Frontiers

The chemistry of subduction-zone fluids

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Received 24 April 2004; accepted 27 April 2004

Abstract

Subduction zones generate voluminous magma and mediate global element cycling. Fluids are essential to this activity, yet their behavior is perhaps the most poorly understood aspect of the subduction process. Though many volatile components are subducted, H₂O is the most abundant, is preferentially fractionated into the fluid phase, and, among terrestrial volatiles, is by far the most effective solvent. H₂O therefore controls the chemical properties of subduction-zone fluids. Rising pressure (P) and temperature (T) along subduction paths yield increased H₂O ionization, which enhances dissolved solute concentrations. Under appropriate conditions, silicate solubilities may become so high that there is complete miscibility between hydrous melts and dilute aqueous solutions. Miscible fluids of intermediate composition (e.g., 50% silicate, 50% H₂O) are commonly invoked as material-transport agents in subduction zones; however, phase relations pose problems for their existence over significant length scales in the mantle. Nevertheless, this behavior provides a key clue pointing to the importance of polymerization of alkali aluminosilicate components in deep fluids. Aqueous aluminosilicate polymers may enhance solubility of important elements even in H₂O-rich fluids. Subduction-zone fluids may be surprisingly dilute, having only two to three times the total dissolved solids (TDS) of seawater. Silica and alkalis are the dominant solutes, with significant Al and Ca and low Mg and Fe, consistent with a role for aqueous aluminosilicate polymers. Trace-element patterns of fluids carrying only dissolved silicate components are similar to those of primitive island-arc basalts, implying that reactive flow of H₂O-rich, Cl-poor, alkali-aluminosilicate-bearing fluid is fundamental to element transport in the mantle wedge. Better understanding of the interaction of this fluid with the mantle wedge requires quantitative reaction-flow modeling, but further studies are required to achieve this goal.

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Keywords: subduction zones; subduction-zone fluids; mantle wedge; metasomatism

1. Introduction

The segments of subduction zones extending from trenches to beneath volcanic arcs are sites of profound chemical change. Incoming lithosphere is stripped of elements [1–4] which are transferred to the overlying mantle [5–9] in a process that ultimately generates volcanic arcs. The chemical work done in this “subduction factory” is fundamental to the Earth’s evolution, because it leads to prolific volcanism and degassing, it mediates the global cycling of elements, and over time it produced the continental crust.

The transfer of material in subduction zones occurs in steps, and the agents of transfer vary. It is generally thought that H₂O-rich fluid (Table 1) is initially...
responsible for leaching elements from the slab [8,10–13]. The fluid is liberated by metamorphic devolatilization in subducting lithosphere and carries solutes as it migrates into the overlying wedge of mantle, resulting in chemical modification, or “metasomatism,” of slab and wedge. As the process continues, the fluid triggers melting, yielding voluminous magma that is chiefly basaltic in composition. These magmas serve as the second agents of mass transfer, bringing slab- and mantle-derived components toward the surface as they rise.

Although melting in subduction-zone settings can be quite complex [8,9,12,14], study of the sources and evolution of the magmas is well advanced. In contrast, we know little about the fluid that begins the process of material transfer in subduction zones. In general, understanding the compositional evolution of a moving fluid and its host rocks requires techniques that couple fluid flow with chemical reaction. Reactive flow characterizes many terrestrial environments—e.g., ore deposits, crustal metamorphic systems, sedimentary basins, mid-ocean ridges—and modeling element transport in such systems has met with success. The same cannot be said of subduction zones, primarily because we lack basic information on fluid composition and how it is controlled. No direct, pristine fluid sample can be collected from this environment and working backwards from evolved magmatic products yields insight into only a part of the flow system. In addition, experimental study of fluids at the requisite high pressure ($P$) and temperature ($T$) has proven to be a singular challenge. As a result, fundamental questions remain: are the fluids dilute solutions or silicate-rich mixtures intermediate between $\text{H}_2\text{O}$ and melt? What is the role of ligands such as Cl? How does mineral solubility and element partitioning change along the flow path? Answering these questions requires a better understanding of the chemical behavior of the fluid phase at great depth.

In this paper, I highlight recent advances that offer preliminary insights into the chemical behavior of subduction-zone fluids. I first review the physical controls that operate along the paths of fluid flow. This is followed by discussion of the chemical properties of solutions at high $P$ and temperature ($T$) has proven to be a singular challenge. As a result, fundamental questions remain: are the fluids dilute solutions or silicate-rich mixtures intermediate between $\text{H}_2\text{O}$ and melt? What is the role of ligands such as Cl? How does mineral solubility and element partitioning change along the flow path? Answering these questions requires a better understanding of the chemical behavior of the fluid phase at great depth.

In this paper, I highlight recent advances that offer preliminary insights into the chemical behavior of subduction-zone fluids. I first review the physical controls that operate along the paths of fluid flow. This is followed by discussion of the chemical properties of solutions at high $P$, which gives context to a summary of constraints on composition. The new results are an initial step toward a chemical foundation for investigating one of Earth’s most important fluids.

### 2. Physical controls on subduction-zone fluids

As summarized in Fig. 1, rising $P$ and $T$ during subduction drive mineral reactions that yield a discrete fluid phase. The fluid is probably rich in $\text{H}_2\text{O}$ relative to other volatiles (e.g., $\text{CO}_2$) because of greater abundance, favorable partitioning and low thermal

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Glossary of terms</th>
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<tbody>
<tr>
<td>Compatible element</td>
<td>An element that is preferentially partitioned into the solid in a solid–fluid mixture</td>
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<tr>
<td>Critical end point</td>
<td>Point defined by the intersection of a critical curve and a solubility curve</td>
</tr>
<tr>
<td>Critical curve</td>
<td>Curve linking critical points of end members in a two-component system. It is the boundary between stability fields of a supercritical fluid and a two phase mixture of liquid + vapor</td>
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<tr>
<td>Critical point</td>
<td>The termination of the liquid + vapor field in a one component system</td>
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<tr>
<td>Fluid</td>
<td>A disordered, non-crystalline phase consisting of particles in motion and possessing unspecified composition</td>
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<tr>
<td>Incompatible element</td>
<td>An element that is preferentially partitioned into the fluid in a solid–fluid mixture</td>
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<td>Liquid</td>
<td>High-density, subcritical fluid</td>
</tr>
<tr>
<td>Metasomatism</td>
<td>The process by which rock is compositionally modified</td>
</tr>
<tr>
<td>Miscibility</td>
<td>The property enabling discrete phases to mix completely to form a single phase</td>
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<tr>
<td>Polymerization</td>
<td>In the context of this paper, the linking of cations by shared (bridging) oxygens</td>
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<tr>
<td>Solubility curve</td>
<td>In a simple two component system, a curve linking the eutectic with the triple point of an end member. In the system $A-\text{H}_2\text{O}$, the high-$T$ solubility curve is the line along which solid $A$ may coexist with liquid and vapor</td>
</tr>
<tr>
<td>Solidus</td>
<td>Reaction boundary denoting the first appearance of melt with increasing $T$ in a multicomponent system</td>
</tr>
<tr>
<td>Supercritical</td>
<td>State of a system in which a single fluid phase is stable</td>
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<tr>
<td>Supercritical fluid</td>
<td>A fluid stable at $P$ and $T$ greater than the critical point or curve in the system of interest</td>
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<tr>
<td>Total dissolved solids</td>
<td>The sum of masses of all dissolved solutes, in g/kg $\text{H}_2\text{O}$</td>
</tr>
<tr>
<td>Vapor</td>
<td>Low-density, subcritical fluid. Synonymous with “gas”</td>
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stability of hydrated silicates [15–20]. Experiments, simulations, and isotope geochemistry indicate that volatile liberation from subducted lithologies is more or less continuous [13,21–28], although instances of episodic fluid production may be indicated by slab seismicity [18,29].

The slab’s capacity to produce fluid diminishes with depth, as minerals progressively transform to less volatile-rich assemblages (Fig. 1a). Although much of the slab H₂O is lost, a fraction is retained and recycled into the deep Earth in residual phases [13,30–37]. The liberated fluid is buoyant, and upon formation begins ascending toward the surface. Some fluid may move upward within the slab [38], but most evidently migrates into the overlying mantle wedge (Fig. 1a).

Movement into the mantle wedge is an important step for subduction-zone fluids. Not only is there a sudden shift in host-rock composition, but in addition mantle minerals are strongly undersaturated in volatiles. At equilibrium, free H₂O cannot exist in the mantle until formation of a fully hydrated mineral assemblage (serpentine, chlorite, talc, and amphibole) or hydrous mineral stability is exceeded. Fig. 1b illustrates that near the slab, the mantle H₂O content can increase to >5 wt.%, but this changes with position. The consumption of fluid by the mantle begins at the shallowest levels in the forearc [18,39,40], and continues to at least sub-arc depths. During subduction, the slab and overlying mantle become mechanically coupled, causing mantle material to be dragged downward (Fig. 1). This provides a mechanism to continually supply fresh, volatile-poor mantle for the uptake of fluid. Despite the efficiency of this process, some slab-derived fluid may travel great distances before being consumed, and locally may even reach the surface (Fig. 1a). A natural example of the surficial venting of slab-derived fluids may occur at Kinki, Japan [41].

The down-going mantle cuts across mineral stability boundaries, causing volatile-bearing minerals to regenerate the fluid they earlier absorbed (Fig. 1b). Several such events may occur, until stable minerals become nominally anhydrous. From this point on, the mantle can no longer consume the fluid phase, leaving it free to flow upwards to trigger arc-magma production. Hence, a given fluid “particle” in subduction zones experiences a stepwise history, with discrete flow events interspersed between periods of movement while dissolved in solids.

Subduction-zone fluids migrate along paths of increasing temperature with decreasing pressure (Fig. 1c). The fluid path illustrated in Fig. 1 involves a T increase from ~ 500 °C at 100-km depth (~ 3.2 GPa), to ~ 1150 °C, at 80-km depth (~ 2.4 GPa). The increase of 650 °C over ~ 20 km (32.5 °C/km) probably represents a maximum gradient because the model system on which the figure is based represents an old, cold subduction system [42]. Nevertheless, increasing T during decompression is unusual among terrestrial fluid-flow systems, and it has important consequences for chemical behavior.

Fluid flow in the mantle wedge may involve porous flow, channeled flow, or a combination [20,43–45]. During porous flow, fluids intimately interact with the rock matrix, leading to a strong potential for continuous equilibration with the host material along the flow path, such that the rock matrix controls the composition of the fluid. By contrast, channelized flow produces zones in which the fluid-to-rock ratio is much higher. The fluid interacts with less rock per unit volume and retains more of its initial source composition. Porous and channelized flow will have significantly different velocities of several m/year vs. hundreds of m/year, respectively. U–Th–Ra disequilibria in arc magmas [46,47] and distances implied by Fig. 1 suggest 2.5 to ~ 100 m/year, which is consistent with both channelized and porous flow. It is quite possible that the velocity variations are real, reflecting a spectrum of hydrologic settings.

3. Chemical controls on the composition of subduction-zone fluids

The key step in identifying the chemical controls on fluid composition in the slab and mantle wedge is the direct measurement of mineral solubility at subduction-zone P and T. Recent advances have provided the first systematic observations of this kind [48–57]. Results demonstrate large solubility increases with increasing pressure. What governs this behavior? Important factors include the properties of solvent H₂O, the association/dissociation behavior of dissolved solutes, ligand concentrations, and the extent of formation of dense, solute-rich fluids that are intermediate between melts and H₂O.
3.1. H₂O at subduction-zone conditions

Because of its volumetric dominance, H₂O controls the properties of subduction-zone fluids. Water’s polar character and greater tendency to dissociate make it a substantially more powerful solvent than other volatiles. The solvent properties of H₂O depend on the density, ordering, hydrogen bonding, and dissociation of H₂O molecules. H₂O density is 1.2–1.4 g/cm³ at sub-arc depths (Fig. 1).
and 1.0–1.1 g/cm$^3$ at the thermal maximum in the mantle wedge [58–62]. With increasing temperature, the short-range-ordered, tetrahedral packing of H$_2$O molecules begins to break down. Supercritical H$_2$O is largely disordered [63,64], and the hydrogen-bond network is disrupted. Although H$_2$O remains a predominantly molecular solvent up to at least 10 GPa and 1000 °C [65,66], its extent of dissociation increases significantly during subduction (Fig. 2). At sub-arc depths, pure H$_2$O has between 0.01 and 0.1% H$^+$ (neutral pH is 3–4). This is significant because, other things being equal, it will raise the concentration of solutes due to an increase in ionic strength.

### 3.2. Dissociation/association and ion hydration

Pressure and temperature have opposite effects on dissociation of salts in H$_2$O. At constant $P$, increasing $T$ causes ions to associate as the dielectric constant of H$_2$O increases and ion-hydration shells become less stable. At constant $T$, increasing $P$ causes dissociation by enhancing electrostriction of H$_2$O around ions, which reduces the volume of solvation. Fig. 2 shows that changes in $P$ and $T$ along subduction paths promote association in dilute solutions; however, dissociation increases dramatically with small increases in concentration. This further increases the capacity to carry dissolved solutes, and magnifies the effects of additional ligands. For example, high Cl in arc lavas [67,68] implies that slab-derived fluids carry chloride. Increasing chlorinity favors solution of metals because of metal-chloride complexing. Calcite and anhydrite solubilities in H$_2$O–NaCl are dramatically enhanced with increasing NaCl concentration at high pressure [50,69], and

![Fig. 2. Extent of dissociation (as measured by equilibrium constant, $K$) of H$_2$O, NaCl, and MgSO$_4$ as a function of pressure. H$_2$O and NaCl dissociation are computed along the high-$T$ part of the envelope of subduction paths in Fig. 1c. See [92] for data sources and methods. MgSO$_4$ is taken from experiments [120] at H$_2$O density of 1.0 g/cm$^3$.](image-url)

![Fig. 1. (a) Results of a numerical simulation of H$_2$O production and migration in a model subduction zone [42]. The model includes solid mantle flow, assumes mineral-fluid-melt equilibrium in the slab (MORB – H$_2$O) and mantle wedge (peridotite – H$_2$O) system, and approximates aqueous fluid migration by porous flow along a pressure gradient. Subduction rate is 6 cm/year and the slab is 130 million years old. Colors represent rock H$_2$O content, solid lines denote isotherms (200 °C intervals, surface is 0 °C), and dashed lines indicate mantle flow paths. The location of the volcanic front is schematic. Water content decreases with depth in the slab due to H$_2$O loss from hydrous minerals. Large portions of the mantle in the model are at least partly hydrated by this fluid; some fluid crosses the moho and locally reaches the surface. High H$_2$O contents in the high-$T$ region of the mantle beneath the volcanic arc signify sites where melting can be expected. This conceptual framework is consistent with high-precision studies of trace elements and isotopic variations across and along volcanic arcs [109–118]. (b) Enlargement of region in (a) showing schematic path of a slab-derived fluid, with mantle H$_2$O contents (wt.%). The fluid migrates into the mantle wedge (solid orange arrows), where it is absorbed through formation of hydrous minerals. Downward flow of solid mantle (dashed arrows) causes dehydration. After multiple hydration–dehydration steps, the fluid enters a region where it is stable with anhydrous minerals, which allows greater travel distances. (c) Pressure–temperature diagram showing representative conditions at the slab–mantle interface (shaded) in northwest and southeast Japan [119], the coolest of which (left side) corresponds to results in (a). The fluid-flow trajectory represents the schematic path from (a) and (b). Also shown are $P$ and $T$ of fluid compositions discussed in the text. Open symbols designate fluids equilibrated with crustal mineral assemblages; filled symbols designate fluids equilibrated with mantle (Table 1). Abbreviations: CR, Costa Rica [83]; MP, Marianas (Pacman Seamount) [82,85]; MC, Marianas (Conical Seamount), [85,108]; EC, MORB eclogite [92]; JP, jadeite peridotite [94]; GP, garnet–orthopyroxene [95]; LH, Lherzolite [96]. (d) Schematic changes in fluid composition along the flow path in (c). Concentration of silicate components derived from the rock matrix increase downstream because of rising temperature, leading to dilution of the slab component. Slab-derived compatible elements are lost close to the slab, but incompatible elements may be transported to the magma source region. In detail patterns will be more complicated owing to changes in mineral stability and fluid flux.](image-url)
minor-element patterns may be strongly affected [70].

3.3. Critical curves and second critical end points

Interpretation of subduction-zone fluids is made difficult by uncertainty about melt-vapor miscibility and the existence of critical end-points in rock–H₂O systems. It is commonly assumed that fluids in subduction zones may attain properties intermediate between hydrous silicate liquid and H₂O. The expected enhancement of transport has led to the frequent indictment of intermediate fluids as metasomatic agents. However, such fluids exist only under quite restricted conditions, and their direct relevance to subduction zones is problematic. Part of the uncertainty derives from complex relationships whose depiction may seem inscrutable to all but the most ardent fan of phase diagrams. A brief overview is offered here.

3.3.1. Background

One-component systems, such as H₂O or mineral A (Fig. 3), possess a critical point marking termination of the distinction between the liquid and the vapor phase. At T and P above this point, there is only one phase, a supercritical fluid. In a system with both A and H₂O, the two critical points are linked together by a “critical curve” that marks the boundary between the stability region of a single supercritical fluid, and that of two fluids, a denser liquid and a less-dense vapor. Another curve extends from the point at which ice and mineral A coexist with liquid and vapor (the A-ice eutectic). This is the “solubility curve,” which marks the stable coexistence of mineral A with liquid and vapor (Fig. 3A). Compositions of liquid, gas, and supercritical fluid vary along critical and solubility curves. For example, with increasing T along the solubility curve, the liquid composition changes continuously from nearly pure H₂O to pure A. At high T, where the liquid is rich in A, the solubility curve is equivalent to the hydrous melting curve (the “solidus” in the simple system shown). Two-component systems of geologic interest exhibit a range of behaviors. In some, the critical curve and solubility curve remain separated over their entire lengths (Fig. 3A); in others, they intersect (Fig. 3B) to yield two “critical end-points.” The lower (first) critical end-point typically lies near the critical point of H₂O. The upper, or second, critical end-point lies at high P and T, potentially near subduction paths.

3.3.2. Critical behavior in albite–H₂O

The curious relationships indicated by critical behavior, in particular of the second type (Fig. 3B), have led to speculation about a role for supercritical fluids in subduction zones. This can be explored using the system albite–H₂O for illustration. The critical curve of albite–H₂O lies at relatively low P (Fig. 4a) [71], below that of melt generation in the mantle wedge. The albite–H₂O critical curve probably intersects the solubility curve at ~ 670 °C, 1.6 GPa (Fig. 4a) to generate a second critical end-point. Two schematic subduction paths illustrate the effects. Along Path 1 (Fig. 4a,b), albite solubility in H₂O increases until, at the solubility curve (H₂O-saturated melting, 685 °C, 1.5 GPa), it is ~ 10 wt.%. Above this point, only hydrous, albite-rich liquid can coexist with H₂O-rich vapor (Fig. 4b). Liquid + vapor become fully miscible at ~ 700 °C, 1.55 GPa (Fig. 4a), which signifies passage of the crest of the liquid + vapor field (Fig. 4b). Along Path 2 (Fig. 4a,c), C2 produces a narrow PT region (shaded) in which albite solubility increases continuously in a coexisting fluid that varies from dilute solution to hydrous albite-rich liquid— there is no discrete melting point.

3.3.3. Critical behavior and subduction zones

Despite the intriguing behavior of albite–H₂O, the significance of intermediate fluids in subduction zones is questionable. Critical behavior is strongly dependent on composition. Complete miscibility occurs at low to moderate P in many Na-rich systems [73]. Fluorine, boron and excess Na shift the critical curve to dramatically lower T at a given P [74]. A natural example has been reported in a pegmatite, in which complete miscibility of F-, B- and P-rich fluid occurs at 712 °C, 21.5 wt.% H₂O [75]. However, the critical curves of potassic and mafic systems lie at much higher P [76]; e.g., 12 GPa, ~ 1150 °C in MgO–SiO₂–H₂O [77]. The system K₂O–SiO₂–H₂O remains subcritical to at least 2 GPa at 1100 °C [78]. Critical curves for mantle systems (lherzolite–CO₂–H₂O) are likely 7.5 GPa or higher at ~ 1000 °C [79]. More information
is needed, but it appears that critical curves for compositions more closely approximating subduction-zone rocks lie at substantially higher $P$ than that for albite–H$_2$O.

Even if phase relations can be approximated by albite–H$_2$O, Fig. 4 reveals important restrictions on supercritical fluids. Other phases in the same chemical system may play an important role. Albite transforms to jadeite + quartz at high $P$, with different solubility and a change in solidus slope (Fig. 4a). No sign of a supercritical fluid has been noted above the solidus at high $P$, possibly pointing to structural changes in the silicate-rich liquid that could cause the liquid + vapor field to reappear.

Another problem for intermediate fluids is that the critical curve seems to be a feature of very H$_2$O-rich compositions at the $T$ of interest. Again taking albite–H$_2$O as a model, a closed system with several wt.%
H2O (Fig. 4b,c) will contain albite everywhere along both paths at conditions in Fig. 4. Such a system on Path 1 will encounter neither the L + V field (which requires more than \( \sim 20 \) wt.% H2O) nor the critical curve (\( \sim 50 \) wt.% H2O; Fig. 4b) [71,72]. On Path 2, albite coexists with an intermediate fluid only over a narrow PT interval (Fig. 4a,c). Comparison of the compositions of the albite-saturated fluid phase (vapor, liquid, or supercritical fluid; Fig. 4d) shows that little difference results from passage of the critical point. Contrasts between supercritical and subcritical solubility behavior will increase with greater separation of the subduction path from C2, but only if “fanning” of iso-concentration lines increases and there are no other phases (e.g., jadeite and quartz) to change the bulk solubility of the solids (Fig. 4a).

Even if they form, intermediate fluids can only be important metasomatic agents if they separate from their source and travel significant distances. However, with movement comes a new chemical environment...
and different \( P \) and \( T \). Rocks encountered along the mantle-wedge flow path will not be in equilibrium with the fluid, which will likely result in precipitation of much of the solute load over short length scales.

### 3.4. Polymerization of silicate components

Although I have argued that fully miscible fluids are not as important in subduction zones as has been supposed, the critical behavior holds a clue to what may be a fundamental aspect of deep fluid chemistry. A supercritical fluid can change continuously from pure \( \text{H}_2\text{O} \) to hydrous melt; i.e., solutes can evolve from hydrated ions or molecules, through small clusters, to the polymerized network of a hydrous silicate liquid. Fully miscible behavior requires that polymerization and network formation by silicate components in the aqueous phase is an important aspect of the chemistry even of dilute \( \text{H}_2\text{O} \)-rich subduction-zone fluids.

An early, direct observation of silicate polymers in dense fluids was by Mysen [78], who identified aqueous silica dimers and trimers coexisting with \( \text{K}_2\text{O} \)--\( \text{SiO}_2 \)--\( \text{H}_2\text{O} \) melts. These structures are marked respectively by two and three \( \text{Si} \) cations linked by bridging oxygens. Phase-equilibrium and in-situ studies confirm that silica polymerization is significant at subduction-zone \( P \) and \( T \) [49,52–54]. At 800 °C, 1.2 GPa, aqueous silica is a mix of monomers and dimers, with dimers increasing from 0 to 70% from pure \( \text{H}_2\text{O} \) to quartz saturation [51].

Its predominance in solution and ability to polymerize mean that silica plays a central role in the generation of solute polymers. However, liquid–vapor miscibility in aluminous compositions [71,73] suggests that \( \text{Al} \) solubility is enhanced because it participates in polymerization, as in silicate liquids. This is consistent with the stability of aqueous \( \text{Al} \)--\( \text{Si} \) complexes in crustal fluids [80]. Other elements may have solubility enhanced by participation in aqueous polymerization reactions (e.g., \( \text{Mg} \) [33,81]). Thus, complexing among \( \text{Si} \), \( \text{Al} \), and other elements via formation of polymerized solute may play an important role in controlling fluid composition at depth in subduction-zone settings. The partitioning of elements between the bulk fluid, the silicate polymer network in the fluid, and the rock matrix likely controls the overall compositional evolution of subduction-zone fluids. It is possible that concentrations of nominally insoluble elements may be enhanced through their participation in polymers, even in fairly dilute solutions.

### 4. Composition of subduction-zone fluids

The discussion of physical and chemical controls on subduction-zone fluids provides a framework for examination of constraints on composition. Fig. 1 shows that fluids may be produced over a range in depth, and that their upward flow into the mantle carries them across a major compositional boundary on a path of increasing temperature with decreasing pressure. Fig. 1d illustrates the expected behavior of typical silicate components from the matrix along this path. Solubility will increase because the increase in \( T \) has a larger effect than the drop in \( P \). For a slab-derived fluid in the mantle, this leads to dilution of the slab component as the mantle minerals progressively dissolve into the fluid. However, the relative solubility of a given element in the rock vs. the fluid will be a complex function of \( P \), \( T \), and composition. Those elements that are strongly partitioned into the rock relative the fluid (“compatible elements”) will be lost from the fluid very near its source. The result is (1) a metasomatic zone near the slab–mantle interface that is rich in compatible elements such as \( \text{Si} \), \( \text{Ca} \), \( \text{Al} \), etc., and (2) difficulty in transferring the slab signal for such elements far into the mantle. In contrast, elements that are strongly partitioned into the fluid relative to the rock (“incompatible elements”) will remain in the fluid as it travels away from the slab, allowing the compositional signature of the slab to travel well into the mantle wedge. Within this framework, we can examine existing constraints on fluid composition in subduction zones (Table 2).

#### 4.1. Major elements

Direct samples of subduction-zone fluids are available from the Costa Rica and the Izu-Bonin/Mariana convergent margins [82,83]. At Costa Rica, décollement fluids are sourced at 10–15 km, 100–150 °C [84]. They have 28 g/kg \( \text{H}_2\text{O} \) total dissolved solids (TDS), dominated by \( \text{Na} \) and \( \text{Cl} \), with low \( \text{Si} \) and a modest load of alkali and alkaline earth metals. Chlorinity is below seawater.
Pore fluids from Mariana forearc serpentine-mud volcanoes originate at the slab–mantle interface at 15–25 km \([82,85,86]\). Cl concentration is lower than seawater, consistent with a dehydration origin. Ca decreases and CO\(_2\) (alkalinity) increases with depth, reflecting the onset of slab decarbonation between 15- and 25-km depth\([85]\). Mg and Ca are lower than in Costa Rica fluids, perhaps due to lower chlorinity. Lower Si probably reflects the serpentinite host.

Direct samples have unknown reaction-flow history prior to sampling, so conditions of last equilibration are unclear. Fluid inclusions in exhumed blueschist- and eclogite-facies oceanic mafic rocks represent an alternative sample. Fluids show 1–7 wt.% NaCl equivalent \([87]\), or TDS = 10–75 g/kg H\(_2\)O. This spans values of shallow fluids. Increases in TDS with depth over 25–50 km are no more than about a factor of two, and low salinity and CO\(_2\) indicate that H\(_2\)O is the dominant solvent.

It is important to note that primary fluid inclusions in eclogites associated with continental subduction may have much higher N\(_2\), CO\(_2\), and salinity (up to 50 wt.% NaCl equivalent) \([11,88–91]\). There appears to be a fundamental difference in the salinity of fluids associated with oceanic and continental subduction, as recorded by fluid inclusions. Cross-comparisons between fluids in these two environments require extreme caution.

In the absence of direct samples of unreacted fluids or fluid inclusions, we must rely on estimates from mineralogic phase relations and experiments. Theoretical analysis of deeper fluid produced at the blueschist-eclogite transition indicates yet higher TDS (48 g/kg H\(_2\)O) \([92]\). This is largely due to the dramatic increase in silica at similar Cl content. Na and Ca concentrations are broadly similar. Al concentration, which is so low in low-pressure fluids that it is rarely analyzed, is higher than Mg. Varying Cl does not have a large effect. The high Al suggests that deep fluids leaving the slab are Na–Ca–Al–Si rich, but low in Mg and Fe, unless significant Cl is present. This is consistent with vein minerals in blueschists and eclogites \([87,93]\).

Insights into the compositional evolution of slab-derived fluids as they react with the mantle wedge can be gained from experimental studies. Aqueous fluid in equilibrium with jadeite–peridotite at conditions near the slab–mantle interface (Fig. 1) is Si- and Na-rich, and contains substantial Al \([94]\). Total dissolved solids is similar to the results for eclogite with Cl-free fluid. The higher Si reflects the strong dependence of Si concentration on temperature at high pressure \([49]\).

In the region of arc-magma generation, a 5-molal NaCl solution in equilibrium with garnet and orthopyroxene at 900 °C, 2 GPa, has nearly twice the silica, as well as higher Mg, Ca, and Al, low Na and

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<th>Table 2</th>
<th>Comparison of major-element compositions of subduction-zone fluids</th>
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<tr>
<td></td>
<td>Costa Rica décollement</td>
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<tr>
<td>Depth to slab (km)</td>
<td>0.36</td>
</tr>
<tr>
<td>P (GPa)</td>
<td>486</td>
</tr>
<tr>
<td>T (°C)</td>
<td>0.084</td>
</tr>
<tr>
<td>Cl</td>
<td>411</td>
</tr>
<tr>
<td>Si</td>
<td>3.3</td>
</tr>
<tr>
<td>Mg</td>
<td>23</td>
</tr>
<tr>
<td>K</td>
<td>12.9</td>
</tr>
<tr>
<td>Ca</td>
<td>28</td>
</tr>
<tr>
<td>Al</td>
<td>4</td>
</tr>
<tr>
<td>Na</td>
<td>31</td>
</tr>
<tr>
<td>Source</td>
<td>([83])</td>
</tr>
</tbody>
</table>

Concentrations in millimol per kg H\(_2\)O, except alkalinity = meq/kg H\(_2\)O, total dissolved solids (TDS) = grams total solute per kg H\(_2\)O.
Fe, TDS of 83 g/kg H₂O [95]. H₂O equilibrated with spinel lherzolite and Ti phases at 2.0 GPa, 1000 °C, shows lower Na, Al, and Si, but is otherwise similar to the garnet–orthopyroxene fluid [96]. Differences in Na between the fluids at 900–1000 °C and those at 550–660 °C are probably due to varying Na in the buffering assemblage.

Three important conclusions can be drawn from Table 2 about the major-element compositions of subduction-zone fluids. First, the deeper fluids are similar to the direct, shallow samples in one important way: total solute concentrations are low, regardless of pressure, temperature, or chlorinity. Deep fluids have TDS only two to three times that of seawater, and no more than 50% higher than shallow fluids from near the entrance to the subduction zone. TDS does not approach tens of wt.%, as would be associated with supercritical fluids of intermediate composition (Fig. 4). This is consistent with the independent arguments developed above.

The second conclusion is that TDS, though generally modest, nevertheless increases with depth. This arises from changes in solubility of rock-forming minerals due to the PT-enhancement of the solvent power of H₂O; additional ligands such as Cl may be present, but are not required.

Finally, there are important changes in major elements with depth. The dominant solutes in deep fluids are Si and Na. Al concentrations are higher than Ca, Fe, and usually Mg. Thus, the solutes in H₂O-rich fluids in subduction zones are dominated by alkali and alumino-silicate components. This contrasts with fluids from shallow environments, where alkali and other metals predominate and Al is virtually insoluble. It is quite likely that this change with increasing depth in subduction zones simply reflects the ability of alkali alumino-silicate components to form aqueous polymers at high P and T.

4.2. Trace elements

Trace-element patterns support the inference that the continental crust formed by island-arc magmatism (Fig. 5), but enrichments and depletions preclude simple anhydrous melting and fractionation. Instead, arc-magmas form by hydrous melting of a source metasomatized by slab-derived components. Trace-element patterns of low-Cl subduction-zone fluids are broadly similar to island-arc basalt (IAB) and continental crust, with enrichments in LILE and Pb, and depletions in high field strength elements (Fig. 5). However, >1 molal chloride changes some aspects of the patterns [70]. While Pb enrichment, Nb depletion, and a decreasing abundance with compatibility remain, the 5 molal brine shows very high Rb/Ba, Th/U ~ 0.01, enhanced Pb enrichment, and extreme differences between adjacent elements. This would appear to suggest that the trace-element signature of the IAB source is controlled by flow of low-Cl fluid [97,98]. However, the simple model compositions require more H₂O in the IAB source than is typically inferred [12], and isotopic studies support a wide range of chlorinity [99–102]. While there are likely to be real variations in Cl content in the global subduction system, there is also a need for more sophisticated reactive flow models that account for changes in mantle mineral assemblage along with evolution of fluid and rock composition.

5. Prospects for the future

This paper has attempted to highlight some of the open questions regarding the chemistry of
subduction-zone fluids. H₂O is a powerful solvent at subduction zone conditions, and it can effectively mobilize many rock components. Although supercritical, intermediate fluids are commonly invoked to explain mantle-wedge metasomatism, the hypothesis is problematic in detail. Supporting this, theoretical and experimental constraints on fluids at sub-arc conditions are surprisingly dilute. It may be that the chemistry of aqueous silicate polymers, as controlled by P, T and the mineralogy of the host, plays a fundamental role in the composition of subduction-zone fluids. The effects of Cl, and by inference other ligands, may be profound, but it is unclear at this time what role is required of them.

There are excellent prospects for progress on these issues in the near future. Experimental advances, including the hydrothermal diamond anvil cell [103–106] and solid media solubility techniques [48–51], promise to provide fundamental data on mineral solubility and fluid composition. Advances in molecular dynamics and ab-initio molecular dynamics simulations are already giving unprecedented insight into the nature of H₂O and ion hydration, geometry and structure [64,107]. Finally, new geochemical tools, including light element isotope ratios (e.g., Li and B), new spectroscopic methods and ICP-MS, are being applied to fluid and melt inclusions in subducted oceanic rocks, mantle-wedge xenoliths and arc magmas. Results give new insight into the details of slabs and arc-magma sources, which constrain the initial and final conditions in the reacting system of interest here.

The new data will provide the foundation necessary to develop sophisticated reaction-flow models of this complex metasomatic environment. Once this is under way, we will have made a major advance toward unraveling the chemical workings of this part of the subduction factory.

Acknowledgements

This study was supported by NSF EAR 9909583 and 0337170. Reviews by J. Ayers, G. Bebout and an anonymous reviewer improved the manuscript. I am also indebted to R. Newton, J. Davidson, and S. Peacock for insightful critiques. [AH]

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