Pattern formation and coarsening dynamics in three-dimensional convective mixing in porous media

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Pattern formation and coarsening dynamics in three-dimensional convective mixing in porous media

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Geologic carbon dioxide sequestration entails capturing and injecting CO$_2$ into deep saline aquifers for long-term storage. The injected CO$_2$ partially dissolves in groundwater to form a mixture that is denser than the initial groundwater. The local increase in density triggers a gravitational instability at the boundary layer that further develops into columnar plumes of CO$_2$-rich brine, a process that greatly accelerates solubility trapping of the CO$_2$. Here, we investigate the pattern-formation aspects of convective mixing during geological CO$_2$ sequestration by means of high-resolution three-dimensional simulation. We find that the CO$_2$ concentration field self-organizes as a cellular network structure in the diffusive boundary layer at the top boundary. By studying the statistics of the cellular network, we identify various regimes of finger coarsening over time, the existence of a nonequilibrium stationary state, and a universal scaling of 3D convective mixing.

Key words: nonequilibrium flow, cellular network, Rayleigh–Bénard instability, CO$_2$ sequestration.

1. Introduction

Geologic carbon sequestration refers to the capture of carbon dioxide (CO$_2$) from the flue stream of large stationary sources like coal- or gas-fired power plants, and the compression and injection of the captured CO$_2$ into deep geologic strata like deep saline aquifers for long-term storage (IPCC, 2005). It has been proposed as a promising technology for reducing atmospheric CO$_2$ emissions and mitigating climate change (Lackner, 2003; Orr, Jr., 2009; Szulczewski et al., 2012). While
CO₂ is less dense than water for all depths in onshore geologic reservoirs, when CO₂ dissolves into water, the density of water increases. This phenomenon leads naturally to a Rayleigh–Bénard-type, gravity-driven hydrodynamic instability that greatly enhances the rate of dissolution of the CO₂: the mixing of water and CO₂ is controlled by convection and diffusion rather than diffusion alone (Weir et al., 1996; Lindeberg & Wessel-Berg, 1997; Ennis-King & Paterson, 2005; Riaz et al., 2006). This process of CO₂ sinking away as it dissolves in brine—known as solubility trapping—increases the security of geological CO₂ storage in deep saline aquifers (MacMinn et al., 2011; Szulczewski et al., 2012). Convective mixing may also play a role in the dissolution of halites or other soluble low-permeability rocks overlying groundwater aquifers (Evans et al., 1991; Van Dam et al., 2009), leading to high dissolution rates that can exert a powerful control on pore-water salinity in deep geologic formations (Ranganathan & Hanor, 1988; Garven, 1995).

Gravity-driven convection in porous media has been studied extensively (see, e.g., Nield & Bejan, 2006), and has received renewed attention in the context of CO₂ sequestration, including linear and nonlinear stability analysis of the onset of convection (Ennis-King et al., 2005; Riaz et al., 2006; Rappa et al., 2008; Slim & Ramakrishnan, 2010), nonlinear simulations of the unstable flow in two dimensions (Riaz et al., 2006; Hassanzadeh et al., 2007; Hidalgo & Carrera, 2009; Neufeld et al., 2010) and three dimensions (Pau et al., 2010), and experimental systems reproducing the conditions for convective mixing in a stationary horizontal layer (Kneafsey & Pruess, 2010; Neufeld et al., 2010; Backhaus et al., 2011; Slim et al., 2013). Much of the previous work has focused on upscaling the dissolution flux (Pau et al., 2010; Kneafsey & Pruess, 2010; Neufeld et al., 2010; Backhaus et al., 2011; Hidalgo et al., 2012). Here we focus, instead, on the formation of intricate patterns in the diffusion boundary layer as a result of the gravitational instability (Pau et al., 2010; Slim et al., 2013). We describe the entire evolution of the convective-mixing instability in 3D, and the 2D emerging patterns in this boundary layer. We identify and characterize several regimes. We pay especial attention to the emergence of a cellular-network structure, and address fundamental questions on the morphology and dynamics of this pattern: What is the evolution that leads to this pattern morphology?
Does this pattern reach a pseudo steady-state characterized by a universal length scale? If so, how does this length scale depend on the system parameters? What are the mechanisms responsible for this nonequilibrium stationary state? Are the coarsening dynamics also universal? Here, we address these questions using 3D high-resolution simulation of convective mixing in porous media, which—in addition to important visual observations—enable quantitative analysis of the pattern-forming process.

2. Simulating convective mixing in 3D

The equations governing gravity-driven convective mixing are the Darcy–Boussinesq equations of variable-density flow in porous media, which for a homogeneous porous medium, and in dimensionless form, are (Riaz & Meiburg, 2003; Riaz et al., 2006):

\[ \nabla \cdot \mathbf{u} = 0, \quad (2.1) \]

\[ \mathbf{u} = - (\nabla P' - C \hat{z}), \quad (2.2) \]

\[ \frac{\partial C}{\partial t} + \nabla \cdot (\mathbf{u} C - \frac{1}{Ra} \nabla C) = 0. \quad (2.3) \]

Equation (2.1) is the incompressibility constraint, Eq. (2.2) is Darcy’s law, and Eq. (2.3) is the advection–diffusion equation governing solute transport. The computational domain is the unit cube \([0, 1]^3\), made dimensionless with respect to a length scale \(H\) taken here to be the depth of the porous layer. In the equations above, \(\mathbf{u}\) is the dimensionless Darcy velocity, \(C\) is the normalized concentration of CO\(_2\) dissolved in water, \(P'\) is the dimensionless pressure with respect to a hydrostatic datum, and \(\hat{z}\) is a unit vector pointing in the direction of gravity. The density of the groundwater–CO\(_2\) mixture is a linear function of the CO\(_2\) concentration: \(\rho = \rho_0 + \Delta \rho C\), where \(\rho_0\) is the density of the ambient brine and \(\Delta \rho\) is the density difference between CO\(_2\)-saturated groundwater and CO\(_2\)-free groundwater. The only controlling parameter of the system is the Rayleigh number,

\[ Ra = \frac{\Delta \rho g k H}{\phi D \mu}, \quad (2.4) \]
where $k$ is the intrinsic permeability, $\phi$ is the porosity, $g$ is the gravitational acceleration, $\mu$ is the fluid dynamic viscosity, and $D$ is the diffusion–dispersion coefficient.

The boundary conditions are no-flow in the $z$-direction and periodic in the $x$- and $y$-directions. We impose a fixed concentration at the top boundary of the cube ($z = 0$), $C(x, y, z = 0, t) = 1$, to simulate contact with buoyant free-phase CO$_2$. Initially, the CO$_2$ concentration is zero almost everywhere. We trigger the density-driven instability by introducing a small perturbation on the initial condition. For fixed $(x, y)$ coordinates, concentrations along the vertical axis follow an error function, quickly approaching $C = 1$ and $C = 0$ above and below the front, respectively. We perturb the front by vertical shifting the isoconcentration contours using a small white-noise perturbation (an uncorrelated Gaussian random function). We have confirmed that our results are independent of the precise magnitude of the perturbation.

We solve equations (2.1)–(2.3) sequentially: at each time step, we first update the velocities, and with fixed velocities we update the concentration field. We adopt the stream function–vorticity formulation of equations (2.1)–(2.2) (Tan & Homsy, 1988; Riaz & Meiburg, 2003). The components of the stream vector are solved for with an eighth-order finite difference scheme, implemented as a fast Poisson solver (Swarztrauber, 1977). For the transport equation (2.3), we use sixth-order compact finite differences (Lele, 1992) in the vertical direction, and a pseudo spectral (Fourier) discretization along the horizontal directions, which we assume to be periodic. We integrate in time using a third-order Runge-Kutta scheme with automatic time-step adaptation (Ruith & Meiburg, 2000).

### 3. Results

We solve the governing equations for Rayleigh numbers up to $Ra = 6400$ on a grid of $512^3$, for which we have approximately 400 million degrees of freedom to be solved at each time step. We have confirmed that the results from the simulations are converged results and, therefore, independent of grid size. In this section,
Figure 1. (Online version in color.) Simulation of convective mixing with Ra = 6400 on a 512³ grid. (a) Snapshot of the concentration field at a slice near the top boundary (z = 0.01) at t = 0.5, showing a pattern of disconnected islands of high concentration. (b) Snapshot of the same slice at t = 1, showing a partially-connected maze structure. (c)-(e) Snapshot of the 3D concentration field at t = 2; (c) is a complete view of the computational domain; (d) is a view of a partial volume (0.01 < z < 0.3) from the top, illustrating the cellular network structure that emerges at the boundary layer; (e) is a view of the same volume from the bottom, illustrating the columnar pattern of CO₂-rich fingers that sink away from the top boundary. See also Movies S1 and S2 in Supplementary Material.

we describe the 3D dynamics of the system and, in particular, the 2D emerging patterns at the top boundary layer.

(a) Pattern formation

The fixed concentration C = 1 at the top boundary leads to a Rayleigh–Bénard-type hydrodynamic instability, in which the initial diffusive boundary layer becomes unstable and gives rise to gravity-driven convection. In our simulations, we perturb the initial concentration with random uncorrelated Gaussian noise to accelerate the onset of this instability. This diffusive boundary layer then reflects a series of patterns that evolve in time.
1. Islands. During the very early stages of the instability, the minute perturbations of the boundary concentrations give rise to protrusions such that a wavy 3D isoconcentration surface develops. A cut near the top boundary reflects these protrusions in the form of disconnected islands of higher concentration, surrounded by a sea of near-zero concentration [Fig. 1(a)]. Our high-resolution simulations illustrate the columnar pattern in this initial regime of the instability, with a characteristic length that is in good agreement with the predictions of a linear stability analysis, $l_{\text{onset}} \sim Ra^{-1}$ (Riaz et al., 2006).

2. Maze. The initial columnar pattern morphs by developing bridges between the islands, giving rise to an increasingly connected maze structure [Fig. 1(b)]. The emergence of the maze pattern observed in 3D is not obvious from the 2D simulations: it is unclear how the bridges between fingers observed in 2D would self-organize in the third dimension. Our 3D simulations show that the bridges connect to form a maze that later develops into an hexagonal cellular network.

3. Cellular network. The maze structure evolves in two ways: making its walls thinner, and reorganizing itself in space to form a globally connected polygonal network of cells of near-zero concentration separated by sheets of high concentration [Fig. 1(d)]. The thinning process of cellular walls is controlled by the balance between vertical downward advection through the wall and lateral diffusion within the cell, similar to the diffusion-and-advection controlled boundary layer (Riaz et al., 2006). A careful analysis indicates that the thickness of the boundary layer and the thickness of the cell wall both scale with $\sim Ra^{-1}$. Underneath the diffusive layer, the nature of this pattern is different. The vertices of the cellular network are the locations of maximum downward flux of CO$_2$, and this leads to a columnar pattern of CO$_2$-rich fingers that sink [Fig. 1(e)]. However, finger roots exhibit faster temporal dynamics (due to horizontal zipping and merging) than the long-lived fingers in the interior. Thus, while the boundary-layer network contributes to the organization of the interior region, the
Figure 2. Concentration field at $t = 10$ for the 3D simulation with Ra=6400, at different depths. (a) $z = 0.001$, (b) $z = 0.04$, (c) $z = 0.12$, and (d) $z = 0.43$.

morphology and the evolution of the characteristic scale in the interior
do not correspond to those of the network structure at the boundary layer
(Fig. 2) (Backhaus et al., 2011; Slim et al., 2013; MacMinn & Juanes, 2013).

(b) Coarsening dynamics

Once it has been formed at $t \approx 2$, the cellular network coarsens through
merging and collapsing of small cells while columnar fingers migrate downward
[Fig. 1(e)]. This early-time coarsening regime persists until $t \approx 8$, when the
characteristic size of the cells reaches a nonequilibrium stationary state. This
statistical steady state lasts for an extended period of time during which two
mechanisms act to balance the characteristic size of the cells.
1. **Cell growth.** In the first mechanism, small cells in the network progressively shrink and large cells expand. The shrinking cells eventually vanish from the network, leaving space for large cells to grow. To understand this coarsening process, one must consider the velocity field induced by convection. Cell centers correspond to upwelling currents of fresh fluid that impinge onto the boundary layer and deviate laterally towards the cell edges, charging themselves with CO$_2$ in the process, and then migrating downwards at the cell edges. Cell coarsening is due to a positive feedback, in which larger cells promote larger vertical upward flow, which then tend to push the cell edges outwards, causing the cell size to increase (Fig. 3).

2. **Cell division.** The inflating large cells then trigger the second mechanism, in which new cell boundaries are born in the middle of large cells. The newborn links are often immediately pushed sideways towards existing cell boundaries; however, past a certain cell size, some newly-born sheets persist to give rise to cell boundaries and permanently divide the mother cells (Fig. 3).

The first mechanism promotes cell growth while the second mechanism penalizes oversized cells. These two mechanisms emphasize the nonequilibrium nature of the convective mixing process. At long-enough times ($t \approx 20$), the domain starts to become saturated with CO$_2$, and the influence of the bottom boundary is felt at the top boundary. After this time, the cellular network can no longer sustain its characteristic size and enters a regime of late-time coarsening.

To demonstrate quantitatively the existence of these three periods (early-time coarsening, nonequilibrium stationarity, and late-time coarsening), we plot the power spectrum density $E(k)$ of the concentration field at a slice near the top boundary ($z = 0.01$) for the system with $Ra = 6400$, at various times (Fig. 4). We confirmed that the network patterns are isotropic by analyzing the 2D Fourier transform of the network images, which indeed exhibit concentric circular isocontours in all cases. Thus, we define the 2D isotropic horizontal wavenumber $k$ as $k^2 = k_x^2 + k_y^2$, where $k_x$ and $k_y$ are the wave numbers in $x$- and $y$-directions, respectively. Note that from our definition of the wavenumber, the corresponding
Figure 3. (Online version in color.) (a) Snapshot of the velocity field at a depth \( z = 0.01 \) at time \( t = 13 \) for \( \text{Ra} = 6400 \), showing upward flow at the cell centers (grayscale) and downward flow at the cell edges (white), and horizontal flow from the center to the edges of individual cells (red arrows). (b) Zoomed view of a small area of the same slice (blue square) at different times, illustrating cell growth and disappearance of small cells (\( t = 11.6 \) to \( t = 11.7 \)), and cell division from the emergence of sheets of high concentration within cells (\( t = 11.7 \) to \( t = 11.8 \)).
Figure 4. (Online version in color.) Evolution of the power spectrum density for the concentration field of a horizontal slice ($z = 0.01$) of the simulation with $Ra = 6400$. The onset wave number inferred from the numerical simulations is $k \approx 40$, corresponding to the maximum energy content for the solution at $t = 0.2$. While this number should be understood as a plausible range rather than a hard value, it does agree nicely with the result of a linear stability analysis (as extrapolated from Fig. 11 in Riaz et al. (2006)).

The length scale is $1/k$ (and not $2\pi/k$). The power spectrum density is calculated using the square of the 2D Fourier transform of the concentration field. Initially, there is a shift in the maximum of the power spectrum towards lower wavenumbers, indicating an increase in the characteristic length (red curves, corresponding to $t = 0.2$ and $t = 1$). Later, for a wide range of times, the power spectra at different times exhibit perfect overlap, strongly suggesting a statistically stationary state (blue curves, $t = 10$ and $t = 14$). At later times, the power spectrum decays more rapidly at higher wavenumbers, indicating that the smaller cells are removed from the system (black curves, $t = 16$ and $t = 22$).

We confirm the transition from an early-time coarsening to a statistical steady state by evaluating the representative cell length of a network,

$$l_{\text{cell}} = \frac{1}{\sqrt{N_{\text{fing}}}}$$  \hspace{1cm} (3.1)

where $N_{\text{fing}}$ is the number of fingers that root within the network, which corresponds to the number of network joints [Fig. 5(a)]. We assume that the number of joints is linearly related to the number of cells in the network—an
Figure 5. (a) Snapshot of the concentration field at $t = 10$, $z \approx 0.01$ for a 3D simulation with $Ra=6400$. (b) The dark lines mark the binary skeleton representation of the same network shown in (a). The red circles are the network joints identified by image processing tool. (c) Snapshot of the concentration field near the top boundary of a 2D simulation with $Ra=10000$ at $t = 10$. The black dotted line indicates $z \approx 0.005$, the depth at which we extract the 1D concentration. (d) The black solid line is the 1D concentration signal obtained from (c); the red circles are the peaks identified by the peak-finding tool.

From this observation, we propose to estimate the average cell area $A_{\text{cell}} \sim l_{\text{cell}}^2$ as proportional to the total area of the network ($1 \times 1$ square) divided by the number of joints ($N_{\text{fing}}$).

A plot of $l_{\text{cell}}$ as a function of time illustrates the growth of the characteristic length scale during an initial period ($t < 8$), and a fluctuating, mean-reverting length scale during the quasi-steady period ($8 < t < 20$) (Fig. 6). The details of this analysis are discussed in section (c) below.
Figure 6. Time evolution of cell size ($l_{cell}$) in 3D simulations for different Rayleigh numbers. The two dashed lines indicate the time averaging window ($10 < t < 15$) used to calculate the characteristic cell length during the nonequilibrium steady state regime of the network.

The characteristic length in the system exhibits three dynamic regimes: early-time coarsening, nonequilibrium steady state and late-time coarsening. It is natural to ask whether the coarsening regimes of the length scale near the boundary layer are reflected in the time evolution of dissolution flux. Indeed, the dissolution flux exhibits three dynamic regimes as well: diffusive, convection-dominated and saturation (Pau et al., 2010; Hidalgo et al., 2012; Slim et al., 2013; Hewitt et al., 2013). Here we compare these two quantities—characteristic length scale and dissolution flux—for both a 3D simulation with Ra=6400 and a 2D simulation with Ra=25,000 (Fig. 7). The dynamics of these two quantities appear to be highly correlated in time. The magnitude of the dissolution flux, however, is uninformative with respect to the length scale. The nondimensional flux is independent of Ra (Hidalgo et al., 2012), and clearly this is not the case for the characteristic length scale (Fig. 6).

(c) Universality of coarsening dynamics

The fact that the characteristic length scale of the process reaches a stationary value during an extended period of time raises the question of what sets that length scale. Our hypothesis is that, in the absence of any external length scale in
Figure 7. Time evolution of non-dimensional flux (blue) and cell length near the boundary (green). (a) 3D simulation with $Ra=6400$. (b) 2D simulation with $Ra=25,000$. 
the problem, this characteristic length is set by a balance between advection and
diffusion, \( l_{\text{diff}} \sim D/U \), where \( U = (\Delta \rho g k) / (\phi \mu) \) is the characteristic density-driven
fluid velocity. From the definition of the Rayleigh number, Eq. (2.4), we have that
\( l_{\text{diff}} \sim H/Ra \). This suggests a linear scaling of cell size with the inverse of Ra,

\[
l_{\text{cell}} \sim Ra^{-1}
\]  

To test this hypothesis, we perform a study of the evolution of cell sizes of the
network. We threshold the concentration field to obtain a binary image that can
then be reduced to a skeleton representation of the network [Fig. 5(b)], using open-
source image processing software (Schneider et al., 2012). We count the number
of vertices, or joints, in the skeleton network using a commercially available image
processing tool (Matlab, 2012), and then estimate the cell length \( l_{\text{cell}} \) defined in
Eq. (3.1).

In Fig. 6 we plot the time evolution of \( l_{\text{cell}} \) for nine different Rayleigh numbers,
ranging from 1600 to 6400. We identify the three coarsening regimes described in
section 2(b), although finite-size effects prevent achieving the pseudo-steady state
for the smaller values of Ra (1600 and 2000). We choose the overall characteristic
length, denoted \( \bar{l} \), as the time average of \( l_{\text{cell}} \) during the nonequilibrium stationary
state, taken here as \( 10 < t < 15 \). This average length scale \( \bar{l} \) exhibits a power-law
dependence with Rayleigh number, with exponent \(-1\) [Fig. 8(a)], supporting the
scaling relation in Eq. (3.2).

We recognize that it would be useful to extend the study of 3D convective
mixing to higher Rayleigh numbers. However, the computational cost would be
significant. Instead, we confirm the proposed scaling \( \bar{l} \sim Ra^{-1} \) with 2D simulations,
where it is computationally tractable to perform simulations with \( Ra = 40,000 \). In
2D, the domain is the unit square \((1 \times 1)\), \( N_{\text{fing}} \) is the number of finger roots in the
boundary layer [Fig. 5(c)], and the characteristic length is the average finger root
spacing: \( l_{\text{cell}} = 1 / N_{\text{fing}} \). We use a robust peak-finding tool (Yoder, 2009) to identify
the number of finger roots, which are the peaks in a 1D concentration signal
[Fig. 5(d)] taken near the boundary [Fig. 5(c)]. In Fig. 8(b), we plot the time-
averaged 2D characteristic length \( \bar{l} \) with Ra in log–log scale, and again observe the
Figure 8. Characteristic length $\bar{l}$ plotted against Rayleigh number. (a) 3D simulations; (b) 2D simulations. This characteristic length scale exhibits a power-law dependence with Rayleigh number $\bar{l} \sim Ra^{-1}$.

The same $-1$ exponent. This strongly suggests that the scaling relation $l_{\text{cell}} \sim Ra^{-1}$ is universal, both in 2D and 3D, in the regime of large Rayleigh numbers.

4. Discussion

In this paper, we have studied the pattern-formation aspects of convective mixing in porous media, a phenomenon of relevance in CO$_2$ sequestration in deep saline aquifers. We have analyzed the process by means of high-resolution simulations in a simplified geometry. Our key observation is the emergence of a cellular network structure in the diffusive boundary layer at the top boundary. Theoretical arguments and statistical analysis of the evolving pattern allowed us to discern the fundamental scaling properties of this pattern in space and time. In particular, we have identified a period of coarsening followed by a nonequilibrium steady state, and explained the detailed mechanisms—cell growth and cell division—responsible for this behavior.

We are currently investigating how the detailed 3D simulations and theory presented here may guide the development of nonequilibrium 2D models of the pattern-forming process, in the spirit of surface-growth models (e.g., Kardar et al.,
1986; Barabási & Stanley, 1995). This will inform our ability to model and predict
the properties of other pattern-forming processes that lead to cellular structures
(Stavans, 1993), such as foams (Weaire & Hutzler, 1999), elastocapillary assembly
(Chakrapani et al., 2004), desiccation cracks (Shorlin et al., 2000), columnar
jointing (Goehring et al., 2006, 2009) and mantle dynamics (Tuckley, 2000).

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