
Publisher's PDF, also known as Version of record

License (if available):
CC BY

Link to published version (if available):
10.2138/am-2021-7752

Link to publication record in Explore Bristol Research

PDF-document

This is the final published version of the article (version of record). It first appeared online via Mineralogical Society of America at https://doi.org/10.2138/am-2021-7752. Please refer to any applicable terms of use of the publisher.

University of Bristol - Explore Bristol Research

General rights

This document is made available in accordance with publisher policies. Please cite only the published version using the reference above. Full terms of use are available: http://www.bristol.ac.uk/red/research-policy/pure/user-guides/ebr-terms/
The glass transition and the non-Arrhenian viscosity of carbonate melts

DONALD B. DINGWELL1, KAI-UWE HESS1,* MARTIN C. WILDING2,5, RICHARD A. BROOKER3, DANILO DI GENOVA3, JAMES W.E. DREWITT4, MARK WILSON4, AND DANIEL WEIDENDORFER1

1Earth and Environmental Sciences, Ludwig-Maximilians-Universität München, Theresienstraße 41/III, 80333 München, Germany
2Materials and Engineering Research Institute, Sheffield Hallam University, Howard Street, Sheffield, S1 1WB, U.K.
3School of Earth Sciences, Wills Memorial Building, Queens Road, Bristol, BS8 1RJ, U.K.
4Department of Theoretical Chemistry, University of Oxford, South Parks Road, Oxford, OX1 1K
5U.K. Catalysis Hub, Research Complex at Harwell, Rutherford Appleton Laboratory, Harwell,OX11 0DE, U.K.

Abstract

We report the first calorimetric observation of the glass transition for a carbonate melt. A carbonate glass [55K2CO3-45MgCO3 (molar)] was quenched from 780 °C at 0.1 GPa. The activation energy of structural relaxation close to the glass transition was derived through a series of thermal treatments comprising excursions across the glass transition at different heating rates. Viscosities just above the glass transition temperature were obtained by applying a shift factor to the calorimetric results. These viscosity measurements (in the range of 109 Pa·s) at supercooled temperatures (ca. 230 °C) dramatically extend the temperature range of data for carbonates, which were previously restricted to super-cool temperatures well below 1 Pa·s. Combining our calorimetrically derived results with published alkaline-earth carbonate melt viscosities at high temperatures yields a highly non-Arrhenian viscosity-temperature relationship and confirms that carbonate liquids are “fragile.” Based on simulations, fragile behavior is also exhibited by Na3CO3 melt. In both cases, the fragility presumably relates to the formation of temperature-dependent low-dimensional structures and Vogel-Fulcher-Tammann (VFT) curves adequately describe the viscosity-temperature relationships of carbonate melts below 1000 °C.

Keywords: Viscosity, glass transition, fragility, carbonate melt, carbonatite, shift factor

Introduction

Melt viscosity has a fundamental control on many Earth processes, from how melts are transported through the deep mantle and crust, to its influence on effusive or explosive eruption styles at the Earth surface (McKenzie 1985; Dingwell 1996). Although physical properties (e.g., viscosity, density, surface tension) are becoming well-established for multicomponent silicate systems of importance in earth sciences (Lange and Carmichael 1987; Knoche et al. 1995; Bagdassarov et al. 2000; Giordano et al. 2008), carbonate melts are less well-studied in the geosciences, due to relatively rare volcanic expression at the Earth’s surface (Keller 1989; Dawson 1966). Carbonate volcanism is nevertheless a part of the Earth system and may be more common on other planets. Carbonatite volcanism has been invoked for so-called “canali”—channels on Venus, which reach up to 6800 km in length, implying a very low-viscosity flow (Kargel et al. 1994). Highly mobile carbonate melts migrating within the Earth likely play a major role as metasomatizing agents (Blundy and Dalton 2000; Hamouda and Laporte 2000) and in the subduction-related recycling of carbon back to Earth’s surface as part of the global carbon cycle (Dasgupta and Hirschmann 2010; Thomson et al. 2016). Their ability to scavenge high concentrations of RRE and rare metals make them increasingly important as multi-commodity exploration targets for the green energy revolution (Simandl and Paradis 2018). The extreme physical and chemical properties (e.g., density, viscosity, conductivity, reactivity), which set carbonatites apart from most silicate melts, must have a strong influence on the interaction and migration dynamics of buoyant carbonate melts throughout the lithosphere after separating from their source region (Hunter and McKenzie 1989; Minarik and Watson 1995; Brooker 1998; Hamouda and Laporte 2000; O’Leary et al. 2015). The relationship between low viscosity and electrical properties of carbonate-rich melts has also been linked to regions of high conductivity in the upper and lower mantle (Stfere et al. 2014; Gaillard et al. 2008).

These physical properties also have commercial applications, particularly in the design of molten carbonate fuel cells, which generally offer a prospect of greener electricity but may now also be designed with a capacity to efficiently capture CO2 derived from the liquid, gaseous, or solid hydrocarbon (i.e., coal) fuel used for generating electricity (Cassir et al. 2012). Such molten carbonates generally provide excellent performance at low cost in thermal energy storage or heat transfer fluid systems (Wu et al. 2011).

Recent investigations of carbonate melt at the molecular scale have revealed a temperature- and pressure-dependence of structure that likely affects physical properties (Wilding et al. 2016, 2019a, 2019b; Wilson et al. 2018). Thus, a more complete understanding of the relationship between carbonate melt structure and physical properties is required. Here, we provide the first ever calorimetric determination of the glass
transition of a carbonate glass and generate a direct measurement of the viscosity of a carbonate liquid just above the glass transition interval. In this manner we are able to constrain the non-Arrhenian temperature-dependence and thereby the fragility of molten carbonates, providing a parameterization valuable for the extrapolation and interpolation in temperature of the transport properties of carbonates.

Viscosity data: The carbonate challenge

Natural carbonatites are mainly composed of alkaline-earth (Ca, Mg, and Fe) alkali (Na and K) carbonates. Rheological measurements on the most geologically relevant end-member carbonates have proven difficult. This is especially true for alkaline earth carbonates as they do not melt congruently at atmospheric pressure (1 atm). For example, CaCO$_3$ decarbonates to CaO+CO$_2$ below 0.004 GPa and 1240 °C and then melts incongruently (to CaCO$_3$+CaO or CaCO$_3$+CO$_2$-rich melts) up to 0.3 GPa and 1375 °C. Only above this pressure and temperature is congruent melting observed (Smyth and Adams 1923; Wyllie and Huang 1976). For MgCO$_3$ and FeCO$_3$ the conditions are even more extreme (see Wyllie and Huang 1976; Weidner 1972). Viscosity measurements are less complicated for the alkali carbonates, as they melt congruently at atmospheric pressure, although they can be subject to dissociative CO$_2$ loss whose intensity decreases in the order Li >> Na > K (Janz and Lorenz 1961).

In Figure 1a, the ambient pressure, single alkali carbonate measurements imply a clear compositional effect of the different alkali cations, but large scatter also exists even within a single study. As is typically observed for very restricted ranges of temperature and viscosity, all data [except (L,N,K)C] can be described within error by an Arrhenian temperature dependence (a linear fit on a log$_{10}$ η vs. 1/T plot). As later noted by Janz et al. (1989), their initial “NIST standard” viscosity data were incorrect (and therefore are not included in Fig. 1a). For Na$_2$CO$_3$, they defer to the data of Ejima et al. (1984), which has a less steep slope, almost identical to the Sato et al. (1999) data in Figure 1a. Janz et al. (1989) suggest that the error and differences in general arise from comparisons of various measurement methods, experimental assemblies, and working equations, as well as matters of sample pre-treatments and impurities. Also included in Figure 1a are green stars, representing new 50Na$_2$CO$_3$-50K$_2$CO$_3$ data from this study (see Results), measured using the same equipment as described by Di Genova et al. (2016). This data set covers a larger temperature range, and so has a better-defined slope (green dotted line). The higher pressure Na$_2$CO$_3$ data of Stagno et al. (2018) in Figure 1a perhaps illustrates the challenge of performing accurate high-pressure measurements, and it remains unclear whether they define any systematic effect of pressure. Also included in Figure 1a are the data from Kim et al. (2015) for a 22Li$_2$CO$_3$-33Na$_2$CO$_3$-45K$_2$CO$_3$ composition (purple stars). This eutectic in the Li$_2$CO$_3$-bearing ternary system allows an even greater temperature range in measured viscosities. With such a large temperature range considered, some curvature clearly becomes evident, indicating a non-Arrhenian temperature dependence and raising questions regarding the validity of a linear fit to the other data.

Figure 1b contains a much more diverse range of mainly mixed carbonate compositions. Many involve alkaline-earth
cations and therefore require pressure to ensure congruent melting. It is difficult to untangle the effect of composition from that of pressure, but the Dobson et al. (1996) study appears to include a range of data points on single lines for two compositional data sets covering a range of pressures from 2.5 to 5.5 GPa. These two lines generally bracket the different compositions of Kono et al. (2014), which cover an even larger pressure range from 0.9 to 5.3 GPa. In terms of composition, the green dotted line is representative of the alkali carbonates at 1 atm, when transferred to Figure 1b and could indicate either a lower viscosity for the alkali carbonates or some pressure effect with respect to the other high-temperature data. The lower viscosity of high-pressure measurements for Na₂CO₃ (yellow hexagons) compared to other compositions, but above the 1 atm data (green dotted line), appear to suggest both effects may be operating. Note that the K₂CO₃ high-pressure (4.0 GPa) data point of Dobson et al. (1996) was subsequently discredited (Liu et al. 2007; and therefore are not included in Fig. 1b), again illustrating the challenges in measuring physical properties using high-pressure experiments (Kono et al. 2014; Stagno et al. 2018). All the other high-temperature data in Figure 1b indicate the effect of pressure is less than half a log₁₀ unit between 0.9 and 5.5 GPa. The measured data appears at odds with the simulations of Vuilleumier et al. (2014), Du et al. (2018), and Desmaele et al. (2019a), which suggest a clear and more pronounced pressure effect. The seminal CaCO₃ data of Vuilleumier et al. (2014) is included in Figure 1b as an example, although these simulations may have overestimated the pressure effect to some extent and the role of forces such as van der Waals interactions require further refinement for these ab initio calculations (Vuilleumier, personal communication).

The glass transition and viscosity of “fragile” melts

As an aid to resolving some of the issues described above, an extension of the temperature range for viscosity determinations of carbonate liquids is a prime experimental goal. Most commercial glasses involve liquids with “covalent” network-forming components such as silicate, aluminate, or borate, and when cooled their increase in viscosity hinders nucleation and results in a glass with a characteristic, calorimetric glass transition reflecting this kinetic arrest (Moynihan 2019). Glass formation is however not restricted to network-forming liquids (Tangeman et al. 2001; Dingwell et al. 2004; Kohara et al. 2011). Glasses can even be formed from organic liquids and metallic systems. Of possible relevance to this study are glasses formed in simple ionic molten salts such as sulfate (Förland and Weyl 1950; MacFarlane 1984) and nitrate systems (Van Uitert and Grodkiewicz 1971). For these ionic compositions, the temperature-dependence of viscosity, a vital parameter in glass-formation, rarely exhibits Arrhenian behavior. This departure from Arrhenius law behavior is at the heart of the concept of liquid “fragility,” a term introduced by Angell (1985) to describe and compare the viscosity-temperature relations of a variety of glass-forming liquids. The quantification of fragility has been provided by the model-independent, so-called melt fragility index m (Böhmer et al. 1993). This index describes the rate of viscosity change of a melt with temperature at T_g, defined as the temperature where the viscosity is equal to 10¹² Pa·s. The exact mechanism is thought to be related to medium-range structural heterogeneity, with faster and slower relaxing regions developing as liquids are supercooled. There are formal links to the changes in the structure of supercooled liquids via the Adam-Gibbs model of structural relaxation (Gibbs and Adam 1965; Angell et al. 2000a, 2000b). Viscosity measurements made close to the glass transition can be used to evaluate liquid fragility (Gottsmann et al. 2002).

The linear Arrhenius law for viscosity (η) is expressed as:

\[ \log(\eta) = A + B/E_a/T \]  

and has two parameters to be fitted, E_a, the activation energy is considered as the energy barrier that must overcome for examples as bonds are broken allowing molecules to “flow” from one position to another, and A is a numerical constant.

Thus, although viscosity-temperature data such as those in Figure 1 are commonly approximated over limited temperature ranges to an Arrhenian fit, it has long been known that for many materials the value of E_a changes with temperature yielding a non-Arrhenian curvature on a reciprocal absolute (Arrhenian) plot (Angell 1985; Böhmer et al. 1993). This is particularly evident as data are added from lower (supercooled) temperatures. One method of fitting such curvature is using the Vogel-Fulcher-Tammann equation (Vogel 1921; Fulcher 1925; Rault 2000) equation of the form:

\[ \log(\eta) = A + B/[T(K) - C] \]  

and B defined as: \( B = (E_a \times 2.303)/R \) with R denoting the gas constant. This has the potential to reconcile a steep reciprocal temperature-dependence of low-temperature viscosity data with a lower reciprocal temperature dependence of high-temperature viscosity data and give the expected intersection of a log₁₀ value near –4 (in Pa·s) at infinite temperature.

Carbonate glasses

At the accessible quench rates of most experimental equipment (hundreds to thousands of K/s), carbonate melts rarely survive undercooling sufficient to produce glasses. As a result, direct investigation of their physico-chemical properties in the supercooled regime is currently limited to just the few known systems where glasses can be readily formed, MgCO₃-K₂CO₃ (Eitel and Skaliks 1929; Datta et al. 1964; Ragone 1966; Genge et al. 1995) and La(OH)₃-Ca(OH)₂-CaCO₃-CaF₂-BaSO₄ (Jones and Wylie 1983; Genge et al. 1995). Eitel and Skaliks (1929) were apparently the first to report a carbonate glass, quenched from ~50:50 mol% MgCO₃-K₂CO₃ melts at 0.12 GPa. They also observed rapid devitrification of the glasses on an hour-timescale when held at a temperature between 205–300 °C. This is a common response to crossing the glass transition within poorly glass-forming systems such as nitrates, sulfates and carbonates and even basaltic silicates (Böhmer et al. 1993; Angell et al. 2000a; Wilding et al. 2000; Nichols et al. 2009). This implies the presence of T_g in this region of temperature and that the glass transition temperature for K₂Mg(CO₃)₃ might be located near 200 °C. To the best of our knowledge, this estimate has not been confirmed or used in the intervening 90 years, although Datta et al. (1964) did return to the system and mapped the glass-forming region.

Here, we report the first quantitative determination of a glass
transition for a carbonate liquid. Specifically, we demonstrate the
temperature and viscosity of a mixed K$_2$CO$_3$-
MgCO$_3$ carbonate melt by performing scanning calorimetry and
viscometry on samples of a supercooled carbonate phase. Using
this low-temperature viscosity data combined with published
higher temperature (high pressure) data as well as both classical
and ab initio molecular dynamics simulation estimates, we
determine whether a non-Arrhenian Vogel-Fulcher-Tammann
(VFT) law fit is possible and quantifies the extent of fragility.

**Materials and Methods**

**Glass synthesis**

Carbonate glass was prepared using a starting mixture of 55 mol% K$_2$CO$_3$, and
45 mol% MgCO$_3$, (subsequently referred to as 55K$_2$CO$_3$–45MgCO$_3$) using a rapid
quench, cold-seal pressure vessel at the University of Bristol. This composition
is directly above the eutectic (at ~460 °C) on the binary join (Ragone 1966), and
glasses can be formed easily at the quench rates achieved (~200 K/s) at a pressure
of 0.1 GPa. Reagent-grade K$_2$CO$_3$ (>99.9%) was dried at 400 °C. The MgCO$_3$
was in the form of a natural, optically clear, and inclusion-free magnesite crystal
(Brumado, Brazil), and transmission FTIR was used on thin cleavage fragments
to confirm that this material was virtually water free. The starting materials were
ground together and loaded in 3.8 mm long gold capsules, which
were then welded shut and loaded into a Tuttle-type cold-seal pressure vessel with
a rapid quench rod extension (Ihinger 1991). The experiments were run at 780 °C,
0.1 GPa for ~10–15 h and quenched (>200 K/s). The resulting glass was removed
from the gold capsule mostly as a single solid slug, representing the central part
of the quenched melt, with an outer section that tends to spall off. This was stored
in a desiccator. A small section was immediately set in dental resin and polished
of the measurement setup (some seconds) and is assumed to be uncorrupted by
inspection after the measurement revealed a surficial tarnishing of the sample
and error were calculated (Hess et al. 2019b). This was also confirmed by H NMR on similar samples (Wilding et al. 2019b). Wilding et al. (2019b) suggest this water is most likely introduced during loading or because the powders were improperly dried (in the case of the $^{13}$C-enriched NMR sample). A comparison with the IR spectra of Genge
et al. (1995) suggests our samples contain significantly less water although it is
very difficult to quantify. There is no evidence to suggest that the water content
of the glass changes over the course of at least 24 h as demonstrated by a time
series of spectra published in Wilding et al. (2019b). However, over the course of several weeks, samples can deteriorate even when stored in a desiccator and
change in appearance from a transparent, pristine glass to an opaque white pow-
der. This particular sample was transferred in one piece from Bristol to Munich
within a few days of production and broken to extract a fresh interior sample for the
measurements. The degree of hydration is unlikely to change over the course of
the subsequent calorimetry experiments.

In addition, a 50Na$_2$CO$_3$-50K$_2$CO$_3$ (molar) carbonate mix was produced from
Merck high-purity carbonates (Na$_2$CO$_3$–99.9%; K$_2$CO$_3$–99.5%) to compare with the
previously published end-members in Figure 1a, but over a larger temperature
range than pervious data sets to better define the slope on a 1/T plot.

**Thermogravimetry, calorimetry, and viscometry**

Combined thermogravimetric analysis (TGA) and differential scanning calorimetry (DSC) was performed using a Netzsch STA 449 C/3/G Jupiter simul-
taneous viscometry analyzer at LMU. A single, transparent chip of the carbonate
glass, of about 10 mg, was heated up to 920 °C at a constant rate of 5 K/min in a Pt-crucible under a purging, high-purity Argon atmosphere (30 mL/min; see
Fig. 2). The temperature calibration was based on the melting points of indium,
zinc, Ba-carbonate, and gold. The measurements were corrected for thermal drift.
Differential Scanning Calorimetry measurements at greater sensitivity were made
using a Netzsch DSC 404 C/3/P Pegasus calorimeter. This also involved a chip of
glass (40 mg), placed in a platinum crucible and heated with constant
rates of 5, 10, and 15 K/min up to a temperature of 250 °C under a high-purity
Argon atmosphere (30 mL/min; see Fig. 3). The sample remained transparent until
the end of the heating cycles. The melting points of indium, zinc, baryum carbonate,
and gold were also used to calibrate temperature for this instrument.

In addition, one shear viscosity measurement has been performed at 230 °C
using the micro-penetration method under a purging, high-purity argon atmosphere (30 mL/min). For this measurement, the sample was held for 15 min at this dwell
temperature prior to the initiation of indentation to establish thermal equilibration.
A hemispherical iridium-indenter was used with a force of 1 N into the sample,
and the penetration depth was monitored with time. The corresponding viscosity
do not appear to have been significantly influenced by this.

The viscosity value was immediately taken after the mechanical relaxation of the measurement setup (some seconds) and is assumed to be uncorrupted by
crystallization effects.

![Figure 2](image)

**Figure 2.** DSC/DTA measurements of a single 55K$_2$CO$_3$–45 MgCO$_3$
glass chip heated to 750 °C. The onset of the glass transition is seen at
about 225 °C (see inset). The relaxed glass is then devitrified shown by
two exothermic crystallization peaks at ~300 and ~340 °C. On further
heating the sample partly decomposes and forms a mixture of carbonates
and oxides, which cannot be quenched to for a glass. The stable mixture
melts at 900 °C.

![Figure 3](image)

**Figure 3.** High-sensitivity DSC measurement of a 40 mg sample of
55K$_2$CO$_3$–45 MgCO$_3$ glass with a series of excursions across the
glass transition with heating rate matched to prior cooling rate at the
rates 5–15 K/min.
High-temperature 50NaCO₃-50K₂CO₃ viscosity measurements were made using an adapted commercially available concentric cylinder rheometer (see Di Genova et al. 2016) equipped with a graphite furnace and featuring an air-bearing-supported synchronous motor and a specially designed Pt-Au concentric cylinder crucible and spindle assembly. With this adaptation, high-accuracy viscosity measurements of highly fluid melts can be achieved at high temperatures, up to 1273 K, and at extremely low torques. This allows accurate viscosity measurements as low as 10⁻¹³ Pa·s (and up to 10⁻⁴ Pa·s) at shear rates up to 10⁴ s⁻¹. The apparatus was calibrated with distilled water, silicone oils, and the DGG-1 standard glass (see Di Genova et al. 2016). Super-liquids 50Na₂CO₃-50K₂CO₃ melt was measured from 1023 to 1198 K at a shear rate of 20 s⁻¹ (see Table 2).

### First-principle (ab initio) molecular dynamics

The ab initio MD simulations for the 55K₂CO₃-45MgCO₃ melt composition were performed using the Vienna Ab Initio Software Package (VASP) (Kresse and Furthmüller 1996a, 1996b). The electronic interactions described by projector-augmented wave (PAW) pseudopotentials (Blöchl 1994; Joubert 1999) were computed at the Brillouin zone Γ-point only with an energy cut-off value of 550 eV and the Perdew-Burke-Ernzerhof (PBE) formulation of the generalized gradient approximation (GGA) exchange correlation functional (Perdew et al. 1996). Molecular dynamics trajectories were computed in the canonical (NVT) ensemble with periodic boundary conditions and a Nosé thermostat (Nosé 1984).

The system was simulated using a randomly generated cubic cell of 222 atoms (44 Mg, 40 C, 120 O) with initial closest approaches defined from the lowest crystallographic interatomic bond lengths. The starting configuration was super-heated to 10,000 K, was equilibrated at the Brillouin zone Γ-point only, and converted in error in viscosity.

### DTA glass transition determination

The results of differential thermal and thermogravimetric analysis (DTA-TGA) and differential scanning calorimetry (DSC) are shown in Figures 2 and 3 and tabulated in Table 1 for the 55K₂CO₃-45MgCO₃ glass. The DTA analysis (Fig. 2) shows clearly the glass transition onset at ~220 °C (493 K). There is a small peak at 120 °C, which we interpret as the loss of surface water from these samples. Above 260 °C (533 K) the DTA curves show a series of exothermic peaks, which result from crystallization, followed by phase transitions (above 400 °C) and finally at higher temperatures, endothermic peaks that represent a solid-state reaction and partial decarbonation resulting in a mixture of carbonates and oxides. The onset of melting occurs at 900 °C (1173 K). On cooling, crystallization occurs at the same temperature, and thus liquidus and solidus temperatures are consistent. This liquidus temperature for the 55K₂CO₃-45MgCO₃ composition is much higher than the 500 °C expected from the 0.1 GPa phase diagram of Ragone (1966). The reported eutectic in the

### RESULTS

### DSC and DTA glass transition determination

<table>
<thead>
<tr>
<th>Method</th>
<th>T (°C)</th>
<th>Quench rate (K/s)</th>
<th>Viscosity (Pa·s)</th>
<th>Error</th>
<th>Pressure (atm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>DSC¹</td>
<td>237</td>
<td>-0.60</td>
<td>8.55</td>
<td>0.09</td>
<td>1 atm</td>
</tr>
<tr>
<td>DSC²</td>
<td>234</td>
<td>-0.78</td>
<td>8.73</td>
<td>0.09</td>
<td>1 atm</td>
</tr>
<tr>
<td>VISC-MP</td>
<td>230</td>
<td>-1.08</td>
<td>9.03</td>
<td>0.09</td>
<td>1 atm</td>
</tr>
<tr>
<td>VISC-FS³</td>
<td>800</td>
<td>-1.44</td>
<td>0.43</td>
<td>3 GPa</td>
<td></td>
</tr>
<tr>
<td>VISC-FS³</td>
<td>900</td>
<td>-1.66</td>
<td>0.43</td>
<td>3 GPa</td>
<td></td>
</tr>
<tr>
<td>VISC-FS³</td>
<td>1200</td>
<td>-2.22</td>
<td>0.43</td>
<td>5 GPa</td>
<td></td>
</tr>
</tbody>
</table>

¹ Using a shift factor of 7.95; error in T converted in error in viscosity.
² From Dobson et al. (1996).

### Table 2. Diffusion coefficients obtained from simulations and used to derive viscosities

<table>
<thead>
<tr>
<th>Potassium</th>
<th>Carbon</th>
<th>Sodium</th>
</tr>
</thead>
<tbody>
<tr>
<td>T (K)</td>
<td>log D (cm²/s)</td>
<td>log η (Pa·s)</td>
</tr>
<tr>
<td>--------------------</td>
<td>-----------------</td>
<td>-----------------</td>
</tr>
<tr>
<td>1500°</td>
<td>-4.875</td>
<td>-2.789</td>
</tr>
<tr>
<td>1800°</td>
<td>-3.998</td>
<td>-2.949</td>
</tr>
<tr>
<td>2100°</td>
<td>-3.857</td>
<td>-3.023</td>
</tr>
<tr>
<td>2500°</td>
<td>-3.538</td>
<td>-3.267</td>
</tr>
<tr>
<td>1850°</td>
<td>-4.000</td>
<td>-2.935</td>
</tr>
</tbody>
</table>

² Values derived from ab initio calculations.
³ Values derived from The classical simulations presented in Wilding et al. (2019a) where only the K diffusivity is provided.
⁴ Values derived from Wilson et al. (2018) for a favored ‘q’ value of 2.28 (see Online Materials 1).

---

American Mineralogist, vol. 107, 2022
binary system (57K2CO3–43MgCO3 at 0.1 GPa) is at 460 °C, although the liquidus rises very steeply with composition. This may suggest a change in composition and the loss of CO2 at ambient pressure consistent with the observation of Eitel and Skaliks (1929) that high pressure is required to ensure melting of K2CO3 and MgCO3 constituents without decomposition, but we note that any such effect is post-Tpeak determination.

The results of the high-sensitivity DSC measurements are shown in Figure 3. The as-quenched glass was heated through the glass transition into the supercooled liquid regime at 250 °C (523 K) where the liquid is fully relaxed. The glass sample was then subjected to a series of thermal cycles comprising excursions across the glass transition at different heating rates, with the heating rate matched to the prior cooling rate (i.e., for example, a heating rate of 5 K/min is matched with a previous cooling rate of 5 K/min) (Eastoe et al. 1977). With this approach of matching cooling and heating rates, the so-called onset of the calorimetric glass transition corresponds to the fictive temperature (Moynihan et al. 1993; Yue et al. 2004) determined by the enthalpy-match method (Moynihan et al. 1976; Yue 2008). As the sample is heated at rates of 5, 10, and 15 K/min, the temperature of the glass transition (taken as the peak position corresponding to the heat flow peak overshoot, hereafter referred to as Tpeak) shifts systematically with matched heating/cooling rate (q) from 229 to 237 ± 1 °C (502–510 K) for 5 to 15 K/min. The combined thermo-gravimetry (TG) and calorimetry (DSC) measurements demonstrate the thermal stability of the glass sample; the glass transition is clearly identified at 229 ± 1 °C (for a cooling rate of 15 K/min), which is consistent with the original study of Eitel and Skaliks (1929) based on rapid devitrification of the glass between 200–300 °C.

**Shift factor**

The shift factor $K$ (here $K_{peak}$) (Scherer 2006; Stevenson et al. 1995; Gottsmann et al. 2002; Yue et al. 2004; Al-Mukadam et al. 2020) can be derived from the DSC measurements (Fig. 3) by matching the characteristic glass transition temperature $T_{peak}$ for a given cooling/heating ($q$, K/s) cycle to the temperature of the viscosity measurement:

$$K_{peak} = \log_{10}q \left( T_{peak} \right) + \log_{10}q$$

(3)

The value (7.95) obtained at cooling/heating rates of 15 K/min (where the temperatures overlap with the accuracy of the measurements) is then used for the other cooling/heating cycles:

$$\log_{10}q \left( T_{peak} \right) = K_{peak} - \log_{10}q$$

(4)

to calculate the viscosity for the 55K2CO3–45MgCO3 liquid (Table 1) ranging from log$_{10}$ 9.03 ± 0.09 Pa·s at 229 °C to log$_{10}$ 8.55 Pa·s ± 0.09 at 237 °C (error in temperature was converted to error in viscosity). We note that this shift factor is lower than that observed for silicate melts but is generally consistent with high compressibility as expected for carbonate melts.

**A viscosity–temperature relationship for carbonate melts**

For the 55K2CO3–45MgCO3 sample, the three viscosities derived using DSC measurements and the shift factor together with the one data point from the micro-penetration measurement are plotted in Figure 4. These low-temperature data can be fitted to a single Arrhenian trend, but the slope greatly exceeds that of an Arrhenian fit solely through the high-temperature 50K2CO3–50MgCO3 data of Dobson et al. (1996) or Sifré et al. (2015), or the trends of simulation data plotted in Figure 4. It is clear that both Arrhenian subsets of data exhibit slopes that are Arrhenian only as artifacts of the very restricted temperature ranges of the individual data sets. Taken together, the data describe a strongly non-Arrhenian temperature-dependence of viscosity. An unweighted Vogel-Fulcher-Tammann (VFT) fit through both data sets is shown in Figure 4. The resultant fit using the Dobson et al. (1996) data demonstrates the “fragility” of this system. At high temperatures, this fit also agrees well with the 50K2CO3–50MgCO3 data of Sifré et al. (2015), who measured the electrical conductivity and demonstrated how this high-temperature data could be converted to a viscosity. It must be borne in mind, however, that both Dobson et al. (1996) and Sifré et al. (2015) data are at high pressure (3–6 GPa). As the magnitude of any pressure effect remains controversial and it is not possible to conduct high-temperature viscosity measurements at 1 atm to directly complement the DSC data, we have further extended (for further comparison) the 55K2CO3–45MgCO3 data set using ab initio simulation data in Table 2. These 1 atm results are included in Figure 4, where they lie close to the Dobson et al. (1996) high-pressure data. In fact, our simulations indicate that from 1 atm to 3 GPa, there is less than half log$_{10}$ unit increase in viscosity. Also included in Figure 4 is an additional simulation data point for “ambient” pressure 55K2CO3–45MgCO3 melt derived from the classical simulations of Wilding et al. (2019a, 2019b), which are consistent with high-energy XRD data used to elucidate the 55K2CO3–45MgCO3 glass structure. When evaluating the reliability of the Dobson et al. (1996) 50K2CO3–50MgCO3 viscosities, Kono et al. (2015) implied that they should lie 0.5 log$_{10}$ unit higher to match the 50K2CO3–
50CaCO₃ Dobson et al. (1996) data as no compositional effect was expected. Desmaele et al. (2019a) also concluded that the compositions should have similar viscosity but also indicated that the effect of pressure would reduce this value by a \( \log_{10} \) unit at 1 atm. Given all these unresolved discrepancies, we have chosen to fit to the Dobson et al. (1996) 50K₂CO₃–50MgCO₃ data accepting they are possibly too high and that a reduction in pressure brings them down and closer to our 1 atm simulated data for 55K₂CO₃–45MgCO₃. Regardless of which high-temperature data set is used, it is clear the 55K₂CO₃–45MgCO₃ melt is highly fragile. The VFT fitting also indicates a value between –3 and \(-4 \log_{10}\) (in Pa·s) at an infinite temperature consistent with other theoretical and statistical estimates (Angell et al. 1989). A calculated VFT for Na₂CO₃ is also illustrated for comparison in Figure 5. This represents the viscosity obtained from the diffusion data obtained from the classical MD simulations of Wilson et al. (2018), which are tabulated in Table 2. The “q” value, the degree of charge separation across the carbonate anion (explained in Online Materials) is 2.28, and the simulated diffraction pattern at this value gives the best fit to the data obtained from high-energy XRD measurements on molten Na₂CO₃. The diffusion (and viscosity) data obtained at this value of charge separation shows fragile (non-Arrhenian) behavior, correlated with the temperature-dependent abundance of CO₃⁻ rings and other complexes. Using this value of charge separation gives the best VFT fit to the experimental Na₂CO₃ viscosity data of Di Genova et al. (2016) in Figure 5 and illustrates the robustness of this simulation methodology.

Figure 5 also contains our new viscosity measurements for 50Na₂CO₃–50K₂CO₃, listed in Table 3. This composition is consistent with the Di Genova et al. (2016) data set plotting between viscosity data curves for Na₂CO₃ and K₂CO₃ when measured under an argon atmosphere (see Fig. 1a). In the context of comparing to the Na₂CO₃ VFT curve, this data has the advantage of a larger temperature range than the Na₂CO₃ measurements. The 50Na₂CO₃–50K₂CO₃ binary viscosity data set can be fitted with a linear trend within error, but a slight positive curvature would also be permitted. The curved trend for the 22Li₂CO₃–33Na₂CO₃–45K₂CO₃ eutectic composition of Kim et al. (2015), as noted in Figure 1, is reproduced in Figure 5 and is also consistent with our Na₂CO₃ VFT curve.

The other synthetic composition data shown in Figure 5 are simulations performed by Desmaele et al. (2019b) that reveal the effect of adding calcium to the sodium-potassium system. Starting from the 50Na₂CO₃–50K₂CO₃ composition of this study, there is a clear increase in viscosity with the addition of CaCO₃. Although Desmaele et al. (2019b) provide only two data points, they are consistent with a decrease in fragility with CaCO₃ addition.

### Table 3. High-temperature viscometry data for synthetic alkali carbonate melts

<table>
<thead>
<tr>
<th>Temperature (°C)</th>
<th>Viscosity (10^5 Pa·s)</th>
<th>Log Viscosity (Pa·s)</th>
<th>Shear rate (s⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>750</td>
<td>7.975</td>
<td>0.00598</td>
<td>20</td>
</tr>
<tr>
<td>775</td>
<td>9.542</td>
<td>0.00554</td>
<td>20</td>
</tr>
<tr>
<td>800</td>
<td>9.320</td>
<td>0.00515</td>
<td>20</td>
</tr>
<tr>
<td>825</td>
<td>10.97</td>
<td>0.00481</td>
<td>20</td>
</tr>
<tr>
<td>850</td>
<td>8.905</td>
<td>0.00453</td>
<td>20</td>
</tr>
<tr>
<td>875</td>
<td>8.711</td>
<td>0.00429</td>
<td>20</td>
</tr>
<tr>
<td>900</td>
<td>8.525</td>
<td>0.00409</td>
<td>20</td>
</tr>
<tr>
<td>925</td>
<td>8.347</td>
<td>0.00385</td>
<td>20</td>
</tr>
</tbody>
</table>

*Error is ±0.01 log (Pa·s).*

**Figure 5.** The sodium carbonate VFT fit has been derived from the simulations of Wilson et al. (2018), but with selected parameters to best fit the sodium carbonate data of Di Genova et al. (2016). Other published data for synthetic alkali carbonate mixes ranging from pure alkalis, to variable amounts of added calcium carbonate. Data from: † Desmaele et al. (2019b); ‡ Kim et al. (2015); † Di Genova et al. (2016); ‡ Sato et al. (1999).

**Discussion**

**Carbonate melt structure**

The structure of levitated alkali carbonate liquids has been studied directly using high-energy X-ray diffraction (Wilding et al. 2016; Wilson et al. 2018). These diffraction data have been combined with classical molecular dynamic simulations and used to evaluate the changes in the liquid structure with temperature and accordingly the carbonate liquid fragility. Central to the simulation methodology is the flexibility of the carbonate anion geometry. Spectroscopic studies of the K₂CO₃–MgCO₃ glass (Sharma and Simons 1980; Genge et al. 1995; Wilding et al. 2019a, 2019b) have suggested the presence of two structurally distinct populations of carbonate anion, one of which is more distorted, and it is this distortion of the carbonate that has been used as a basis for the molecular dynamics simulation of the alkali carbonate liquids. In the simulation of Na₂CO₃, flexibility is imposed by employing springs between the O-O and C-O pairs in the anion, and the stiffness of these springs is constrained by comparing the liquid diffraction patterns at high and low values of scattering vector with the simulated liquid structure, the inter- and intra-molecular contributions, respectively. Once constrained, the simulated liquid structure can be explored as a function of temperature. The flexibility of the molecular carbonate anion allows the central carbon atom to be drawn out of the triangular plane and the anion becomes polarized. One consequence of this is that there is the development of a secondary length scale and the development of carbonate chains and other carbonate complexes, the extent of which is strongly temperature dependent. In a more recent study (Wilson et al. 2018), the fluctuation of charge across the molecular anion was introduced into the simulation to further explore the dynamics of sodium carbonate liquids and demonstrates a correlation between the development of carbonate complexes and liquid fragility, with
the connectivity of these emergent structures dependent on the mean charge separation (see Online Materials).

This modeling approach, with the flexibility of the molecular anion, has also been applied to the $\text{K}_2\text{CO}_3$-$\text{MgCO}_3$ system where the simulations are used to identify the pressure-dependent changes in structure of the same $5\text{K}_2\text{CO}_3$-$4\text{MgCO}_3$ glass studied here. At ambient pressure, there is no evidence for formation of carbonate chains or other complex structures in the $5\text{K}_2\text{CO}_3$-$4\text{MgCO}_3$ carbonate liquids. However, as pressure is increased, there is the development of a carbonate network associated with an increase in the mean coordination number of the carbon with the development of a CO$_{3+4}^-$ configuration (Wilding et al. 2019a). The response to pressure in these liquids again reflects the flexibility of the carbonate and the complex interaction between the oxygen atoms in the CO$_{3+4}^-$ anion and their strong electrostatic interaction with potassium cations. The ambient pressure glass does not show the development of this network, but $^{13}$C NMR confirms a distorted carbonate anion (Wilding et al. 2019b), while infrared spectroscopy confirms the presence of two structurally distinct populations of carbonate anions identified in earlier studies. In fact, the simulation of the liquids suggests different degrees of flexibility of the carbonate anion rather than two distinct populations with the stronger interactions with Mg$^{2+}$ and K$^+$ cations associated with the more distorted carbonate. Nonetheless, the simulation of the $\text{K}_2\text{CO}_3$-$\text{MgCO}_3$ liquid shows distorted carbonate with magnesium and potassium cations occupying irregular channels and with both types of cations adopting a network-forming role by bridging isolated carbonate anions (Wilding et al. 2019b). It has been suggested that glass formation in sulfate and nitrate systems also requires the presence of two different cations with different field strengths and different degrees of polarizability (Förland and Weyl 1950; van Uitert and Grodkiewicz 1971; MacFarlane 1984; Wilding et al. 2017).

The ab initio molecular dynamics simulations that form part of this study can be used to evaluate the changes in the ambient pressure liquid structure as a function of temperature. As expected, there is no formation of a Na$_2$CO$_3$ style carbonate network in $5\text{K}_2\text{CO}_3$-$4\text{MgCO}_3$. However, the main changes in structure occur in the local environment surrounding the K$^+$ cations. There is an increase in the mean K-O coordination number, which increases as the temperature is decreased, but no change in the average C-O coordination at ambient pressure. The distortion in the local environments for potassium is shown in the changes in the partial contributions to the pair distribution functions for K-O and K-K (Online Materials). The coordination environments for both potassium and magnesium differ significantly from those in the equivalent crystalline phase, illustrated in a snapshot from the ab initio simulation in Figure 6.

As noted above, the $\text{K}_2\text{CO}_3$-$\text{MgCO}_3$ composition is not a naturally occurring carbonatite liquid. However, this and related studies show that combining experimentally derived structures and structure-related properties with classic and ab initio simulation provides insight into the viscosity-temperature relations of mixed carbonate liquids and that the same modeling approach can be extended to naturally occurring systems. In contrast to the alkali carbonates, there is no evidence for the formation of a carbonate network in the $\text{K}_2\text{CO}_3$-$\text{MgCO}_3$ liquids. However,

**Figure 6.** A snapshot of the high-temperature liquid configuration, carbon atoms are shown in black, oxygen atoms red, potassium atoms blue, and magnesium atoms yellow, obtained directly from the ab initio MD (VASP) simulation trajectories for liquid $5\text{K}_2\text{CO}_3$-$4\text{MgCO}_3$ at 2500 K.

the liquid dynamics still reflect the underlying flexibility of the carbonate anion.

### The fragility of carbonate melts

The carbonate melt viscosities are compared in reciprocal absolute temperature (Arrhenian) space with other geologically relevant liquids in Figure 7 using the data in Table 4. This comparison confirms that under similar conditions, the carbonate liquids have very low viscosity and are potentially very mobile, at least at the

### Table 4. Fit parameters describing the temperature dependence of the viscosity (VFT), the glass transition temperature $T_g$, the activation energy $E_a(T_g)$ and the fragility index (m) for carbonates in this study and selected silicate liquids of geological relevance.

<table>
<thead>
<tr>
<th>Composition</th>
<th>A (K)</th>
<th>B (K$^{-1}$)</th>
<th>C (K$^{-1}$)</th>
<th>$T_g$ (K)</th>
<th>$E_a(T_g)$ (kJ/mol)</th>
<th>Fragility index (m)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\text{K}_2\text{CO}_3$</td>
<td>-7.219</td>
<td>15950</td>
<td>232.2</td>
<td>1108</td>
<td>489</td>
<td>23</td>
</tr>
<tr>
<td>$\text{Na}_2\text{CO}_3$</td>
<td>-5.550</td>
<td>10261</td>
<td>263.8</td>
<td>884</td>
<td>399</td>
<td>24</td>
</tr>
<tr>
<td>$\text{SiO}_2$</td>
<td>-4.167</td>
<td>15336</td>
<td>508.5</td>
<td>1457</td>
<td>693</td>
<td>25</td>
</tr>
<tr>
<td>$\text{K}_2\text{CO}_3$</td>
<td>-4.550</td>
<td>6101</td>
<td>567.0</td>
<td>936</td>
<td>753</td>
<td>42</td>
</tr>
<tr>
<td>$\text{SiO}_2$</td>
<td>-4.167</td>
<td>4500</td>
<td>508.5</td>
<td>1457</td>
<td>693</td>
<td>25</td>
</tr>
<tr>
<td>$\text{K}_2\text{CO}_3$-$4\text{MgCO}_3$</td>
<td>-4.010</td>
<td>1914</td>
<td>356.0</td>
<td>476</td>
<td>580</td>
<td>64</td>
</tr>
<tr>
<td>$\text{Na}_2\text{CO}_3$</td>
<td>-4.310</td>
<td>3703</td>
<td>761.7</td>
<td>989</td>
<td>1345</td>
<td>71</td>
</tr>
<tr>
<td>$\text{K}_2\text{CO}_3$-$4\text{MgCO}_3$</td>
<td>-3.602</td>
<td>798</td>
<td>324.4</td>
<td>376</td>
<td>824</td>
<td>115</td>
</tr>
</tbody>
</table>

Notes: Glass transition temperature $T_g$ at log$_{10} 12$ Pa s, activation energy $E_a$ at $T_g$, derived from VFT parameter.

$^a$ Data from Dorfman et al. (1996).
$^b$ Data from Giordano et al. (2009).
$^c$ Data from Hetherington et al. (1964) and Hofmaier and Urbain (1968).
$^d$ Data from Dingwell et al. (2004).
$^e$ Data from Tweer et al. (1971) and Weiler et al. (1969).
generally considered to be an archetypal fragile liquid. In Figure 7b, these are compared in an Angell plot (Böhmer et al. 1993; Angell and Moyniha 2000) with the temperature normalized to $T_g$ and the liquids show progressive departure from the Arrhenius behavior of “strong liquids” such as the network-forming liquid SiO$_2$, seen as a series of curves of increasing fragility. The different degrees of fragility reflect differences in the temperature-dependence of liquid structure through their relative contributions to the configurational entropy (Richet 1984). The temperature-dependent structural elements might be locally favored structures, dynamic heterogeneities, or density fluctuations within the supercooled liquid state.

**Implications**

It will be apparent that the fragility of the carbonate liquid and its high-temperature viscosity are very similar to those of a peridotite melt (Dingwell et al. 2004). Liquid peridotite is one of the most fragile silicate compositions ever measured and is extremely difficult to quench to a glass. Peridotite is an example of a “fully depolymerized” silicate melt, where the silicate network structure is broken into isolated units surrounded by metal cations (cf. Kohara et al. 2011). Peridotite and carbonate melts lie at higher fragility than all the other petrologically and volcanologically relevant silicate melts, including basalts (Giodano and Dingwell 2003), extremely peralkaline phonolites and pantellerites (Whittington et al. 2001; Di Genova et al. 2013), and water-rich calc-alkaline rhyolites (Hess and Dingwell 1996). Peridotite melt may have formed the magma ocean in early Earth history when carbonate content could also have been relatively high. In fact, at high enough pressures, there is a continuum from carbonate melt to “depolymerized” high-CO$_2$ silicate melt compositions (such as mellite) generated during high-temperature carbonated mantle melting (e.g., Brey and Green 1976; Gudfinnsson and Presnall 2005). Initially, SiO$_2$ that is added to dilute the carbonate melt will remain as isolated “SiO$_2$” units with little effect on the silica structure and, therefore, viscosity. But as SiO$_2$ (and Al$_2$O$_3$) reaches 15–30 wt% these transitional melt compositions (Brooker et al. 2011) form new structures with two sub-networks, one consisting of regions with polymerized silicate compositions (such as melilitites) generated during high-temperature carbonated mantle melting (e.g., Brey and Green 1976; Gudfinnsson and Presnall 2005). Initially, SiO$_2$, which is added to dilute the carbonate melt will remain as isolated “SiO$_2$” units with little effect on the silica structure and, therefore, viscosity. But as SiO$_2$ (and Al$_2$O$_3$) reaches 15–30 wt% these transitional melt compositions (Brooker et al. 2011) form new structures with two sub-networks, one consisting of regions with polymerized silicate structures and the other ionic carbonate. These may be precursors to silicate-carbonate immiscibility (Brooker et al. 2001; Morizet et al. 2017). The presence of two sub-networks at the molecular level could lead to a complex viscosity temperature dependence related to this medium-range structural heterogeneity, with the faster and slower relaxing regions that are often cited as a cause of fragility. However, the viscosity determinations of Morizet et al. (2017) and those inferred by the conductivity measurements of Sifré et al. (2014) suggest that various amounts of CO$_2$ in transitional or basaltic silicate melts and thus the proportion of the two sub-networks, has little effect on the viscosity, which remains similar to the CO$_2$-free silicate composition. This is consistent with our observations here that all very high-temperature interactions between extremely fragile very-low-silica silicate melts (e.g., peridotite, mellite, kimberlite) and carbonatites will be interactions between two liquids of similar low viscosities.

The absolute values of the glass transition temperatures, together with the restricted range of glass-forming ability in carbonatite, means that the likelihood of encountering glassy behavior in natural magmatic systems is vanishingly small.
The more rapid increase in viscosity at lower temperatures is obviously more important in silicate systems, particularly for silica-rich eruptions where volatiles are being exsolved. Gas bubbles are unable to escape, instead building up internal pressure that results in explosive behavior and fragmentation of a supercooled melt (i.e., glass) as the viscosity approached some critical value (usually considered around 10⁷ Pa·s; e.g., Namiki and Manga 2008). The stoichiometry of the carbonate components within carbonate melts suggests that the main volatile component (CO₂) is not necessarily exsolved, at least in alkali-rich systems. Even if other volatiles are exsolved, the general low viscosities at eruptive temperatures (see Fig. 7a) will allow bubbles to escape through the melt more efficiently than in a silicate melt. The exception to this could be the apparently explosive alkali-earth carbonatite volcanism that produced rounded lapilli melt droplets, preserved at the Kaiserstuhl volcanic complex (Keller 1989). These have an almost pure CaCO₃ composition, and the shape and internal structure suggest these are airborne quenched melt. Pure CaCO₃ should dissociate at pressures below 0.004 GPa, so how this composition could erupt molten droplets is still a mystery. This has prompted the usual idea that minor amounts of alkalis or fluorine allow the melt to exist at low pressure and these are subsequently leached away (Gittins and Iago 1991; Brooker and Kjarsgaard 2011). However, one could speculate that a rapid increase in viscosity and cooling within some “CO₂ confining pressure” in the vent could combine to preserve a CaCO₃ melt as droplets. Zimanowski et al. (1986) have demonstrated droplet formation for 50Na₂CO₃–50K₂CO₃ and Oldoinyo Lengai carbonatites by the interaction of the melt with injected water. As Zimanowski et al. (1986) point out, these processes are perhaps also important for modeling explosive situations involving industrial molten salt cells where a full understanding of the physical processes becomes important as regards hazard mitigation.

Zimanowski et al. (1997) have indeed directly observed the fracturing of a high-temperature carbonate liquid upon interaction with external water. They effectively induced the glass transition in the carbonate liquid and recorded the results by high-speed video. These observations are entirely consistent with the observation here that a glass transition can be encountered in molten carbonates. With knowledge of the viscosity–temperature curve observed here, together with the effective temperature of the melt–water interaction in the experiments of Zimanowski et al. (1997), one should be able in principle to estimate the volume strain rates at the point of brittle failure of the carbonate liquid. Alternatively, if the thermal stresses are well-estimated, then one should be able to predict the effective temperature of the brittle failure-induced melt–water interaction. That would have been impossible to achieve accurately with a degree of certainty by simply using extrapolations of super-liquidus viscosity-temperature relations.

A nonlinear rate change in viscosity is also an important parameter in understanding the nature of carbonate lava flows, particularly the length and terminal velocity at Oldoinyo Lengai. The same applies to the calculations of Treiman and Schell (1983), who calculated the carbonate melt properties in magma chambers and their effect on turbulence and the rapid setting velocities of crystals and growth rates on chamber walls, with implications also for the settling rate in lava flows (Norton and Pinkerton 1997).

The most widespread occurrence of carbonate melts is most likely deep in the Earth’s mantle. Dalton and Wood (1993) demonstrated that carbonatites generated by melting a depleted carbonate mantle source are almost alkali-free, ranging from 25MgCO₃–75CaCO₃ to 8MgCO₃–92CaCO₃ (with some minor FeCO₃), but sodium can reach 15Na₂CO₃ for a fertile mantle source and subsequently be increased or decreased due to metasomatic wall–rock reactions. The range of compositions derived from subducted altered basalt (eclogite) in Thomson et al. (2016) is even more diverse, ranging to higher contents of iron- and alkalis (14–24FeCO₃, 9–15MgCO₃, 43–66CaCO₃, 2–29Na₂CO₃, 1–3K₂CO₃). At these conditions, the viscosity can be reasonably well approximated by a linear temperature dependence trend. The possible compositional effect of Na₂CO₃ inferred from Figures 1b and 5 suggest this is an important component when considering the viscosity of mantle carbonatites, although these mantle melts are all produced at high temperatures (>1200°C) and even 1 atm sodic melts would have viscosities within 0.5 log₁₀ units of the other compositions in Figure 1b. However, the hydrous “fertile pyrolite” composition of Wallace and Green (1988) produced an alkali-rich carbonate melt (∼5K₂CO₃–33Na₂CO₃–62CaCO₃) at temperatures between 930 and 1080°C at 2.1 GPa. Wallace and Green (1988) suggest an even more alkali-rich carbonate melt may exist at even lower temperatures, perhaps forming before the water is released by hydrous silicate minerals. These low temperatures are realistic for the shallow lithosphere beneath the old continental crust, and this is getting into the temperature range where a linear fit based on the high-temperature data will become inaccurate, and the high fragility of the Na₂CO₃ component may become important (Fig. 5). This would have implications for modeling the transport properties of these melts to their surface expression or as metasomatizing agents.

Acknowledgments and Funding
We thank Fabrice Gaillard and Rodolphe Vuilleumier for fruitful suggestions that greatly improved the manuscript. R.A.B was funded by the NERC Thematic Grant consortium NE/M000491/1. J.W.E.D acknowledges support from NERC under Grant NE/P002951/1. D.B.D. acknowledges the support of ERC 2018 ADV Grant 834255 (EAVESDROP).

References cited
silicate liquids at pressure? Constraints from the fusion curve of K$_2$CO$_3$ to 3.2 GPa. Contributions to Mineralogy and Petrology, 153, 55–66.


