Causes and consequences of flow organization during melt transport: The Reaction Infiltration Instability in compactible media

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Abstract. Geochemical and field evidence suggest that melt transport in some regions of the mantle is localized into meso-scale “channels” that have widths of 0.1–100 meters or larger. Nevertheless, the mechanisms for formation of such channels from a grain-scale distribution of melt are poorly understood. The purpose of this paper is to investigate one possible mechanism for channel formation: the Reaction Infiltration Instability (RII). We present numerical solutions of the full equations for reactive fluid flow in a viscously deformable, permeable medium. We show that dissolution in a compactible solid with a vertical solubility gradient can lead to significant flow localization such that over 90% of the melt flux is channelized in less than 20% of the available area. In particular, the ability of the solid to compact enhances the instability by forming impermeable regions between channels. The combination of reaction, diffusion and solid compaction leads to strong selection of preferred length scales with channel spacing smaller than the compaction length ($\delta \sim 10^2–10^4$ m). We explore the evolution of dissolution channels over parameter space and show that the behaviour of the full non-linear solutions is consistent with predictions from linear stability analysis. We also briefly consider the behaviour of the instability in the presence of melting due to adiabatic decompression and demonstrate that significant localization can occur even in the presence of uniform melting and compaction. Using the linear analysis to extend these results for parameters expected in the Earth’s mantle suggests that robust channel systems could form through the RII from a homogeneous system in $\sim 100,000$ yr with dominant channel spacing of 1–200 m.

Introduction

Understanding how melt flow is organized on scales ranging from the grain scale to the size of the partially molten region is a crucial problem in mantle and magma dynamics because the local distribution of melt affects the rates of melt transport, the composition of mantle melts, and the bulk rheology and dynamics of partially molten regions. There is geochemical and field evidence that, although melting begins on the grain-scale, melt transport must somewhere organize into “channels” that could range in scale from 10cm spaced veins [Spiegelman and Kenyon, 1992] to episodic dikes [Sleep, 1988; Nicolas, 1989, 1990] to multiscale “fractal trees” [Hart, 1993]. Nevertheless, the causes and consequences of channel formation remain virtually unexplored. The purpose of this paper is to investigate the...
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physics, scaling and observable consequences of one possible mechanism, the Reaction Infiltration Instability [e.g. Chadam et al., 1986; Ortoleva et al., 1987; Aharonov et al., 1995], where high-porosity channels form by a feedback between melt flow and corrosive dissolution. Previous authors have considered the behaviour of this instability in rigid porous media and multi-component water rock systems [e.g. Steefel and Lasaga, 1990, 1994; Aharonov et al., 1997] and have demonstrated it experimentally [Hoefner and Fogler, 1988; Daccord, 1989; Kelemen et al., 1995b]. Here we extend the problem to viscously deformable media, appropriate to the mantle, through a series of theoretical calculations for a simplified physical system.

The principal geochemical argument for flow-localization in the mantle comes from the observation that mid-ocean ridge basalts (MORB) are strongly out of chemical equilibrium with residual peridotites by the time they reach the crust/mantle transition zone. In terms of major elements, MORB is not saturated in orthopyroxene (opx) at Moho pressures. However, opx is a major constituent of all residual mantle peridotites (~20%) [e.g. O’Hara, 1965; Stolper, 1980] and dissolves very rapidly in undersaturated mantle melts [Kuo and Kirkpatrick, 1985a, b; Brearley et al., 1986; Zhang et al., 1989]. Thus the observation that MORB appears to have last equilibrated with mantle peridotite at a pressure of 1.0 GPa or more suggests that melts have traveled through at least the top 30km of oceanic upper mantle without equilibrating with residual mantle peridotite. Similar arguments can be made from the observation that MORB is also not in trace element equilibrium with most abyssal peridotites [Johnson et al., 1990; Johnson and Dick, 1992]. It is usually interpreted to be a mixture of melts produced from mantle peridotite over a range of pressures that preserves information about pressures greater than that at the base of the crust [Klein and Langmuir, 1987; Salters and Hart, 1989].

These observations place important constraints on the melt transport process and are usually explained by a near-fractional melting process beginning at some depth, with only limited interaction between melt and residue during transport. While the depth at which disequilibrium transport begins is still an open question, Spiegelman and Kenyon [1992] showed that a minimum requirement for disequilibrium transport is for melts to be transported in veins or channels spaced \( \geq 10 \text{cm} \) apart assuming negligible inter-channel porosity (or \( \text{e.g.} > 1 \text{~m} \) apart for an inter-channel porosity of 0.1%).

Beyond chemical arguments, there is also in-situ field evidence for flow localization preserved in ophiolites which record melt migration features from both the adiabatically rising asthenosphere and the conductively cooled lithosphere [e.g. Nicolas, 1989, 1990; Kelemen et al., 1995a; Ceuleneer et al., 1995]. Kelemen et al. [1995a] provide a brief summary of these different features and identify a set of “replacive dunite” bodies in the Oman ophiolite as melt conduits [see also, Hirschmann, 1995]. These dunites are regions where all of the orthopyroxene has been dissolved away, leaving a relatively unreactive olivine residue. The most important observation is that small, rare clinopyroxenes (cpx) in the dunites are chemically in equilibrium with MORB-like liquids, such as those that formed the igneous crust in the ophiolite, while similar pyroxenes in the adjacent harzburgites are not. These harzburgites are strongly depleted in light rare earth elements [Kelemen et al., 1995a] and have pyroxene compositions similar to pyroxenes in abyssal peridotites [Johnson et al., 1990; Johnson and Dick, 1992; Dick and Natland, 1996; Ross and Elthon, 1995].

The sharp geochemical contrast between residual harzburgites and replacive dunites suggests that the dunites are conduits for melt transport. Similar reactive dunites are seen in other ultramafic bodies and occur on scales from 0.1–100m [e.g., Quick, 1981; Kelemen et al., 1992; Nicolas, 1989; Kelemen and Dick, 1995]. Some of these dunites are associated with shear zones and dikes, but others have contact relationships which indicate that they formed as purely porous dissolution channels. While the dunites are clearly a locus for melt migration, questions remain as to how they form and how much melt has been channeled through them.

Previously suggested mechanisms for flow localization fall into two end-member categories. The first set of instabilities are purely mechanical, where local variations in matrix strength lead to flow localizations. These mechanisms include nucleation of “veins” into dikes or hydrofractures [Nicolas, 1989, 1990; Sleep, 1988; Connolly and Podladchikov, 1998] and the localization of melt by viscous shear of a material with porosity weakening viscosity [Stevenson, 1989; Richardson, 1998; Hall and Parmentier, 1998]. An alternative physico-chemical mechanism, “the reactive infiltration instability” (RII) [Chadam et al., 1986; Ortoleva et al., 1987] has also been suggested as a means of melt extraction [Kelemen et al., 1995b; Aharonov et al., 1995, 1997]. In this mechanism, channeling arises from a feedback between dissolution and melt flux. In a system where the solubility of a mineral phase (e.g. pyroxene) in the melt increases with decreasing pressure in the melting column [e.g. Figure 3 Kelemen et al., 1995b], regions of larger melt flux cause increased dissolution which increases the permeability, which in turn increases the flux in a runaway process.

Both mechanical and reactive instabilities are expected to occur in the mantle and should probably reinforce each other; however, because the dunite channels are clearly as-
sociated with dissolution features, in this study we begin by asking how much localization can be attributed to the reaction infiltration instability alone. The purpose of this paper is to investigate the simplest fully non-linear problem appropriate for understanding the consequences of reactive flow in a compactible matrix. The discussion will consider additional process and compare our results to other studies.

Model description

The simplest problem that demonstrates much of the physics of flow in reactive, deformable media is the flow of a solvent through a static (i.e. not upwelling) but compactible medium with a single soluble phase, in which the solubility of the phase increases with decreasing pressure (see Figure 1). This problem is discussed in detail in Aharonov et al. [1995] and is similar to that calculated for a rigid media by Aharonov et al. [1997]. This problem is also related to problem of flowing water through a box of salt and glass beads as modeled by Kelemen et al. [1995b].

Governing Equations

Appendix A provides a general derivation of the equations for reactive flow in viscously deformable porous media based on the original formulation of McKenzie [1984]. For the problem considered here, the equations can be written for four scalar variables: the porosity \( \phi \), the excess fluid pressure \( P \) (or compaction rate \( C \)), and the concentration of the soluble phase in the solid, \( c^s \) and the melt, \( c^f \). The dimensionless equations can be written

\[
\frac{D_s \phi}{Dt} = (1 - \phi_0 \phi)C + \Gamma
\]

(1)

\[
-\nabla \cdot k_0 \nabla P + \frac{P}{(\zeta + 4\eta/3)} = -\nabla \cdot [k_0(1 - \phi_0 \phi)k] + \frac{\Delta \rho}{\rho_f} \Gamma
\]

(2)

\[
\frac{D_s c^s}{Dt} = \frac{-\phi_0}{(1 - \phi_0 \phi)}(c^f_{eq} - c^s) \Gamma
\]

(3)

\[
\frac{D_f c^f}{Dt} = \frac{1}{\rho_f} \nabla^2 c^f + \frac{\rho_s}{\rho_f} (c^f_{eq} - c^f) \Gamma
\]

(4)

where \( \phi \) is the volume fraction of melt present (\( \phi_0 \) is a reference porosity), \( C = \nabla \cdot \mathbf{V} \) is the “compaction rate” i.e. the rate of expansion or compaction of the solid phase [see Spiegelman, 1993a]. The compaction rate is related to the excess fluid pressure by

\[
P = (\zeta + 4\eta/3)C
\]

(5)

where \( (\zeta + 4\eta/3) \) is the viscosity that controls volume changes of the solid matrix. Additional notation is given in Table 1.

Figure 1. Cartoon showing geometry of reactive flow model for a static but compactible matrix. In this problem, the solution domain is \( 4 \times 5 \) compaction lengths (129 \( \times \) 161 grid points) where the upper \( \delta \) are reactive with \( c^f_{eq} \) increasing linearly from 0 to 0.1. The lower boundary is near constant flux inflow for melt with dimensionless \( \phi = 1, w = 1 \), upper boundary is free-flux outflow and the sides are periodic (wrap-around) boundaries. In this problem the solid velocity is \( \mathbf{V} = 0 \) to zeroth order. Initial conditions are \( \phi = 1, c^f = c^f_{eq} \) and \( c^s = 0.95 \pm \epsilon \) where \( \epsilon \) is Gaussian white noise with variance of \( 10^{-4} \). Inset graph shows the equilibrium solubility curve \( c^f_{eq}(z) \) and the initial dissolution rate curve which for large values of \( Da \) approaches \( \frac{\partial c^f_{eq}}{\partial z} = 0.025 \) in the reactive zone.

The full scaling of these equations is given in Appendix A but, for reference, all distances are scaled by the compaction length \( \delta \), which depends on the permeability and the ratio of solid and melt viscosities (see Eq. A14). The compaction length governs the distance over which fluid pressure variations are transmitted through the solid phase [see Spiegelman, 1993a, b]. In rigid media, \( \delta \rightarrow \infty \); however, estimates for the compaction length in the mantle are \( \delta \sim 10^2-10^4 \) m. In Eqs. (1)–(5), time is scaled by the time it takes the melt to move one compaction length at the reference melt velocity \( w_0 \) (Eq. A15). For the problems in the next section, however, it is more convenient to scale by the time it takes melt to cross the entire system. For reference, estimates of melt velocities in mantle systems range from \( 1-100 \) m yr\(^{-1}\) (or faster) and thus transit times for a 60km high melting region are \( \sim 600-60,000 \) years.
Γ is the rate of mass transfer from solid to liquid which, for this problem, is the dissolution rate of the soluble phase. For tractability, and high resolution, we neglect multi-component effects [see Steefel and Lasaga, 1994] and assume linear kinetics such that the dimensionless dissolution rate can be written

$$\Gamma = DaA'(c^f_{eq} - c^f)$$

where

$$Da = \frac{R \delta}{\rho_s \phi_0 w_0}$$

$$A' = \frac{c^s(1 - \phi_0 \phi)}{c^s_0(1 - \phi_0)}$$

Da is the Damköhler number, which governs the amount of reaction that occurs in the time it takes the fluid to move one compaction length. \(R\) is the reaction rate constant (in \(\text{kg m}^{-3}\text{s}^{-1}\)) and \(w_0\) is the melt velocity at porosity \(\phi_0\). A large Damköhler number implies rapid reaction relative to transport time scales, while a small \(Da\) implies little reaction. \(A'\) is the available surface area of the soluble phase (and goes to zero when the phase is exhausted). \(c^f\) is the concentration of the soluble phase in the liquid and \(c^f_{eq}\) is the equilibrium solubility of that phase. For this problem, the dissolution rate depends on the Damköhler number and the distance from equilibrium of the melt composition. For undersaturated melts, \(c^f < c^f_{eq}\), \(\Gamma > 0\) and the solid dissolves. \(\Gamma < 0\) implies precipitation from a supersaturated melt. The permeability of the solid phase, \(k_\phi = \phi^n\) is assumed to be a power-law in porosity. \(c^f_0\) is the concentration of the soluble phase entering the fluid due to the dissolution reaction (and here is just 1) and

$$Pe = \frac{D^f \delta}{w_0}$$

is the Peclet number which controls the amount of diffusion or dispersion that occurs in the time it takes to advect one compaction length. A large Peclet number implies negligible diffusion.

Finally, the expressions \(D_s/Dt\) and \(D_f/Dt\) are the material derivatives for the solid and melt respectively. For example

$$\frac{D_f c^f}{Dt} = \frac{\partial c^f}{\partial t} + \mathbf{v} \cdot \nabla c^f$$

is the time rate of change of melt concentration in a frame traveling at the melt velocity

$$\mathbf{v} = \mathbf{V} - \frac{k_\phi}{\phi} \left[ \nabla P - (1 - \phi_0 \phi) \mathbf{k} \right]$$

For small porosities, and no imposed solid flow field, the solid velocity in this problem is approximately zero. (Appendix A provides a more general derivation).

Equation (1) governs the time rate of change of porosity which is controlled by the balance of physical compaction (or decompaction) and dissolution of the solid as measured in the frame of the solid. Equation (2) governs the excess fluid pressure (or compaction rate) due to variations in melt flux driven by gravity (first term RHS) and the volume change on melting. This equation is discussed in detail by Spiegelman [1993a, c] and controls both compaction and the excess pressure gradients required to drive horizontal flow (e.g. Eq. (10)). Equation (3) governs the change in concentration of the soluble phase in the solid. Equation (4) controls the change in melt concentration as measured in a frame moving with the melt.

Because these equations are dominated by advective terms and the balance between advection, diffusion and reaction, they require highly accurate advection schemes. Here we have used a semi-implicit semi-Lagrangian algorithm [Staniforth and Cote, 1991] for the purely advective equations and a modified 2-D semi-Lagrangian Crank-Nicholson scheme for the combined advection-diffusion-reaction problem. Time stepping uses an iterative predictor-corrector method which converges rapidly. These schemes are unconditionally stable and have no Courant stability condition [see e.g. Press et al., 1992]. Thus, time steps and spatial resolution are decoupled. Accuracy tests for the static runs suggest that time steps of \(2-4\) times the Courant condition are good values. At Courant numbers greater than \(4\), a standing wave artifact becomes noticeable in the static runs (although it does not change the overall physics) and we use a Courant number of \(2\) for the runs shown here. This artifact does not appear in the upwelling melting runs (Fig. 9) and for these we use a Courant number of \(4\). In 2-D, Crank-Nicholson style differencing results in large sparse sets of simultaneous linear equations as does the differencing of the elliptic equation (2). These equations are solved rapidly with multi-grid methods, which scale linearly with the number of gridpoints. The combination of multi-grid and semi-Lagrangian techniques allows efficient solution of these equations on very fine uniform grids in reasonable time.

**Model geometry and details**

Figure 1 shows the geometry and setup for the model we will investigate. For moderate resolution runs, we use a box that is \(4\) compaction lengths wide by \(5\) high with a reactive zone that occupies the upper \(4\) compaction lengths of the box. The bottom compaction length of the box is an unreactive buffer to minimize the influence of the lower boundary. Figure 1b shows the initial equilibrium solubility \(c^f_{eq}\) as a function of height. In the reactive zone the dimensionless equilibrium solubility is a linear function of pressure given...
by
\[ c^f_{eq} = c^f_{eq} \left( (z-1) - P \right). \] (11)

Thus, solubility increases nearly linearly with height \( z \), but excess fluid pressure causes a decrease in solubility at a given depth. The solubility gradient of pyroxene in the mantle is approximately 1%/kb [Kelemen et al., 1995b, Figure 3] such that the maximum solubility over a melting column of height 60km is approximately 20%. We use a maximum solubility of 10% (i.e. \( c^f_{\text{max}} = 0.025 \) per compaction length) for the static runs and 20% for the melting runs. The initial solid concentration is \( c^s_{\text{eq}} = 0.95 \pm \epsilon \) where \( \epsilon \) is Gaussian white noise with variance of \( 1 \times 10^{-4} \).

Following a suggestion by Sleep [1988], this model also incorporates a porosity-dependent bulk viscosity for the solid matrix so that the dimensionless viscosity that controls compaction and fluid pressure is
\[ (\zeta + 4\eta/3) = \left( \frac{\phi_c}{\phi} \right)^m + \frac{4}{3} \left( \frac{\phi_c}{\phi} \right)^n + \frac{4}{3} \] (12)
which is nearly constant for porosities greater than the “compaction porosity” \( \phi_c \) but becomes strongly incompressible at small porosities (i.e. \( (\zeta + 4\eta/3) \rightarrow \infty \) as \( \phi \rightarrow 0 \)). Other initial and boundary conditions are described in Figure 1a. General notation and adjustable parameters in this model are given in Table 1, although in practice we fix most of these parameters except the Peclet and Damköhler numbers.

**Results**

**Basic behaviour**

Figure 2 shows the evolution of porosity with time for a run with \( Da = 40 \), \( Pe = 40 \), and shows the development of a series of high porosity channels that coalesce downstream (up section). By the end of this run (\( t = 116 \)), the calculation has developed significant channels in porosity such that near the top of the box, 94% of the vertical flux is going through only 24% of the available area. Channels are defined as areas where the melt flux is greater than the mean flux at a given height. The maximum porosity in the channels is \( \sim 3.9 \) times the original porosity. More significantly, strong localization arises because the porosity between the channels has been reduced to about 0.1 of the original porosity due to compaction. Because the melt flux scales as \( \phi^2 \), this implies that the maximum difference in flux between the channels and the inter-channel region is a factor of \( \sim 1500 \). The addition of compaction to reactive flow is a new feature in this problem and actually enhances the localization caused by this instability compared to rigid media [e.g. Ortoleva et al., 1987; Aharonov et al., 1997]. In particular, it leads to stronger localization and wavelength selection of channels with spacing smaller than the compaction length (see below).

We calculate the number of channels, the percentage of the total area occupied by channels and the percentage of the total vertical flux at that height that is carried in the channels. A useful measure of the degree of flow localization is the “excess channel flux” \( F_c \):

\[ F_c = \bar{F} \left( \frac{P_F}{P_A} - 1 \right) \] (13)

where \( \bar{F}(z,t) \) is the mean vertical flux at height \( z \), \( P_F \) is the percentage of flux in the channels, and \( P_A \) is the percentage of area covered by channels. For very weak oscillations of flux around the mean, \( P_F \sim P_A \sim 50\% \) and therefore the excess channel flux is close to zero. When a large percentage of flux is carried in the channels and they occupy a very small area \( F_c \) becomes large. Figures 3 and 4 show the evolution of the number of channels and excess channel flux for the numerical results illustrated in Figure 2.

Figure 3 shows the evolution in the number of channels and excess flux vertically averaged over the top region of the box from \( z = 4.5-5\delta \). At the top of the box the maximum number of channels (here \( \sim 3 \) per compaction length) is selected at approximately time \( t = 50 \), long before the excess channel flux significantly exceeds zero. Nevertheless, the excess flux is growing steadily and has two phases. In the earliest phase (\( t \lesssim 100 \)), the excess channel flux grows
Figure 2. Evolution of porosity with time for a run with $Da = 40, Pe = 40$. A dimensionless time of 1 in these runs is the time it takes to traverse the box once at the background porosity. At $t = 116$ the maximum porosity is 3.9% while the minimum porosity is only 0.1% due to compaction. In these runs the melt flux scales as $\phi^2$ so that the flux is $\sim 1500$ times greater in the channels than between them. (a) Absolute porosity with time, the contour is at $\phi = 1.01$. (b) Renormalized porosity with time. In each figure the porosity at each height $z$ is renormalized such that $\phi_n = (\phi - \phi_{min}(z))/(\phi_{max}(z) - \phi_{min}(z))$. (c) Cross section of porosity at $z = 4.5$ for the 4 panels in 2a.
exponentially with a doubling time $\tau$ of about 5.3. This rate is comparable to, but slightly slower than, the growth rate of the fastest growing mode predicted by linear analysis ($\tau = 4.4$). The discrepancy may arise from the resistance to compaction caused by the porosity dependent bulk viscosity (see Fig. 9). Channels grow faster in runs that can compact more easily. When the channels begin to become significant ($F_c \gtrsim 1$), the excess flux grows nearly linearly from $t \sim 100$, with a slope of $\frac{dt}{dF_c} = 11$.

Figure 4 shows the number of channels and excess channel flux for the entire height of the box. At early times, the number of channels ($\sim 30-40$) is determined by the initial noise. The number of channels at the top of the box rapidly decreases to a nearly constant value of about 10 to 12 and grows to a value of about 50 channels at the bottom, thus producing a coalescing network. The overall number of channels is already determined by $t = 56$ although the excess flux is still negligible at this time. Figure 4b shows the evolution of the excess channel flux as a function of height and time, and shows that it grows exponentially until $t \approx 100$ and then grows linearly. Overall, the vertical structure of the channels in this run is quasi-steady state. Once formed the channels tend to stay in the same place although the tips and junctures may migrate (a movie of this and other runs can be found at http://www.ldeo.columbia.edu/~mspieg/SolFlow/).

The change in number of channels with height can be predicted from the linear analysis (Eq. 14) by calculating the fastest-growing wavelength for a series of boxes of increasing height that have the same parameters per compaction length. This calculation is shown in Fig. 4a and agrees well with the non-linear solution. To demonstrate that this approximation is valid, Fig. 5 compares the porosity fields for two runs with the same parameters per compaction length but with reaction zones of length 40 and the 80. It is apparent that the number of channels predicted for a box of height 4 is a good approximation to the number of channels half way up a box of length 8.

The strong channeling is also apparent in other variables. Figure 6 shows the dissolution rate, solid concentration field
Figure 5. Normalized porosity at $t = 80$ for a short box with reactive zone of height $4\delta$ and one with a reactive zone twice as long. The short box is almost identical to the lower half of the long run.

and melt concentration field for $t = 116$. Inspection of the dissolution rate field shows that significant dissolution occurs only in the channels. The solid concentration field shows the integrated dissolution for the entire run and shows strongly corroded regions corresponding to the channels. By design, these runs do not exhaust the soluble phase before the run ends. Thus they only approximate the initial stages in the formation of dunite channels in the mantle, in which orthopyroxene is completely dissolved. Finally, Figure 6 shows the variation in melt concentration, and shows that it is slightly out of equilibrium in the channels. For lower $Da$ the degree of disequilibrium is larger while for larger $Da$ it is smaller. Nevertheless, the instability will occur even under conditions of complete equilibrium transport, $Da \to \infty$, and in fact grows fastest for large $Da$.

**Parameter Space**  Figure 7 shows the behaviour of this model in the parameter space spanned by $Pe$ and $Da$. This figure is effectively a phase diagram of 25 solutions in $\log Pe - \log Da$ space. The individual frames are shown at different times (see figure). However, times are the same for solutions with the same product of $DaPe$ (i.e. northwest-southeast trending diagonals). The choice of this time is driven by the linear analysis (next section) which suggests that the product of $DaPe$ is the principal control on the behaviour of these solutions. For any given value of $Da$, increasing the Peclet number increases the rate of channel growth and allows for more closely spaced channels because diminished diffusion allows more closely spaced chemical fluctuations to react before being diffused away. The spacing of the channels at the top of each box is nearly constant for a constant value of $DaPe$, however, higher values of the Damköhler number allow for more robust channels over a larger portion of the reactive zone. For $Da = 160$, well developed channels extend across the entire reactive zone. Preliminary results suggest that the size of the strongly channelized region depends on the ease of compaction at small porosities.

**Comparison to linear analysis**  Inspection of Figure 7 shows that there is a strong preferred wavelength of channeling in this problem and that the spacing and rate of growth of channels is principally a function of the product $DaPe$. A new linear analysis of this problem for large $Da$, based on the work of Aharonov et al. [1995], supports these claims and provides a useful framework for extending the numerical results to other problems. The principal result is a dispersion relationship for the exponential growth rate of porosity $\sigma$ assuming porosity perturbations of the form $\phi^\ast \propto \exp[\sigma t + ikx + mz]$ where $k = 2\pi/\lambda$ is the horizontal wavenumber for horizontal features of wavelength $\lambda$ and $m$.
Figure 7. Behaviour of the channeling calculation as a function of the two parameters the Dahmköhler number $Da$ and the Peclet number $Pe$. Higher $Da$ implies more rapid reaction. Higher $Pe$ implies less diffusion. NW-SE diagonals show solutions for which the product of $DaPe$ is constant and are shown at equivalent times. The top 5 runs are actually higher resolution runs ($259 \times 321$ grid points) that are periodic in 2 horizontal compaction lengths.
Aharonov et al. 1995 is slightly different from that given in the box in compaction lengths (here \( h = \delta \)). The dissolution term is always negative but goes to zero for high horizontal wavenumbers. The parameters in these numerical solutions were chosen to match the boundary conditions, both terms (and therefore \( \sigma \)) are real. The combination of these two terms is illustrated in Figure 8a which plots \( \sigma \) for the values of parameters used in the 25 runs shown in Fig. 7. The compaction term is always negative but goes to zero for high horizontal wavenumbers because wavelengths much smaller than the compaction length are difficult to compact. The dissolution term is always positive and has a maximum value of \((\rho_f/\rho_s) c_{eq}^f\) for large Da and \( k^2 < DaPe \). For horizontal wavenumbers \( k > \sqrt{DaPe} \), diffusion becomes important and growth of very high-wavenumber features is damped. The combination of compaction and diffusion gives rise to a strong peak in \( \sigma \) with significant wavelength selection. The preferred wavelength increases weakly with DaPe while the growth rate increases considerably. Figure 8b shows the same plot as 8a calculated from the numerical results of the 25 runs in Fig. 7. Figure 8b shows a similar structure to Figure 8a, and is in good agreement with the linear analysis for large values of DaPe (where the analysis is most appropriate) and large wavenumbers. Runs with DaPe \( \lesssim 800 \) have slower growth rates than predicted which appears to be related to the viscosity effects and limitations of the analysis at lower Da. The discussion, however, suggests that values of DaPe for the mantle are larger than the calculations shown here. Thus this analysis will be useful for estimating the behaviour and efficiency of channeling for appropriate mantle parameters.

For example, for very large values of DaPe expected in the mantle, the maximum growth rate from the linear analysis is \( \sigma_{max} = (\rho_f/\rho_s) c_{eq}^f \) or rescaling to box flushing times is

\[
\sigma'_{max} = \frac{\rho_f}{\rho_s} c_{eq}^{max}
\]

where \( c_{eq}^{max} \) is the maximum solubility at the top of the box. For the problems calculated here, \( c_{eq}^{max} = .1 \) and \( n = 2 \) so \( \sigma'_{max} \sim 0.2 \) which implies that at early times the most reactive problems will increase their porosity by a factor of \( e \) after 5 flushings of the box. In the mantle beneath ridges, \( n \) probably ranges between 2–4 and \( c_{eq}^{max} \sim 0.2–0.3 \) for pyroxenes (see Kelemen et al. 1995b) for estimates of \( c_{eq}^f \) and Aharonov et al. 1995 for estimates of effective parameter ranges for the mantle). This range of parameters could reduce the growth time to around 1–2 flushing times.

The long times in these calculations before flow in the channels becomes significantly non-linear stem from the very small initial perturbations that were used in our calculations in order to minimize the effects of the initial noise structure on the eventual wavelength selection. Figures 3&4, however, show that the instability is growing exponentially over the entire period. In a natural system with larger initial heterogeneity, it is expected that saturation will occur much faster but that channel structure might lock onto any large initial permeability variations. The linear analysis also suggests that for extremely reactive systems with significant initial heterogeneity, channels may exist at a range of scales smaller than the compaction length. The discussion explores expected values for the overall parameters and suggest that efficient channel systems could form from this mechanism in order 100,000 years with channel spacings \( \sim 1–100m \).

Discussion

The parameters in these numerical solutions were chosen to span a range that reveals most of the behaviour of this solution while remaining numerically accessible in terms of resolution. Nevertheless, the question remains as to the appropriate values of \( \delta \), Da and Pe for various Earth science problems of interest. The compaction length \( \delta \) occurs in both the Damköhler number and the Peclet number and can be readily estimated for a very permeable one-dimensional adiabatic melting column as

\[
\delta(z) \approx \left[ \frac{(\zeta + 4n/3)W_0F}{\Delta \rho g} \right]^{1/2}
\]

where \( W_0 \) is the solid upwelling velocity and \( F(z) \) is the degree of melting as a function of height. This is the “reduced compaction length” of Ribe 1985 [see also Spiegelman and Kenyon, 1992]. Values of parameters appropriate for melting beneath mid-ocean ridges1 yield a range of compaction lengths from \( \sim 10^2–10^4 \) m.

Given a range of values for the compaction length, estimating the Damköhler and Peclet numbers reduces to determining how reactive and diffusive the mantle is on this scale.

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1E.g. \( W_0 \sim 1–10 \text{ cm yr}^{-1} \), \((\zeta + 4n/3) = 10^{18–10} \text{ Pa s, } \Delta \rho g \sim 5000 \text{ Pa m}^{-1} \text{ and } F \sim 0.1–0.2 \text{ cm} / \text{h} \)
length scale. In general, the Damköhler number can be rewritten as $Da = \delta/L_{eq}$ where $L_{eq}$ is the “equilibration length” i.e. the distance that a parcel of fluid will travel before equilibrating with the matrix. A short equilibration length relative to the compaction length implies a high $Da$. Aharonov et al. [1995] (Appendix A), discuss estimates for $L_{eq}$ in some detail and give values from $10^{-7}$–100 m. The lower values assume reaction rate constants appropriate for the dissolution of pyroxenes in a well-mixed, undersaturated melt [e.g. Kuo and Kirkpatrick, 1985a; Brearley et al., 1986] and slow melt migration whereas the higher values are based on assumptions of more limited interaction between melts and pyroxene [Zhang et al., 1989] and faster melt migration. Either way, the equilibration length is still significantly smaller than the compaction length which implies that, in the mantle, if undersaturated melts have ready access to soluble minerals, then dissolution reactions can occur extremely rapidly.

A similar argument for the Peclet number suggests that diffusion is likely to be negligible and therefore $Pe$ should also be large. The Peclet number can be rewritten as $Pe = \delta/L_{diff}$ where $L_{diff}$ is the length scale over which diffusive/dispersive effects are important. If diffusion is controlled by molecular diffusion within the melt phase, $L_{diff}$ is likely to be very small because liquid diffusivities are of order $10^{-12}$ m$^2$s$^{-1}$. For melt velocities 1–1000 times solid upwelling velocities, the diffusion length due to molecular diffusion alone would still be $L_{diff} \ll 1$ m with Peclet numbers $> 10^6$. In a permeable flow system, however, it is likely that horizontal smearing of chemical fluctuations will be caused by hydrodynamic dispersion as well as chemical diffusion. For a dispersivity of $\alpha = 1$–10m the effective diffusivity would be $D_{eff} = \alpha w_0$ and the Peclet number would be $Pe \sim \delta/\alpha$. Even at quite high melt velocities, these dispersivities give estimates of $Pe \sim 10$–10000. The lower of these values would imply some diffusion. However an upper bound on the diffusion length can probably be placed by the observation that dunite channels and reaction zones around pyroxenite dikes in the mantle section of ophiolites preserve mineral compositions that are distinct from those in surrounding, residual peridotites [e.g., Quick, 1981; Kelemen et al., 1992, 1995a; Takahashi, 1992; Varfalvy et al., 1996]. Moreover, if channels can begin to form by compaction localization, the very low permeabilities between channels should limit the dispersivity and inter-channel diffusion.

All of these arguments taken together suggest that the values of $Da$ and $Pe$ appropriate for melting beneath mid-ocean ridges are significantly larger than the most reactive and least diffusive run shown in Fig. 7. If so, then the channeling instability is likely in the upper mantle and will grow at a rate close to the maximum growth rate (Eq. 15) for wavelengths smaller than the compaction length and larger than $\sim 2\pi\delta/\sqrt{DaPe}$. Using the linear analysis as a guide, we can estimate the growth rate and channel spacing for a range of mantle conditions. As an example, consider a melting column 75km high with a maximum solubility of

![Figure 8](image-url)
0.25, permeability exponent 2, compaction length $\delta = 1\text{km}$, $Da = Pe = 1000$ and melt velocity $w_0 \sim 10\text{m yr}^{-1}$. The e-folding time for the instability is $\sim 2$ flushing times (here $t_{\text{flush}} = 7500\text{yr}$) so we might expect significant channeling to form from a homogenous system, at this growth rate, in 100,000–150,000yr (see below). At this large value of $Da Pe$, the preferred channel spacing is $\sim 17$ channels/compaction length (60m) but all wavelengths between 20m to 160m all grow at rates greater than 95% of the maximum growth rate.

**Adding melting as well as reaction**

The previous analysis has been based on simple scaling arguments for 1-D steady state melting columns. However, the problem we have been considering thus far doesn’t actually include melting. In decompression melting regimes such as the region beneath mid-ocean ridges “background melting” adds porosity to regions that would otherwise compact to near impermeability in systems where additional melt is produced only by reaction (e.g. Figure 7). In addition, the rising solid will have only a finite residence time in the melting region.

In the calculations shown so far, we have assumed that the solid is stationary relative to the melt and can be fluxed indefinitely with a solvent. In an adiabatic melting system, the time-integrated flux of solvent through a piece of solid depends on the size of the system and the relative velocities of melt and solid. If we scale to the time it takes melt to cross the system, then for a low permeability system in which the velocities of melt and solid are the same, the solid will cross the system in a time of 1. In this case, the solid will only see the melt that was produced locally, and channeling will not occur. For a higher permeability system, however, the solid will move more slowly than the melt. In this case, a significant flux of melt will pass through the rock, and channeling will occur. Thus, the numerical results presented in previous sections can be viewed as solutions to the limiting case of an infinitely permeable system. The question remains what happens when melting is included and the permeability is finite.

Appendix A provides the additional equations required for a background melting rate proportional to the solid upwelling rate $W_0$. Figure 9b shows results for porosity for a melting run with $Da = 80$ and $Pe = 160$ and a mean melt velocity about 200 times the solid upwelling velocity. Strong, compaction driven channels are produced with spacings comparable to the static run with the same $Da$ and $Pe$ (Fig. 7). Unlike the static runs, however, this problem evolves to a quasi-steady state where the maximum channel porosity at any height is roughly constant with time. For Fig. 9b, the maximum porosity is $\sim 5\phi_0$ and the minimum porosity is $\sim 0.3\phi_0$. Significant channels are seen at about 15 flushing times and the steady state is achieved by 25 flushing times. For this problem, the channels only occupy the upper two compaction lengths of the box.

However, additional runs (Fig. 9a,c) suggest that both the growth rate and the size of the region with significant channels in the melting runs depends on how compactible the medium is. For the melting runs in Figure 9, the matrix becomes harder to compact for porosities much less than the compaction porosity $\phi_c$ (see Eq. 12) Figure 9a shows the same problem as Fig. 9b but is easier to compact and produces significant channels over a larger region in less time than Fig. 9b. Figure 9c is more difficult to compact than 9b and channels grow slowly and are only significant in the upper compaction length. A run with $\phi_c = 0.5\phi_0$ (not shown) has negligible channels. More work needs to be done to quantify this phenomenon. However, it emphasizes an important feature of this paper, that compaction strongly enhances flow localization by producing low permeability, inter-channel regions.

**Observational constraints on channeling**

Give the behaviour of the simplest reactive channeling problems, here we review and amplify the important observational constraints that can be compared and contrasted with theory.

**Chemical Disequilibrium** We emphasize the central observation that primitive mid-ocean ridge basalts are observed to be in neither major or trace element equilibrium with shallow, residual mantle peridotites [e.g. O’Harra, 1965; Stolper, 1980; Johnson et al., 1990; Johnson and Dick, 1992]. Both orthopyroxene undersaturation and trace element disequilibrium imply that primitive mid-ocean ridge basalts must have last equilibrated with mantle peridotite at a pressure of 1.0 GPa or more. Such liquids must pass through the top 30 km, or more, of the oceanic upper mantle without equilibrating with residual mantle peridotite. This constraint can be satisfied by very rapid transport, for example in melt-filled fractures — so that the diffusion time is short — or by slower porous flow in wide dunite conduits, so that the distance between migrating melt and residual peridotite is longer than the diffusion distance.

**Melt migration features in mantle samples** The characteristics of melt migration features in the mantle section of ophiolites, and in drill core from the oceanic mantle, were reviewed by Kelemen et al. [1997]. They may be classified into two groups: dikes and dunites. Dikes are composed of igneous pyroxenes and plagioclase, as well as minor olivine, which crystallized from melt migrating in an open fracture. Their compositions require conductive cooling of migrating melt, and they are typically undeformed, indicating that they
Figure 9. Comparison of porosity for three melting runs with Da = 80, Pe = 160 and three different values of the compaction porosity \( \phi_c \). The degree of melting \( F \) increases linearly from 0 at the base of the reactive zone \( (z = 1\delta) \) to 20\% at the top \( (z = 5\delta) \), while the solubility of the soluble phase increases linearly from 0.2 to 0.4 across the reaction/melting zone (thus \( c_f^0 = 0.05 \) is twice as large as in the previous figures). The scale porosity is \( \phi_0 = 0.001 \) and thus the scale melt velocity is roughly 200 times faster than the solid velocity. (a) Easily compactible run \( \phi_c = 0.1\phi_0 \) develops robust channel system over more than half of the column in approximately 10 flushing times. The run stopped at \( t = 17 \) due to extreme permeability variations (\( \phi_{\text{min}} = 0.09\phi_0 \)). (b) Moderately compactible run goes to quasi-steady state channel system by \( t = 25 \) (\( \phi_{\text{min}} = 0.3\phi_0 \)). (c) Poorly compactible system produces weaker steady channel network after \( t \sim 30 \) (\( \phi_{\text{min}} = 0.45\phi_0 \)).

have not undergone corner flow beneath a spreading ridge. Thus, it is inferred that they formed off-axis in a conductive boundary layer. Dunites commonly show evidence for extensive ductile deformation along with their host peridotites, and could form from migrating melt in the adiabatically upwelling region beneath a ridge. Many dunites also show contact relationships that indicate that they form by replacement, as a result of dissolution of pyroxene from mantle peridotite in melt migrating by porous flow. Rarely if ever are mantle dunites formed by crystallization of olivine in a melt-filled, open fracture.

Thus, observed melt migration features in ophiolite mantle and drill core from oceanic mantle indicate that dunites, formed as a result of reactive flow of melt, are important conduits for melt extraction. There is no field evidence which requires melt migration in fractures in the adiabatically upwelling region of the mantle, except in the shallowest mantle. Furthermore, Kelemen et al. [1997] showed that focused flow of melt in porous conduits could satisfy melt velocity constraints derived from Uranium-series disequilibria in lavas [e.g. data compilations of Sims et al., 1995; Lundstrom et al., 1995] and from the timing of volcanism due to decompression melting following deglaciation in Iceland (Jull and McKenzie [1996]; Slater et al. [1998]).

These observations are consistent with an important role for the Reactive Infiltration Instability (RII) in mantle melt extraction. However, it does not follow that the geological evidence requires extensive channels formed by the RII. Migration of melt in fractures within the melting region cannot be ruled out on the basis of geological or geochemical evidence. Adiabatic ascent of melt within a fracture in the melting regime would produce no crystallization, so no dike would form. Reaction zones along the walls of such fractures would be composed of replacive dunite.

Thus, it is possible that some or all dunites form as porous reaction zones around melt-filled fractures, as proposed by Quick [1981], Nicolas [1989] and Suhr [1999]. Perhaps such a mechanism, near the transition from the adiabatic region to the conductive thermal boundary layer, forms small dunites, less than one meter wide, which contain medial pyroxenite or gabbro dikes, such as are locally observed in the Trinity, Josephine and Oman ophiolite mantle sections, and in drill core from the shallow mantle from the East Pacific Rise [e.g. Quick, 1981; Boudier and Coleman, 1981; Kelemen and Dick, 1995; Dick and Natland, 1996]. It is less likely that large dunites, up to 100 meters wide in the Oman mantle section, form as reaction zones around a single crack.

Clearly it is possible to envision melt migration mechanisms intermediate between open, melt-filled fractures and porous conduits and combinations of mechanical and chemical instabilities that can form dunites. The RII with compaction leads to significant flow localization by itself, which is probably enhanced by mechanical instabilities and both types of instabilities are likely to occur together in the mantle. Significant work remains, however, to understand these more complex instabilities and their diagnostic observable features.

Focusing of mantle melt extraction to the ridge axis

Despite the fact that fracture-controlled melt migration mechanisms in the melting region cannot be ruled out on the basis of outcrop-scale geologic evidence, plate scale observational constraints can be used to argue that relatively simple porous flow within a viscously compacting solid matrix is probably the dominant mechanism of melt transport in the melting region beneath oceanic spreading ridges. The
combination of two crucial observations, (1) that the igneous ocean crust at fast-spreading ridges reaches 95% of its total thickness within 2 km of the ridge axis [e.g. Vera et al., 1990], and (2) that modeling and seismic data suggest that the region of partial melting is on the order of 100 km wide at its base [e.g. Barnouin-Jha et al., 1997; Forsyth et al., 1998], requires that melt transport occurs via mechanisms which focus melt flow toward the ridge axis. As noted by Sleep [1988], melt extraction in fractures spanning the melting region in mantle undergoing passive, corner flow beneath a spreading ridge cannot satisfy this constraint, because such fractures, formed parallel to the direction of maximum compressive stress, would reach the top of the mantle over a region more than 80 km wide.

If solid mantle upwelling is focused into a narrow column directly beneath the ridge, then perhaps melt flow in fractures could produce focused accretion of igneous crust within 2 km of the ridge axis. However, seismic results from the MELT experiment do not require focused solid upwelling [Forsyth et al., 1998], and may rule out focused upwelling geometries. Also, some extremely focused solid upwelling geometries, such as those required to focus all melt extraction to a narrow region beneath ridges, produce melts with compositions unlike MORB [Spiegelman, 1996]. Thus, it seems that melt migration primarily controlled by fractures cannot produce the observed coalescence of melt extraction toward spreading ridges.

In contrast, a variety of porous flow mechanisms have been proposed to explain coalescence of melt toward ridges: A. “suction” due to corner flow [Spiegelman and McKenzie, 1987; Phipps Morgan, 1987]; B. anisotropic permeability along mineral foliation [Phipps Morgan, 1987] or in stress controlled planes of high porosity [Daines and Kohlstedt, 1997; Zimmerman and Kohlstedt, 1999]; C. channels at the base of the “lithosphere”, beneath a permeability barrier created by melt crystallization [Sparks and Parmentier, 1991; Spiegelman, 1993c]; and D. coalescence of dissolution channels, as illustrated in Figures 2–7 of this paper.

Individually, each of these may be insufficient to explain melt focusing to the ridge. “A” requires mantle viscosities higher than estimated, particularly in high temperature mantle such as beneath Iceland [review in Hirth and Kohlstedt, 1996]; “B” requires foliated peridotites and/or non-hydrostatic stress, which may not be present in much of the melting region; “C” should form “impregnated peridotite” — residual mantle combined with interstitial products of melt crystallization — above dunite in ophiolites, but this is not observed; and ongoing modeling of “D,” presented in this paper, suggests that the aspect ratio of coalescence is insufficient to focus melt from, e.g., a 100 km tall melting region, 100 km wide at the base, into a 5 km wide zone beneath the ridge.

However, combinations of A-D may be sufficient to explain focused melt extraction beneath ridges. Unstable formation of dissolution channels by decompressing melt is enhanced by melt flux, and intrinsically follows pressure gradients in the liquid. High melt flux and lateral pressure drops toward the ridge axis due to A and B will lead to the formation of the largest channels beneath the ridge, and of diagonal dissolution channels converging toward the ridge. Also, a variant of C is likely, particularly beneath slow spreading ridges. Diagonal isotherms near the top of the melting region may drive formation of diagonal dissolution channels converging toward the ridge.

Thus, coalescence of melt transport from a broad melting region to a narrow region of igneous crustal accretion beneath oceanic spreading ridges seems to require that the dominant mode of melt transport is via a combination of porous melt migration mechanisms. If melt migration is dominantly by porous flow then this, in turn, requires that melt migration through the upper 30 km of the mantle beneath ridges occurs in wide, porous dunites conduits, which shield migrating melt from equilibrating with residual mantle peridotite. Thus, the RII may well be an important factor in controlling melt migration in the upper mantle beneath ridges.

Comparison of model results and observations

Because the numerical model considered in this paper is greatly simplified compared to the actual process of melt extraction from the mantle beneath oceanic spreading ridges, it may be premature to directly compare modeling results to field observations. However, here we try to look toward the future, to anticipate how the application of the theory of RII to mantle melt extraction may be tested and refined.

There are two intrinsically different types of models presented in this paper, those with solvent migrating through a static solid matrix (Figures 1 through 8), and those which incorporate an upwelling solid matrix and decompression melting (Figure 9). Each is potentially comparable to different mantle samples. Results of modeling with a static solid matrix may have relevance for environments on Earth in which melt rises into nearly static, overlying mantle peridotite; for example, migration of melt in the shallow mantle beneath subduction-related magmatic arcs might be approximated in this way provided that solubility of pyroxene increases upward in such settings. In many ophiolites, such as the Trinity, Josephine and Ingalls ophiolites near the US West Coast, late- and post-kinematic replacive dunites preserve evidence for melt migration through nearly static mantle [Quick, 1981; Kelemen et al., 1992; Kelemen and Dick, 1995, and our unpublished data]. For mid-ocean ridges, and
for the early features of the Oman ophiolite, formed beneath an active oceanic spreading ridge, the results of modeling including decompression of the solid matrix are more relevant.

Of course, many aspects of the simple models presented in this paper do not adequately simulate dunite formation in the mantle. As noted in a previous section, none of the models shown here reach exhaustion of the soluble phase (but see www.ldeo.columbia.edu/~mspieg/SolFlow), whereas it is common to find mantle dunites which are completely free of pyroxene. Furthermore, division of the solid matrix in the models into “soluble” and “insoluble” components is not strictly correct, since even the refractory minerals olivine and spinel, which compose dunites, are slightly soluble in adiabatically ascending mantle melts. We have also neglected more complex multi-component reactions as considered by Steefel and Lasaga [1994]. These will be investigated in future studies.

Another crucial divergence between dissolution channels in simple models and observed mantle dunites is their three-dimensional morphology. Although the numerical results presented in this paper are two-dimensional, Aharonov et al. [1997] presented three-dimensional results of numerical models of the RII in rigid, soluble porous media. In those results, dissolution channels were roughly cylindrical, whereas most dunites in the mantle section of ophiolites are tabular features. Nevertheless, these 3-D models did not include compaction which clearly enhances the instability in 2-D. As it is easier to compact tabular features than cylindrical ones, it may be that compaction in 3-D leads to tabular features; however, this remains to be tested with further modeling. Alternatively, it may be that natural, tabular dunites result from pre-existing foliation in mantle peridotites, e.g. anisotropic permeability due to crystal shape and/or lattice preferred orientation, or the presence of tabular pyroxene-rich banding [Dick and Sinton, 1979].

Comparison of quantitative model results to field observations is rendered difficult by the fact that the instantaneous and time-integrated values of most calculated parameters, e.g., melt flux, continue to vary throughout the model runs, whereas mantle outcrops preserve only the final, time-integrated effects of melt migration. An exception to this is the quasi-steady state structure of the coalescing channel networks illustrated in Figures 3, 4 and 9a. Because this structure remains nearly constant in time and space in the model results, it has the potential for direct comparison to similar structures formed in the mantle. Currently, this is not straightforward, because the models do not reach phase exhaustion, so the final width of “dunite” channels has not been determined in the numerical results. Conversely, the width of dunite channels in the field is easily measured, but the time-integrated flux of melt through the dunites is much less well-constrained.

We can make a start at comparing numerical results and field observations by assuming that the final width of dunite channels is related to the flux of melt through the channels. In Figure 10, we show flux/frequency and size/frequency statistics for channel networks from models and from mantle outcrops in the Ingalls ophiolite [Kelemen et al., 2000]. Figure 10a illustrates the power law relationship for flux/channel vs. number of channels with a given flux defined by an idealized, coalescing network in which flux is preserved downstream (“Einat’s Castle”, [Fig A1 Aharonov et al., 1995]). Figure 10b shows the same relationship in results of the numerical model illustrated in Figures 2 through 6 of this paper. The slopes in Figures 10a and 10b are the same because nearly all the melt in the numerical model migrates in channels, and flux is conserved downstream through the coalescing network. Figure 10c is a plot of channel width (intercept length) vs. number of channels with a given width (per meter of intercept measurements) in three outcrops from the Ingalls ophiolite in the Washington Cascades, USA which also shows a power law relationship albeit with a different slope. To make the current data on size/frequency relationships for dunites consistent with the flux/frequency relationships in our numerical modeling results, requires that the flux in the dunites is proportional to the dunite width. However, it is possible that other melt migration mechanisms could give rise to a power law relationship between dunite size and frequency, so that the qualitative similarity between observations and our model results cannot be taken as a confirmation that the RII is responsible for forming dunites in the Ingalls ophiolite.

More complete tests of the applicability of the RII to melt extraction from the mantle beneath oceanic spreading ridges await completion of more elaborate numerical models, and further characterization of the spatial relationships and integrated melt flux for dunites in mantle outcrops. Also, it will be important to develop quantitative, predictive models for other melt migration mechanisms, such as hydrofracture, to search for spatial relationships which might be diagnostic of a particular transport mechanism. Nevertheless, these initial results suggest that strong flow localization can form from coupled fluid-solid process associated with melt transport.

Appendix A: Derivation and scaling of governing equations

The equations for flow in viscously deformable, permeable media have been discussed in detail by several authors [see, McKenzie, 1984; Scott and Stevenson, 1984, 1986; Spiegelman, 1993a, b, c]. The equations for conservation of total mass and momentum of the two-phase system of melt
and solid are

\[
\frac{\partial \rho_f \phi}{\partial t} + \nabla \cdot [\rho_f \phi \mathbf{v}] = \Gamma \tag{A1}
\]

\[
\frac{\partial \rho_s (1 - \phi)}{\partial t} + \nabla \cdot [\rho_s (1 - \phi) \mathbf{V}] = -\Gamma \tag{A2}
\]

\[
\phi (\mathbf{v} - \mathbf{V}) = -\frac{k_\phi}{\mu} \left[ \nabla P - \rho_f \mathbf{g} \right] \tag{A3}
\]

\[
\nabla P = -\nabla \times \eta (\nabla \times \mathbf{V}) + \nabla (\zeta + 4\eta/3) \cdot \mathbf{V} + G + \dot{\rho} \mathbf{g} \tag{A4}
\]

\[
k_\phi = \frac{a^2 \phi^n}{b} \tag{A5}
\]

Where \(\rho_f, \rho_s\) are the melt and solid densities, \(\phi\) is the volume fraction of melt (porosity), \(\mathbf{v}\) and \(\mathbf{V}\) are the melt and solid velocities and \(\Gamma\) is the total rate of mass transfer from solid to liquid. \(k_\phi\) is the permeability which is a non-linear function of porosity (Eq. A5), \(\mu\) is the melt viscosity, \(P\) is the fluid pressure and \(\mathbf{g}\) is the acceleration due to gravity. Finally, \(\eta\) is the solid shear viscosity, \((\zeta + 4\eta/3)\) is the combination of solid bulk and shear viscosity that controls volumes changes of the matrix, \(G/n, \mathbf{V}\) are the cross-terms that arise for non-constant shear viscosity (and vanish if \(\eta\) is constant) and \(\dot{\rho} = \rho_f \phi + \rho_s (1 - \phi)\) is the mean density of the two phase system. Equations (A1) and (A2) conserve mass for the melt and solid respectively and allow mass-transfer between the phases. Equation (A3) governs the separation between melt and solid and Eq. (A4) governs stress-balance and deformation of the solid phase.

To close these equations requires functional forms for the mass-transfer rate \(\Gamma\). For simple adiabatic upwelling, the mass transfer rate can be set to be a function of the upwelling rate [for example, see Spiegelman, 1993c, 1996]. Reactive flows, however, require a more general formulation that, at the minimum, requires tracking the compositions of melt and solid. For an \(N\) component system with \(J\) simultaneous reactions, we can write \(2N\) equations for the conservation of mass of component \(i\) in the melt and solid as

\[
\frac{\partial \rho_f \phi c_i^f}{\partial t} + \nabla \cdot \left[ \rho_f \phi D_i^f \nabla c_i^f + \sum_{j=1}^{J} c_{ij}^s \Gamma_j \right] \tag{A6}
\]

\[
\frac{\partial \rho_s (1 - \phi) c_i^s}{\partial t} + \nabla \cdot \left[ \rho_s (1 - \phi) c_i^s \mathbf{V} \right] = -\sum_{j=1}^{J} c_{ij}^s \Gamma_j \tag{A7}
\]

where \(c_i^f, c_i^s\) are the concentration of component \(i\) in the melt and solid respectively. \(D_i^f\) is the combined diffusivity/dispersivity tensor of component \(i\) in the melt (we assume negligible solid diffusion) and \(c_{ij}^s\) is the concentration of component \(i\) in the fluid that is involved in reaction \(j\). \(\Gamma_j\) is the rate of mass transfer for reaction \(j\). Specific forms for different types of reactions will be introduced below. To make Eqs. (A6)–(A7) consistent with Eqs. (A1)–(A2) and the property that the sum of all concentrations in any phase must add to 100% (e.g. \(\sum_{i=1}^{N} c_i^f = \sum_{i=1}^{N} c_i^s = \sum_{i=1}^{N} c_{ij}^s = 1\)) requires that the total mass transfer rate be

\[
\Gamma = \sum_{j=1}^{J} \Gamma_j \tag{A8}
\]

and that \(\sum_{i=1}^{N} \nabla \cdot \rho_f \phi D_i^f \nabla c_i^f = 0\) because only \(N - 1\) concentrations can freely diffuse. The final component must be anti-diffusive to conserve mass. For extensions to more complicated multi-component systems see Steefel and Lasaga [1994].
To make Eqs. (A1)–(A7) more tractable and amenable to analysis, it is useful to rewrite the equations into potential form and make them dimensionless. Spiegelman [1993a] provides a basic recipe for scaling and writing Eqs. (A1)–(A5) in potential form. To get from Eqs. (A1)–(A7) to (1)–(4) we follow the analysis in Spiegelman [1993a] and Aharonov et al. [1995]. To get the dimensional forms of the equations, expand (A2) and define $C = \nabla \cdot \mathbf{V}$ to yield:

$$\frac{\partial \phi}{\partial t} + \nabla \cdot \phi = (1 - \phi)C + \frac{\Gamma}{\rho_s} \cdot \nabla$$

(A9)

Adding (A1) and (A2) and substituting in (A3) and (A4) under the assumptions $\eta = \text{constant}$ and neglecting rotational flow yields

$$-\nabla \cdot \frac{k_0}{\mu} \nabla (\zeta + 4\eta/3)C + C = -\nabla \left[ \frac{k_0 \Delta \rho g}{\mu} (1 - \phi) \right] + \frac{\Delta \rho}{\rho_f \rho_s} \cdot \nabla$$

or substituting $P = (\zeta + 4\eta/3)C$ yields the equation in terms of excess pressure

$$-\nabla \cdot \frac{k_0}{\mu} \nabla P + \left[ \frac{P}{(\zeta + 4\eta/3)} \right] = -\nabla \left[ \frac{k_0 \Delta \rho g}{\mu} (1 - \phi) \right] + \frac{\Delta \rho}{\rho_f \rho_s} \cdot \nabla$$

(A10)

By the definitions for $C$ and $\mathbf{V} = \nabla U$ we get $\nabla^2 U = C$.

To get the dimensional equations for conservation of composition, we expand the RHS of Eqs. (A6) and (A7) and substitute in Eqs. (A1), (A2) and (A8) to get

$$\rho_s(1 - \phi) \frac{D_A c^*}{D_t} = - \left[ \sum_{j=1}^{J} (c_{ij}^* - c_i^*) \Gamma_j \right]$$

(A12)

$$\rho_f \phi \frac{D_f c_i^f}{D_t} = \left( \nabla \cdot \rho_f \phi \Delta_i \nabla c_i^f + \left[ \sum_{j=1}^{J} (c_{ij}^* - c_i^*) \Gamma_j \right] \right)$$

(A13)

To reduce the number of adjustable parameters, it is useful to non-dimensionalize Equations (A9)–(A13) with the following natural scaling:

$$\phi = \phi_0 \phi' \quad t = \frac{\delta}{w_0} t'$$

$$x = \delta x' \quad \nabla = \frac{1}{\delta} \nabla'$$

$$\mathbf{u} = \mathbf{u}(\mathbf{v}, \mathbf{V})' \quad U = \delta w_0 U'$$

$$C = \frac{\phi_0 w_0}{\delta} C' \quad \Phi = \Delta \rho g \Phi'$$

$$\left( \Gamma, \Gamma_j \right)' = \frac{\rho_s \phi_0 w_0}{\delta} \left( \Gamma, \Gamma_j \right)'$$

$$\left( \zeta + 4\eta/3 \right) = \left( \zeta + 4\eta/3 \right)_0 \left( \zeta + 4\eta/3 \right)'$$

where

$$\delta = \sqrt{\frac{k_0 (\zeta + 4\eta/3)}{\mu}}$$

(A14)

$$\phi_0 w_0 = \frac{k_0 \Delta \rho g}{\mu}$$

(A15)

are the compaction length and the melt separation velocity [e.g. McKenzie, 1984; Spiegelman, 1993a] where $k_0$ and $(\zeta + 4\eta/3)_0$ are the permeability and solid viscosity at porosity $\phi_0$. Substituting these definitions into (A9)–(A13) and dropping primes yields

$$\frac{D_A \phi}{D_t} = (1 - \phi_0 \phi)C + \Gamma$$

(A16)

$$-\nabla \cdot k_0 \nabla P + \frac{P}{(\zeta + 4\eta/3)} = -\nabla \cdot \left[ k_0 (1 - \phi_0 \phi) \right] + \frac{\Delta \rho}{\rho_f} \cdot \nabla$$

(A17)

$$\nabla^2 U = \phi_0 C$$

(A18)

$$\frac{D_f c^f}{D_t} = \frac{-\phi_0}{(1 - \phi_0 \phi)} \left[ \sum_{j=1}^{J} (c_{ij}^* - c_i^*) \Gamma_j \right]$$

(A19)

$$\frac{D_A c^*}{D_t} = \frac{1}{\phi} \left[ \frac{1}{\rho_f \phi} \nabla \cdot \phi \nabla c^f + \frac{\rho_s}{\rho_f} \left[ \sum_{j=1}^{J} (c_{ij}^* - c_i^*) \Gamma_j \right] \right]$$

(A20)

where

$$\rho_A = (\zeta + 4\eta/3)C$$

(A21)

$$k_0 = \phi'$$

(A22)

$$\mathbf{V} = \nabla U$$

(A23)

$$\mathbf{v} = \mathbf{V} - \frac{k_0}{\phi} [\nabla P - (1 - \phi_0 \phi) \mathbf{k}]$$

(A24)

The principal simplifications required to get Eqs. (1)–(4) are that the reference porosity $\phi_0$ is sufficiently small such that Eq. (A18) can be neglected (i.e. $\mathbf{V} \approx 0$), that

$$\frac{1}{\rho_f \phi} \nabla \cdot \rho_f \phi \nabla \sim \nabla^2$$

(A25)

and that there is only a single reaction $p_{\text{solid}} \rightarrow p_{\text{dissolved}}$ and $c_{ij}^* = c_i^f = 1$. The rate of this reaction is $\Gamma_R$ which we assume to be controlled by linear kinetics such that

$$\Gamma_R = RA(\phi, c') \left[ c_{i}^* - c_i^f \right]$$

(A26)

where $R$ is a reaction rate per unit area, $A$ is the specific area of the reaction and depends on the porosity and amount of soluble phase. Equation (A26) states that the reactive dissolution rate is proportional to the difference between the melt composition and the equilibrium composition. If the melt is undersaturated in the reactive component ($c_i^f < c_{i}^*$) the
solid will dissolve ($\Gamma_R > 0$) else the solid will precipitate. Using the scaling given in Eq. (A14), Eq. (A26) can be written in dimensionless form as

$$\Gamma_R = D_{\text{ac}} A' (c_{eq}^f - c_f^f)$$ (A27)

In the more general case of adiabatic background melting and dissolution

$$\Gamma = \Gamma_0 + \Gamma_R$$ (A28)

where dimensionally

$$\Gamma_0 = \rho_s W \frac{\partial F}{\partial z}$$ (A29)

is the mass-transfer rate due to adiabatic decompression and is simply proportional to the solid upwelling velocity and the imposed degree of melting $F$ as a function of height. Figure 9 has details of specific melting runs.

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**References**


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