists the fitted parameters for the rough-walled fracture. The error bars represent the possible range of D_L estimates resulting from concentration measurement errors (see the appendix for a discussion of errors).
overestimates flow under a specified gradient in rough-walled fractures, we specified the mean flux to generate the flow fields for transport simulations resulting in simulated mean solute velocities that closely matched our experimentally measured means.

We simulated solute transport using a three-dimensional random-walk particle-tracking algorithm. We used the same discretization of the domain used for solving the Reynolds equation and assumed that the aperture within each grid block was constant. This is consistent with experimental aperture measurements that represent an average of the aperture within each pixel. In addition, our measurement system only provides measurements of the local aperture and not deviations of the local aperture midpoint from the center plane of the fracture. For the transport simulations we assume that the fracture is symmetric about the center plane to approximate the actual fracture geometry. However, it should be noted that when the assumptions underlying the Reynolds equation are satisfied, the flow field depends only on the aperture field and not explicitly on the geometry of the fracture surfaces. Particle displacements in each time step consisted of a two-dimensional advective displacement (in the $x$, $y$ plane) and a three-dimensional random diffusive displacement ($x$, $y$, $z$, where $z$ is across the aperture) reflecting the role of molecular diffusion. We calculated the advective displacements using the local gradients specified by the solution to the Reynolds equation and imposing a parabolic velocity profile across the local aperture. In the absence of diffusion, particles maintained their relative $z$ position, creating a pseudoadvection in the $z$ direction when a particle moved between adjacent grid blocks with different apertures. Particles colliding with the fracture wall because of diffusion across the aperture were reflected back into the fracture. Adaptive time stepping ensured that the three-dimensional velocity field was well sampled by each particle. The minimum of the following three criteria defined the length of each time step: the time required for a mean diffusive displacement of 5% of the local aperture and the times required for a particle traveling at the maximum local velocity (i.e., along the aperture centerline) to traverse 50% of the local grid block in both the $x$ and $y$ directions.

### 4.2. Comparison of Results From Physical Experiments and Computational Simulations

Accurate, full-field measurements of both aperture and concentration allow us to directly compare the results of the computational simulations to the experimental data. We specified initial particle locations based on the initial conditions in our experiments. The initial conditions were generated by taking the initial solute mass field from an experiment and assigning experiments. The initial conditions were generated by taking initial particle locations based on the initial conditions in our computational simulations to the experimental data. We specified concentration allow us to directly compare the results of the computations.

For the transport simulations we assume that the fracture is symmetric about the center plane to approximate the actual fracture geometry. However, it should be noted that when the assumptions underlying the Reynolds equation are satisfied, the flow field depends only on the aperture field and not explicitly on the geometry of the fracture surfaces. Particle displacements in each time step consisted of a two-dimensional advective displacement (in the $x$, $y$ plane) and a three-dimensional random diffusive displacement ($x$, $y$, $z$, where $z$ is across the aperture) reflecting the role of molecular diffusion. We calculated the advective displacements using the local gradients specified by the solution to the Reynolds equation and imposing a parabolic velocity profile across the local aperture. In the absence of diffusion, particles maintained their relative $z$ position, creating a pseudoadvection in the $z$ direction when a particle moved between adjacent grid blocks with different apertures. Particles colliding with the fracture wall because of diffusion across the aperture were reflected back into the fracture. Adaptive time stepping ensured that the three-dimensional velocity field was well sampled by each particle. The minimum of the following three criteria defined the length of each time step: the time required for a mean diffusive displacement of 5% of the local aperture and the times required for a particle traveling at the maximum local velocity (i.e., along the aperture centerline) to traverse 50% of the local grid block in both the $x$ and $y$ directions.

**Figure 6.** Comparison of $D_L/D_m$ versus $Pe$ for experiments and simulations in the Hele-Shaw cell and the rough-walled fracture. The curves through the Hele-Shaw cell data represent (5), the theoretical expression for Taylor dispersion. The curves through the data represent (7) fitted to each data set (neglecting $\tau$). The fitted parameters are presented in Table 4.
simulations closely reproduce the functional dependence of $D_L$ on $Pe$ evident in the rough-walled fracture. We believe this is because over the $Pe$ range for which Stokes flow is valid, errors in the velocity field obtained by solving the Reynolds equation are proportional to the mean velocity (and hence $Pe$). Thus, in this range, estimates of $D_L$ from simulations of transport should reliably reflect the actual $Pe$ dependence of $D_L$ though they may underestimate the magnitude of $D_L$.

### 4.3. Computational Simulations Investigating the Influence of $\sigma_b^2$ and $\lambda$ Over a Wide Range of $Pe$

Though the experiments and simulations in the rough-walled fracture both demonstrate the influence of Taylor dispersion on $D_L$, neither reach the $Pe$ range where macrodispersion becomes negligible (i.e., $D_L \approx Pe^{-2}$) as predicted by the theory presented in section 2. Additionally, our experimental aperture field embodies a single $\sigma_b^2$ and $\lambda$. In this section we use numerical simulations to investigate the role of aperture variability (i.e., $\sigma_b^2$ and $\lambda$) over a wider $Pe$ range than was possible experimentally.

As discussed in section 2, simulations at large values of $Pe$ require fractures that are somewhat longer than $0.4 \langle b \rangle Pe$ (the distance required for Taylor dispersion to become fully developed). To investigate the influence of aperture variability (quantified by $\langle b \rangle$, $\sigma_b^2$ and $\lambda$) on dispersion over a wide range of $Pe$, we generated three correlated random aperture fields that were significantly longer ($7.9 \text{ cm} \times 169.9 \text{ cm}$ and $512 \times 11,000$ grid) than the experimental field: field 1 with statistics (i.e., $\langle b \rangle$, $\sigma_b^2$, and $\lambda$) similar to those of the experimental field and fields 2 and 3, with different values of $\lambda$ and $\sigma_b^2$, respectively (Table 3). These longer fields satisfy the conditions for the full development of dispersion and thus allow meaningful comparison of theoretical results to computational simulations over a wide $Pe$ range.

We generated lognormally distributed aperture fields with a “hole-type” covariance function that captured the negative correlation caused by the slight periodicity of the experimental field. The spectral density function corresponding to this covariance function is

$$S_{bb}(\omega_x, \omega_y) = \frac{16 \sigma_b^2 \omega_x^4 \omega_y^4}{\pi^3} \exp \left( -\frac{4 \lambda^2 \omega_x^2 \omega_y^2}{\pi} \right), \quad (15)$$

where $\omega_x$ and $\omega_y$ are the wave numbers in the $x$ and $y$ directions, respectively, $\omega = \sqrt{\omega_x^2 + \omega_y^2}$, and $\lambda$ is the integral scale of the corresponding correlation function.

We used a fast Fourier transform algorithm to obtain $\beta(x, y)$, a normally distributed, correlated random field. Using this field, we generated the three synthetic, lognormally distributed, aperture fields ($b(x, y)$) with statistics presented in Table 3. Figure 7 compares the aperture distributions of the three synthetic fields and the experimental field. Synthetic field 1 has the same $\sigma_b$ and $\langle b \rangle$ as the experimental field but exhibits a slightly higher peak and less spread than the experimental field, because of the positive skew and long tail of the lognormal distribution. Note that although $\sigma_b$ is equal for the two fields, $\sigma_b$ (and thus $\sigma_{macro}$) is slightly different because of the different shapes of the two distributions. Figure 4 shows the normalized semivariograms of field 1 and the experimentally measured rough-walled fracture, demonstrating that field 1 is stationary and isotropic and has approximately the same correlation scale as the experimental field. Field 2 is identical to field 1, except that the length of the side of each grid block, and thus $\lambda$, was increased by a factor of 50. We generated field 3 by scaling the same $B(x, y)$ field used to generate field 1 to increase $\sigma_b$ by a factor of 7.7 (or $10 \sigma_b^2$). All three fields include a negative correlation at separations of $\sim 2\lambda$ to $\sim 4.5\lambda$, as in the experimental field.

We simulated transport in each of the three random fields in the range $1 \leq Pe \leq 5 \times 10^4$. Note that $Pe = 5.0 \times 10^4$ corresponds to a Reynolds number ($Re = V\langle b \rangle/\nu$) of 31.8. Experimental data presented by Nicholl et al. [1999] suggest that inertial forces in a fracture similar to field 1 will begin to

<table>
<thead>
<tr>
<th>Aperture Field</th>
<th>$\langle b \rangle$, cm</th>
<th>$\sigma_b$, cm</th>
<th>$\sigma_b^2$, cm</th>
<th>$\lambda$, cm</th>
<th>$I(\sigma_b^2)/\sigma_b^2$</th>
<th>$\sigma_{macro}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Field 1</td>
<td>0.022</td>
<td>0.0060</td>
<td>0.073</td>
<td>0.035</td>
<td>1.04</td>
<td>0.47</td>
</tr>
<tr>
<td>Field 2</td>
<td>0.022</td>
<td>0.0060</td>
<td>0.073</td>
<td>1.8</td>
<td>1.04</td>
<td>23</td>
</tr>
<tr>
<td>Field 3</td>
<td>0.022</td>
<td>0.019</td>
<td>0.56</td>
<td>0.035</td>
<td>1.32</td>
<td>3.9</td>
</tr>
<tr>
<td>Experimental</td>
<td>0.022</td>
<td>0.0060</td>
<td>0.096</td>
<td>0.044</td>
<td>1.05</td>
<td>0.78</td>
</tr>
</tbody>
</table>

*For the aperture fields used in this study, $R_{bb}(s) = \sigma_b^2 \left[1 - \left(\frac{\pi r^2}{16 \lambda^2}\right)\right] \exp \left[-\left(\frac{\pi r^2}{16 \lambda^2}\right)\right]$ was used to derive these values.

**Figure 7.** Comparison of aperture distributions for experimental fracture and the three synthetic fields (fields 1, 2, and 3). Synthetic fields 1 and 2 have the same variance as the experimental field. The synthetic fields exhibit a positive skew because of the lognormal distribution, and the experimental field exhibits a negative skew. Field 3 has a larger variance and the same mean as the other fields, resulting in a higher percentage of small aperture values.
Influence flow in the range $1 < Re < 10$. Thus these simulations surpass the range in which Stokes flow is expected to be valid. As a result, the simulation results for $Pe > 2 \times 10^4$ should be interpreted with caution. For all the simulations in the three synthetic fields we used line source initial conditions spanning only the middle 2.5 cm ($\sim 70\lambda$) of the fracture. This guaranteed that no particles reached the lateral boundaries before $D_L$ became fully developed. As expected, for simulations at higher velocities (larger $Pe$) the plume required greater travel distances for Taylor dispersion to become fully developed. At the largest values of $Pe$ reported here ($\sim 5.0 \times 10^4$), $dM_{L}/dt$ reached a constant value at a distance of approximately 70 cm ($40\%$ of the length of the field or $\sim 0.06 (b) Pe$).

Figure 8 shows $D_L/(V(b))$ plotted against $Pe$ for simulations in each of the three synthetic aperture fields, together with theoretical results that will be discussed in detail in section 5. At large $Pe$ all three sets of simulations illustrate the progressive dominance of Taylor dispersion, where $D_L \propto Pe^2$. However, simulations in each field exhibit markedly different behavior at intermediate values of $Pe$ (i.e., $\sim 10^0 < Pe < 10^3$). The simulations in field 2 (large $\lambda$) indicate $D_L \propto Pe$ over almost this entire intermediate range of $Pe$ values, whereas the simulations in field 1 indicate almost no range where $D_L \propto Pe$. Simulations in field 3 (large $\sigma_n^2$) indicate $D_L \propto Pe$ in a $Pe$ range that is between the corresponding $Pe$ range in fields 1 and 2. It is clear that as $\sigma_n^2$ and/or $\lambda$ increase, there is a corresponding increase in the magnitude of $\sigma_{macro}$ and hence a larger range of $Pe$ in which $D_L \propto Pe$.

5. Comparison of Theory to Physical Experiments and Computational Simulations

The results of the transport simulations over the range $O(10^0) < Pe < O(10^5)$ (Figure 8) exhibit the three distinct dispersion regimes suggested by Roux et al. [1998]: molecular diffusion, Taylor dispersion, and macrodispersion. In this section we compare the theoretical estimates of $\sigma_{macro}$ and $\sigma_{Taylor}$ obtained from (5) and (6) to estimates from the experimental and computational results.

To calculate the theoretical estimates of $\sigma_{macro}$ for the experimental rough-walled fracture and the three synthetic fields, we applied (6) to estimates of $I(\sigma_n^2)$ calculated for each field by numerically integrating the corresponding covariance function. Figure 9 shows $D_L/D_m$ plotted against $Pe$ for the experimental field (experimental, computational, and theoretical results) and field 1 (computational and theoretical results). For field 1, theoretical and computational results are almost identical, with a discrepancy of $\sim 1\%$ at $Pe = 300$. This indicates that (6) and (7) are accurate in a variable aperture fracture when the underlying assumptions (local cubic law flow, a stationary, lognormally distributed aperture field, and small $\sigma_n$) are satisfied, as is the case for the simulations in field 1. The theoretical estimates of $D_L/D_m$ for the experimental fracture are smaller (by $\sim 9\%$ at $Pe = 300$) than the estimates from the computational results (Figure 9). This discrepancy, which is larger than that observed for field 1, is likely because the aperture field is not lognormally distributed, as this is the primary discrepancy between the assumptions of the theory and the computational simulations in this field. Another possible reason may be that the simulations were carried out in one realization, whereas the theory predicts ensemble average behavior. However, the relatively large source width ($\sim 28\lambda$) for our single realizations leads us to expect closer agreement.
Table 4. Fitted Values of $\sigma_{\text{macro}}$ and $\sigma_{\text{Taylor}}$ for Experiments and Simulations and Theoretical Values Based on Equations (2) and (6)

<table>
<thead>
<tr>
<th></th>
<th>$\sigma_{\text{macro}}$</th>
<th>$\sigma_{\text{Taylor}}$</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Hele-Shaw</strong></td>
<td>NA</td>
<td>0.00476 ± 0.000004</td>
</tr>
<tr>
<td>experiments</td>
<td>NA</td>
<td>0.00472 ± 0.000002</td>
</tr>
<tr>
<td>simulations</td>
<td>NA</td>
<td>0.00476</td>
</tr>
<tr>
<td>theory</td>
<td>NA</td>
<td></td>
</tr>
<tr>
<td><strong>Rough wall</strong></td>
<td>1.87 ± 0.15</td>
<td>0.00522 ± 0.00026</td>
</tr>
<tr>
<td>experiments</td>
<td>1.01 ± 0.10</td>
<td>0.00481 ± 0.00016</td>
</tr>
<tr>
<td>simulations</td>
<td>0.77</td>
<td>0.00476</td>
</tr>
<tr>
<td>theory</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Field 1</strong></td>
<td>0.41 ± 0.20</td>
<td>0.00472 ± 0.00033</td>
</tr>
<tr>
<td>simulations</td>
<td>0.47</td>
<td>0.00476</td>
</tr>
</tbody>
</table>

NA is not applicable. Tolerances are 95% confidence intervals for the fitted parameters.

with the ensemble average theory. The experimental results for $D_I/D_m$ in the experimental field are substantially larger than the theoretical estimates (by ~51% at $Pe = 300$). Of this 51% discrepancy, about 9% can be explained by the deviation between the computational and theoretical results just noted. Thus, by process of elimination, the remaining portion of the discrepancy between theoretical and experimental values (~42%) reflects the influence of deviations from the Reynolds equation in the flow field.

The theoretical value of $\sigma_{\text{Taylor}}$ for each field is $4.76 \times 10^{-3}$. This value is 9% smaller than that obtained by fitting (7) to the experimental results and is 1% less than that obtained from the simulations (Table 4). The small discrepancy between the theoretical $\sigma_{\text{Taylor}}$ and the value estimated from simulations indicates that calculating an effective $D_L, \text{Taylor}$ based on the mean aperture and flow velocity provides a good estimate of the influence of Taylor dispersion in the rough-walled fracture. The small discrepancy between the experimentally measured $\sigma_{\text{Taylor}}$ and the theoretical value indicates that deviations from the Reynolds equation do not influence $\sigma_{\text{Taylor}}$ as much as they influence $\sigma_{\text{macro}}$.

Figure 8 shows $D_I/(V(b))$ plotted against $Pe$ for both theoretical and computational results for fields 1, 2, and 3. There is excellent agreement between the theoretical results and simulations because both employ the Reynolds equation to describe flow through a rough-walled fracture. Also, the aperture fields are lognormally distributed, as assumed in the derivation of the theoretical results. Even though the simulations were carried out in a single realization, the good agreement with the ensemble average theory is probably due to the large source sizes (~70λ) transverse to the mean flow. It may also be noted that the agreement between simulations and theory is poorer in field 3 than in fields 1 and 2. This is likely because $\sigma_{\beta}^2$ is larger for field 3, and there is an assumption of small $\sigma_{\beta}^2$ implicit in the theoretical results.

In summary, comparing theoretical results to experimental and computational results demonstrated the following: (1) excellent agreement between theory and simulations in synthetic, lognormal, and random fields due to the consistency of the assumptions (i.e., both are based on the Reynolds equation); (2) small discrepancies between simulations and theory for the rough-walled experimental fracture, likely due to the assumption of a lognormal aperture distribution implicit in the theory; (3) poorer agreement between theoretical and experimental results, highlighting the additional influence of deviations of the actual velocity field from that described by the Reynolds equation, on estimates of $D_I$; (4) good agreement between theoretical and experimental/computational estimates of $\sigma_{\text{Taylor}}$ indicating that the “effective” Taylor dispersion coefficient defined by $V^2(b)^2/(210D_m)$ is a good estimate of the influence of Taylor dispersion in a variable aperture fracture; (5) good agreement between theory and numerical simulations for fields with different $\sigma_{\beta}^2$ and λ, indicating that when the assumptions of the Reynolds equation are met, (8) effectively characterizes the $Pe$ regimes where different dispersion mechanisms dominate.

6. Concluding Remarks

We have described a series of physical and computational transport experiments designed to clarify the Peclet number ($Pe$) dependence of the longitudinal dispersion coefficient ($D_L$) in a saturated, variable-aperture fracture. During physical experiments in two analog glass fractures (a parallel plate fracture (Hele-Shaw cell) and a rough-walled fracture), aperture and concentration fields were measured using an accurate light transmission technique. Computational simulations in the measured aperture fields were compared to experimental results and the role of aperture variability on the $Pe$ dependence of $D_L$ was investigated through additional simulations in three synthetically generated random aperture fields. The flow fields for the transport simulations were generated by solving the Reynolds equation in each aperture field. The three-dimensional velocity fields within the fractures incorporated parabolic velocity profiles across the aperture, with magnitudes based on the local aperture-averaged velocity obtained from the Reynolds equation solution. A three-dimensional particle-tracking algorithm was then used to simulate solute transport. This approach enabled us to represent the influence of both macrodispersion and Taylor dispersion in contrast to previous approaches, which have isolated one or the other of these mechanisms.

Excellent agreement between simulations and experiments in the Hele-Shaw cell verified our computational model under simple conditions. In the rough-walled fracture both physical and computational experiments demonstrated a nonlinear relationship between $D_L$ and $Pe$. In particular, our results confirmed two of the distinct dispersion regimes suggested by Roux et al. [1998]. At intermediate values of $Pe$, macrodispersion dominated ($D_L \propto Pe$), and at large $Pe$ values, Taylor dispersion dominated ($D_L \propto Pe^2$). We predicted $D_L$ throughout this range using a simple theoretical model that represents the total longitudinal dispersion coefficient as the sum of a macrodispersion coefficient based on the stochastic analysis of Gelhar [1987, 1993] and a Taylor dispersion coefficient based on the mean solute velocity and the mean aperture. This theoretical model, which also incorporates the assumptions inherent in the Reynolds equation, agreed closely with simulations in synthetic, lognormally distributed aperture fields. However, though the theoretical model and numerical simulations captured the $Pe$ dependence of $D_L$ in the experimental fracture quite well, they underestimated the magnitude of $D_L$. We demonstrated, by process of elimination, that this discrepancy is primarily due to inadequacies of the Reynolds equation. Thus the ability of the theoretical model to describe dispersion will be closely tied to the validity of the Reynolds equation in a given fracture.
The theoretical model shows that the statistics of the aperture field determine the specific nature of the Pe dependence of $D_L$. In particular, the Pe ranges in which different dispersion mechanisms dominate vary with aperture statistics ($\langle b \rangle$, $\sigma_{\beta^2}$, and $\lambda$). The theoretical model assumes that the aperture field can be described as a stationary lognormal random field. Brown [1995] suggested that when two rough surfaces are brought together to form a fracture, there is a length scale (mismatch length scale) above which aperture variability is essentially stationary because of correlation between the two surfaces. Thus, although studies of fracture surfaces have demonstrated variability in topography over multiple length scales [e.g., Brown and Scholz, 1985; Poon et al., 1992; Schmittbuhl et al., 1993; Pourahrabou et al., 1995; Power and Tullos, 1995], the largest being of the order of the sample size (up to $O(1)$ m), the aperture field formed by two of these surfaces may not exhibit the same multiscale characteristics. This hypothesis is supported by a number of measurements made of natural fractures (see Table 5), but it is clearly an area that merits further study.

Our analog, rough-walled fracture was statistically homogeneous, with mean and variance within the range measured in these actual fractures. Additionally, the correlation scale $\lambda_\lambda$ (defined here as the separation distance at which the semivariogram reaches a value equal to $(1 - 1/e)$ times the sill value, which is identical to the integral scale in the case of an exponential covariance function), was much smaller than the dimensions of the fracture. This avoided the influence of channeling at scales of the order of the sample size thus allowing meaningful comparison to theoretical results.

For the previously published aperture statistics for fractures in rock samples detailed in Table 5, we calculated the Pe value ($Pe = 210 \sigma_{\text{macro}}^2$) at which the Taylor dispersion and macrodispersion coefficients are equal (next to last column in Table 5). The values of B used for calculating $\sigma_{\text{macro}}^2$ (6) assumed an exponential covariance function; note that the exact shape of the covariance function has only a mild influence on B, through the integral in (4). The corresponding Reynolds number ($Re$) (last column in Table 5) clarifies whether the high velocities associated with macrodispersion/Taylor dispersion equivalence are within the “Darcian” region (i.e., linear relation between the mean hydraulic gradient and the mean flux).

Values of $Re > \sim 10$ imply that rather than the Taylor dispersion regime, a new dispersion regime is likely, where dispersion will be influenced by non-Darcian flow effects resulting from the inertial terms in the Navier-Stokes equations. We see that in the limestone, granite, and welded tuff fractures characterized by Kumar et al. [1995], Yeo et al. [1998], and Wan et al. [2000], respectively, the influence of Taylor dispersion will be evident (for solute transport in water) at $Re \lessapprox 1.0$. Thus there is a range of flow rates for these fractures in which Taylor dispersion will be the dominant dispersion mechanism, just as we have found in our analog, rough-walled fracture. For the granite fractures measured by Hakami and Larsson [1996] and Keller et al. [1999], the value of Pe at which Taylor dispersion will begin to dominate corresponds to $1 < Re < 10$. In these fractures, Taylor dispersion may not become the dominant dispersion mechanism before the flow becomes non-Darcian, the influence of Taylor dispersion will result in a nonlinear relationship between Pe and $D_L$ at high flow rates.

The occurrence of both macrodispersion and Taylor dispersion regimes in fractures can have significant implications on interpreting tracer tests at the single-fracture scale. For instance, in radial flow tracer tests the mean flow velocity increases as solute approaches the well. As a result, macrodispersion can be the dominant dispersion mechanism far from the well, whereas Taylor dispersion effects become increasingly important close to the well. Commonly used interpretation techniques [e.g., Raven et al., 1988; Malozewski and Zuber, 1990; Cady et al., 1993; Novakowski et al., 1995] incorporate only a dispersion coefficient that is a linear function of velocity, implicitly assuming that Taylor dispersion effects are insignificant. The analysis of Hodgkinson and Lever [1983], however, incorporates only Taylor dispersion. Subtle nonlinear Pe dependence of $D_L$ may also be important, when using multiple tracers with widely differing molecular diffusion coefficients. The Pe values corresponding to the different tracers under the same hydraulic conditions can vary over 1–2 orders of magnitude, placing the different tracers in different dispersion regimes. The Pe dependence of $D_L$, however, is typically ignored in interpreting these tracer tests [e.g., Malozewski and Zuber, 1990].

Another important feature that merits additional study is the

### Table 5. Comparison of Aperture Field Statistics

<table>
<thead>
<tr>
<th>Fracture Type</th>
<th>Fracture Statistics</th>
<th>$\sigma_{\beta^2}$</th>
<th>$\lambda_\lambda$</th>
<th>$Pe = 210 \sigma_{\text{macro}}^2$</th>
<th>$Re$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Variety</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Readers et al. [1993]</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Kumar et al. [1995]</td>
<td>limestone</td>
<td>0.027</td>
<td>0.0051</td>
<td>0.036$^e$</td>
<td>0.3</td>
</tr>
<tr>
<td>Keller et al. [1995]</td>
<td>granite</td>
<td>0.038</td>
<td>0.18$^e$</td>
<td>4</td>
<td>16000</td>
</tr>
<tr>
<td>Hakami and Larsson [1996]</td>
<td>granite</td>
<td>0.036</td>
<td>0.16$^e$</td>
<td>0.3</td>
<td>1100</td>
</tr>
<tr>
<td>Keller et al. [1999]</td>
<td>granite</td>
<td>0.083</td>
<td>0.24$^d$</td>
<td>0.6</td>
<td>1580</td>
</tr>
<tr>
<td>Yeo et al. [1998]</td>
<td>granite</td>
<td>0.0607</td>
<td>0.067$^a$</td>
<td>&lt;0.5</td>
<td>460</td>
</tr>
<tr>
<td>Wan et al. [2000]</td>
<td>tuff</td>
<td>0.0281</td>
<td>0.114</td>
<td>0.2</td>
<td>690</td>
</tr>
<tr>
<td>Rough-walled fracture</td>
<td>analog</td>
<td>0.0221</td>
<td>0.073</td>
<td>0.05</td>
<td>140</td>
</tr>
</tbody>
</table>

Ellipsis indicates not available.

*The $210\sigma_{\text{macro}}^2$ is the value of Pe at which Taylor dispersion equals macrodispersion.

This is the Reynolds number corresponding to $Pe = 210 \sigma_{\text{macro}}^2$ assuming $D_m = 1.5 \times 10^{-5}$ cm$^2$/s (typical for commonly used ionic tracers) and $v = 1.0 \times 10^{-2}$ cm/s.

This is based on numerically combining measurements of individual surfaces.

This is based on reported values of $\sigma_{\beta^2}$ assuming a lognormal aperture distribution.

This is based on the reported value of $\sigma_{\text{macro}}^2$.

These are reported values for 0 displacement between fracture surfaces, based on light transmission measurements made in the Flow Visualization and Processes Laboratory, Sandia National Laboratories, New Mexico.
fundamental difference between the Pe dependence of $D_L$ in variable aperture fractures and porous media. Several studies in porous media indicate a linear relationship between $D_L$ and Pe at high Pe ($Pe > 10^3$) [e.g., Fried and Combarnous, 1971]. This linear relationship breaks down only when the Pe value becomes so large that the flow becomes non-Darcian. We believe that this feature relates to the difference in the topology of the void space in fractures and porous media. In most fractures used in previously reported experiments and in our study, the fraction of contact area between the fracture surfaces is relatively small. The resulting long, simply connected flow paths permit Taylor dispersion to "develop." However, because of the complex tortuous topology of the pore space in porous media, there may not be an opportunity for Taylor dispersion to develop, especially at high Pe, where the travel time through a single pore throat will be very small [e.g., Bear, 1972]. Additional computational and experimental studies in fractures with larger fractions of contact area and in porous media would help to clarify the fundamental reasons for the difference of the Pe dependence of $D_L$ between fractures and porous media.

Appendix: Influence of Concentration Measurement Errors on Estimates of $D_L$

As with light transmission measurements of aperture fields, concentration field measurements are subject to random errors due to CCD image noise and accuracy or systematic errors (e.g., nonlinear dye absorbance, reflections, and refraction). Unlike aperture measurements, concentration measurements can be easily calibrated by sequentially filling the fracture with standards of known concentration and fitting (1) to the measurements at each pixel. Calibration effectively minimizes accuracy errors, leaving image noise as the primary error source. The influence of noise can be quantified by measuring the difference between the actual concentration in the fracture during calibration and the concentration calculated using the calibration curves developed at each pixel. Averaging the difference between the measured concentration at each pixel and the actual concentration over the entire field yielded a maximum mean error of 0.003C₀ (C₀ = 0.25 g/L). This indicates that (11) fits the data well and that errors are predominantly due to random signal noise. The root-mean-square (RMS) errors at individual pixels ranged from 0.001C₀ for the 0 g/L fields in both fractures to 0.023C₀ and 0.019C₀ for the 0.25 g/L field in the Hele-Shaw cell and the rough-walled fracture, respectively. These random errors do not influence our estimates of $D_L$, because in calculating $D_L$, we combine measurements from thousands of pixels which significantly reduces the influence of noise.

Despite calibrating our measurement system, the reflective coating on the surface of the band-pass filter on the camera lens caused additional errors in our concentration measurements when the fracture was not filled entirely with one concentration (i.e., during solute transport experiments). These errors included an artificial, reflected plume that was ~1% of the concentration of the real plume and led the real plume through the fracture by ~7.1 cm. We also observed a trend in background concentrations in some fields that resulted in additional mean errors ranging from 0.003C₀ at the inflow end of the fracture to ~−0.003C₀ at the outflow end of the fracture. This trend in background concentrations may be due to a small shift in the intensity of the light source that was not uniform over the entire field. Because the background trend remained relatively constant over the several days that we performed the experiments, this shift appears to be a one-time occurrence after we obtained the calibration images and before we ran the experiments. Because these two additional sources of concentration measurement error were consistent from image to image, we were able to develop an image processing algorithm to minimize the influence of these errors on estimates of dispersion coefficients.

To remove our observational bias and to streamline the processing of over 700 experimental images, we developed a processing routine that consisted of four primary steps (described here in detail): (1) Subtract the reflected plume from each image. (2) Apply a median filter to the entire concentration field. (3) Define a region of interest (ROI) that completely surrounds the plume but excludes small errors far from the plume. (4) Apply a thresholding routine inside the ROI to isolate the plume from any trend in background concentrations within the ROI. We varied the parameters used in this routine to bound the possible values of $D_L$ measured for each experiment. These bounds are displayed as error bars in Figure 5. Note that the error bars represent the possible range of values of $D_L$ not a standard deviation or confidence interval.

We removed the reflected plume from each image by scaling and shifting the real plume appropriately and subtracting it from the concentration field. Using a median filter, which sets the value at each pixel to the median value of a 5 × 5 pixel box surrounding the pixel, we reduced noise and any anomalous measurements from the field. This filtering step smoothed the concentration field slightly, but since the resolution of our measurements (pixel size equals 0.154 × 0.154 cm) was significantly smaller than the scale of real concentration variations within the field, the effect of this smoothing on our estimates of $D_L$ was negligible. We then defined a small ROI centered at the first moment of the real plume and, using (14), measured the second spatial moment of the ROI. Incrementally increasing the size of this ROI resulted in a plot of second moment versus ROI size which at first grew quickly and then leveled off. The degree to which the slopes of plots of $M_z$ versus ROI size leveled off depended on the magnitude and relative location of the trend in the background concentrations. However, for each experimental image the value of $M_z$ at which the slope first began to level off ($M_z_{est}$) was directly related to the size of the plume. We defined an ROI around the plume that had a width and height of 8√$M_z_{est}$ as we found a region of this size captured the entire plume and a buffer around the plume (this corresponds to the size of the third frame shown in Plate 1b). We then obtained the worst case overestimation of $D_L$ by measuring $M_z$ of this entire ROI. We defined a threshold concentration equal to 0.3% of the maximum mass measured in the solute plume from the initial image of each series. By setting all values within the ROI that were less than this threshold to zero, we obtained the worst case underestimation of $D_L$. We obtained the best estimation of $D_L$ by dilating the region defined by the thresholding routine to include pixels with low mass in the vicinity (approximately eight pixels) of the plume that were below the threshold concentration.

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