

Influence of intensive parameters and assemblies on friction evolution during piston-cylinder experiments

PIERRE CONDAMINE^{1,*}, SIMON TOURNIER¹, BERNARD CHARLIER², ETIENNE MÉDARD³, ANTOINE TRIANTAFYLLOU⁵, CÉLIA DALOU¹, LAURENT TISSANDIER¹, DELPHINE LEQUIN¹, CAMILLE CARTIER¹, EVELYN FÜRI¹, PETE G. BURNARD¹, SYLVIE DEMOUCHEY^{4,†}, AND YVES MARROCCHI¹

¹CNRS, CRPG, Université de Lorraine, F-54000 Nancy, France

²Department of Geology, University of Liège, 4000 Sart Tilman, Belgium

³CNRS, IRD, OPGC, Laboratoire Magmas et Volcans, Université Clermont Auvergne, F-63000 Clermont-Ferrand, France

⁴Géosciences Montpellier, CNRS, Université de Montpellier, F-34095 Montpellier, France

⁵Geology Laboratory of Lyon, Earth, Planets and Environment (LGL-TPE), Université Lyon 1, ENS de Lyon, CNRS, UMR 5276, Villeurbanne, France

ABSTRACT

Piston-cylinder assemblies exhibit inhomogeneous pressure distributions and biases compared to the theoretical pressure applied to the hydraulic press because of the thermal and mechanical properties of the assembly components. Whereas these effects can partially be corrected by conventional calibration, systematic quantification of friction values remain very sparse and results vary greatly among previous studies. We performed an experimental study to investigate the behavior of the most common cell assemblies, i.e., talc [$\text{Mg}_3\text{Si}_4\text{O}_{10}(\text{OH})_2$], NaCl, and BaCO_3 , during piston-cylinder experiments to estimate the effects of pressure, temperature, run duration, assembly size, and assembly materials on friction values. Our study demonstrates that friction decreases with time and also partially depends on temperature but does not depend on pressure. We determined that friction decreases from 24 to 17% as temperature increases from 900 to 1300 °C when using talc cells, indicating a friction decrease of ~2% per 100 °C increase for 24 h experiments. In contrast, friction becomes independent of time above 1300 °C. Moreover, at a fixed temperature of 900 °C, friction decreases from 29% in 6 h runs to 21% in 48 h runs, corresponding to a decrease of friction of 0.2% per hour. Similar results obtained with NaCl cell assemblies suggest that friction is constant within error, from 8% in 9 h runs to 5% in 24 h runs. At 900 °C, possible steady-state friction values are only reached after at least 48 h, indicating that friction should be considered a variable for shorter experiments. We establish that assembly materials (and their associated thermomechanical properties) influence the friction correction more than the dimensions of the assembly parts. Finally, we show that the use of polytetrafluoroethylene film instead of conventional Pb foil does not modify friction but significantly reduces the force required for sample extraction, thus increasing the lifetime of the carbide core, which in turn enhances experimental reproducibility.

Keywords: Experimental petrology, piston-cylinder, friction, assembly, calibration

INTRODUCTION

The piston-cylinder apparatus (Boyd and England 1960) is well established in experimental petrology and mineralogy for the synthesis of high-pressure and high-temperature geomaterials. Typical setups can attain pressures of 0.5–4 GPa and temperatures of 600–2000 °C, and special setups extend these ranges down to 0.3 GPa (e.g., Mirwald et al. 1975; Moore et al. 2008) and up to 2500 °C (Cottrell and Walker 2006). Piston cylinders are thus particularly well suited to investigate material properties at crustal to upper mantle conditions on Earth and even at core conditions on smaller planetary bodies. Furthermore, many thermodynamic models concerning the properties of the mantle and their evolution through geological time rely on experimental databases. It is thus critical to provide accurate experimental data with minimal uncertainties.

During high-pressure and high-temperature (HP-HT) piston-

cylinder experiments, multiple factors can lead to important biases on the pressure applied to the sample. Laboratories worldwide employ various pressurization and heating procedures to reach the P - T conditions of interest, such as the commonly used hot piston-in and piston-out techniques, leading to noticeably different applied pressures (Johannes et al. 1971; Shimizu and Kushiro 1984; McDade et al. 2002). It has also been suggested that a fraction of the hydraulic pressure is not transmitted to the sample due to heterogeneous pressure distributions and/or frictional strain between the carbide core and the cell assembly (Tamayama and Eyring 1967; Edmond and Paterson 1971). These pressure losses can be characterized by the friction value F (in %), defined as the difference between the applied hydraulic pressure P_{app} and the effective pressure on the sample P_{eff} :

$$F = \left(\frac{P_{\text{app}}}{P_{\text{eff}}} - 1 \right) \times 100 \quad (1)$$

The materials used in piston-cylinder cell assemblies vary widely depending on the purpose of the experiments (see Dunn

* E-mail: pierre.condamine@gmail.com

† Orcid 0000-0001-5023-4655

1993 for a review). For instance, different spacer and sleeve materials can be used around the sample, each possessing unique thermal and mechanical properties (Bell et al. 1971; Longhi 2005). Alumina is sometimes substituted for MgO spacers to complete graphite furnaces (Johannes et al. 1971; McDade et al. 2002); however, the different rheological properties of alumina and MgO (elasticity, ductility, hardness) can lead to over- and under-pressured zones around the sample. Furthermore, assemblies made with high-strength materials require greater pressure corrections than those made with low strength materials (Johannes 1978). It is therefore important to understand and quantify the role of material properties such as hardness, density, isothermal compressibility, and thermal expansion during HP-HT experiments.

Several studies have sought to (1) understand and model thermal uncertainties and reproducibility during piston-cylinder experiments (e.g., Watson et al. 2002; Médard et al. 2008) and (2) estimate the friction correction required depending on cell materials, including the commonly used talc, NaCl, and BaCO₃ pressure media (e.g., Fram and Longhi 1992; Bose and Ganguly 1995; McDade et al. 2002; Longhi 2005). However, the reproducibility of pressure and associated corrections (i.e., friction) during HP-HT experiments and which parameters control their evolution remain unclear. For example, Fram and Longhi (1992) claimed that the friction correction is highly dependent on pressure for BaCO₃ cells, whereas McDade et al. (2002) reported that friction is independent of temperature and pressure at 1000–1600 °C and 1.5–3 GPa. Therefore, the main objective of this study is to experimentally quantify the friction induced by the use of talc, NaCl, and BaCO₃ cells during piston-cylinder experiments. Whereas the use of talc cells is known to produce important pressure losses on the sample compared to the applied hydraulic pressure, friction values are only sparsely reported and vary widely (Johannes et al. 1971; McDade et al. 2002). Our second aim, then, is to understand and quantify the roles of experimental parameters (pressure, temperature, and experimental duration) and assembly materials and size on friction and its evolution during short- and long-duration experiments.

METHODS

Experimental strategy

Experiments were performed on three different reactions. First, we chose the quartz-coesite transition for its shallow *P-T* slope (Bose and Ganguly 1995; and references therein), which limits biases due to thermal gradients. This simple reaction precludes problems observed in other commonly used transitions, such as the synthesis of the starting material in the fayalite + quartz = ferrosilite transition (see Bohlen et al. 1980). In addition, the kinetics of this transition are remarkably fast (Lathe et al. 2005), allowing experiments to be performed over both short (2.5 h) and long durations (48 h), in turn allowing us to investigate the evolution of friction over time. We calibrated the quartz-coesite transition over a wide range of typical piston-cylinder temperatures, i.e., at 900 and 1300 °C, to understand the role of temperature on frictional issues in the three cell types (Fig. 1). Second, we investigated the albite (NaAlSi₃O₈) = jadeite (NaAlSi₂O₆) + quartz (SiO₂) reaction at 800 and 1000 °C (Holland 1980) using BaCO₃ cells. Third, we investigated the anorthite (CaAl₂Si₂O₈) + gehlenite (Ca₂Al₂SiO₇) + corundum (Al₂O₃) = kushiroite (3CaAl₂SiO₆) reaction (Hays 1966a, 1966b) at a constant pressure of ~1.3 GPa using talc cells to understand the role of pressure and assembly size (1/2" vs. 3/4") on friction at 1300 °C.

Because low-temperature and short-duration experiments exacerbate friction (Bose and Ganguly 1995), we performed additional experiments on the roles of

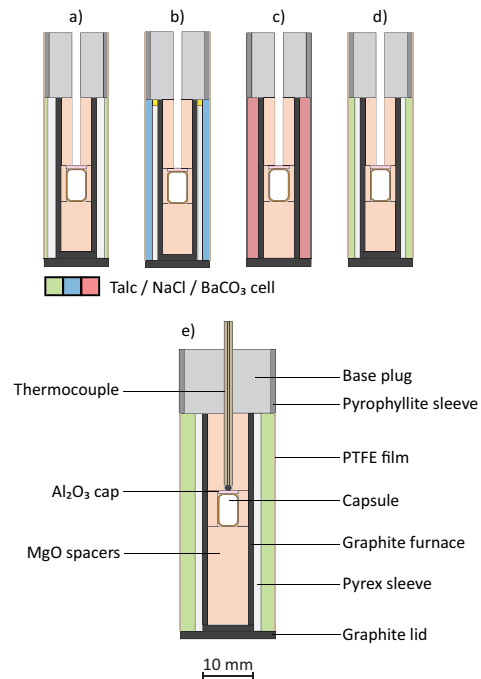


FIGURE 1. Schematic diagrams of the four different piston-cylinder cell assemblies used in this study. (a) Talc cell assembly used at CRPG (Nancy). (b) NaCl cell assembly used at LMV (Clermont-Ferrand). (c) BaCO₃ assembly used at ULG (Liège). (d) Batch 2 of the talc cell assembly used at CRPG, using a thicker cell and thinner Pyrex sleeve compared to a. The talc cell is 0.75 mm thick in a against 1.35 mm in d. Contrary to other assemblies, no Pyrex glass is present in the BaCO₃ cell assembly and a Pb film is used instead of PTFE to wrap the cell. See Online Materials¹ OM1 for additional details. (Color online.)

assembly size and component materials at 900 °C for 6 h. These experiments included replacing the MgO sleeve around the capsule with an alumina sleeve, using a polytetrafluoroethylene (PTFE) film wrapping the cell, and using higher density MgO spacers (2.8 vs. 2.0 g/cm³) in the graphite furnace (Table 1). A last series of experiments was performed on a new batch of assemblies (thicker talc cell, thinner Pyrex glass) at more conventional piston-cylinder conditions (1300 °C, 24 h) to understand the potential role of assembly part sizes on the generated friction (batch 2; Table 1; Fig. 1d).

Starting materials

For the quartz-coesite calibration experiments, we used analytical-grade amorphous SiO₂ (99.9 wt% purity) and added 6 wt% H₂O to each capsule with a micro syringe to enhance the reaction kinetics (Lathe et al. 2005). To study the kushiroite = anorthite + gehlenite + corundum reaction, we prepared a mixture based on the stoichiometry of the kushiroite, using analytical grade SiO₂, Al₂O₃, and CaCO₃ powders, ground in ethanol for 1 h, and then decarbonated in a furnace at 1200 °C during 24 h. The procedure was repeated to ensure complete decarbonation. Oxide powders were stored in an oven at 110 °C prior to capsule assembly. For the albite = jadeite + quartz reaction, the starting material consisted of a mixture of pure powdered crystals, validated by X-ray diffraction measurements on a Bruker D8-Advance diffractometer with CuK α radiations (ULiège). Proportions were set during mixture preparation, with 70 wt% quartz, 20 wt% jadeite, and 10 wt% albite. The powder mixture was stored in an oven at 120 °C prior to capsule assembly.

Experimental techniques

We performed 38 experiments on identical Voggenreiter Mavo LPC 250-300/50 end-loaded 12.7 mm (1/2") and 19.0 mm (3/4") piston-cylinders at three different institutes: the Centre de Recherches Pétrographiques et Géochimiques (CRPG, Nancy, France) for talc cell assemblies, the Laboratoire Magmas et Volcans (LMV,

TABLE 1. Experimental conditions and results

Assembly	T (°C)	Duration (h)	Run	P_{app} (GPa)	Product
Talc cell experiments (CRPG laboratory)					
PTFE-Talc-Pyrex-C-MgO	900	2.7	V130	3.97	qz
			V16	4.06	qz + co
			V10	3.87	qz
		4	V19	3.97	qz + co
			V108	3.77	qz
			V55	3.87	qz + co
		15	V41	3.77	qz
			V38	3.87	qz + co
			V30	3.68	qz
		24	V24	3.77	qz + co
			V33	3.58	qz
			V56	3.68	qz + co
		48	V37	3.77	qz
			V18	3.87	qz + co
			V22	3.77	qz
24	V57	3.87	qz + co		
	V114	1.45	an + gh + crn		
	V113	1.54	kush + an + gh + crn		
PTFE-Talc-Pyrex-C-Al ₂ O ₃	900	6	V32	3.87	qz
			V42	3.97	qz
Pb-Talc-Pyrex-C-MgO	900	6	V107	3.77	qz
PTFE-Talc-Pyrex-C-MgO (2800 kg/m ³)	900	6	V39	3.87	qz + co
			V109	3.68	qz
PTFE-Talc-Pyrex-C-MgO (batch 2)	1300	24	V117	3.77	qz + co
			V110	3.77	qz
PTFE-Talc-Pyrex-C-MgO (¾")	1300	24	V116	3.87	co
			V112	1.46	an + gh + crn
			V106	1.55	kush + an + gh + crn
NaCl cell experiments (LMV laboratory)					
PTFE-NaCl-Pyrex-C-MgO	900	9	PC/2014/11	3.21	qz
			PC/2014/14	3.30	qz + co
		24	Cal-LMV-03	3.11	qz
			Cal-LMV-01	3.21	qz + co
			Cal-LMV-02	3.41	qz
1300	Cal-LMV-04	3.46	qz + co		
BaCO₃ cell experiments (ULG laboratory)					
Pb-BaCO ₃ -C-MgO	800	24	A070	2.35	ab
			A058	2.40	jd + qz
			A073	2.80	ab
			A060	2.90	jd + qz

Clermont-Ferrand, France) for NaCl cell assemblies, and the University of Liège (ULiège; Belgium) for BaCO₃ cell assemblies. The talc-based piston-cylinder assembly (Ceramic Substrates and Components Ltd.) comprised a talc cell wrapped in an outer PTFE film, a Pyrex cylinder, a graphite furnace, and inner MgO spacers (Fig. 1a). The current was conducted to the graphite heater through a stainless-steel plug, which was electrically isolated with a pyrophyllite sleeve. The use of PTFE film instead of conventional Pb foil for the outer sleeve allows covering the base plug, which tends to drastically decrease the force required to extract the assembly at the end of an experiment from ~35 to ~5 kN, which in turn better preserves the carbide core from fracturing. PTFE is also a more environment- and health-friendly alternative compared to Pb and/or Mo-based lubricants. The NaCl cell assembly was similar to the talc cell assembly (Fig. 1b). At ULiège, the assembly is composed of a BaCO₃ cell wrapped in Pb foil, a graphite furnace, and MgO spacers (Fig. 1c). Additional details on assembly materials and dimensions are provided in Online Materials¹ OM1. Calibrated W₇₄Re₂₆-W₉₅Re₅ thermocouples protected by a 4-bore alumina sleeve were used to control temperature to within 1 °C of the setpoint. Temperature gradients throughout the capsule are estimated to be about 20 °C (Sorbadere et al. 2013). The hot spot is located in the center of the capsule, implying a thermal gradient of about 10 °C/mm. To preclude perforation of the capsule, an Al₂O₃/MgO cap 0.6 mm thick was inserted between the capsule and the thermocouple. During experiments, the pressure was first increased to about 0.7 GPa (½" assemblies) or 0.3 GPa (¾" assemblies) at room temperature. In all experiments, the temperature was then increased to 650 °C at 50 °C/min, held for 15 min to reach the target pressure, and then heated to the target temperature.

Several series of experiments were conducted to understand the role of assembly

parts on friction in talc cells (Table 1). In experiments V32 and V42, we replaced the MgO sleeve with an alumina sleeve, a method used to preserve the experimental charge because alumina sleeves are harder and denser than MgO sleeves. We also investigated the role of the density of MgO spacers by replacing the conventional spacers (2.0 g/cm³) with higher density spacers (2.8 g/cm³) in experiments V109 and V117. In experiments V110 and V116, we tested a last batch of assemblies (batch 2) with thicker talc cells (1.35 mm instead of 0.75 mm) and thinner Pyrex glass cylinders (1 mm instead of 1.6 mm; see details in Online Materials¹ OM1). Finally, in experiments V39 and V107, Pb foil was used instead of the PTFE film.

For quartz-coesite transition experiments, Au₅₀Pd₅₀ capsules (4 mm high) were used to limit water loss (Gaetani and Grove 1998; Kägi et al. 2005). For the kushiroite reaction, Pt capsules (4 mm high) were employed. For the albite = jadeite + quartz reaction, graphite containers (3.75 mm high) were used. At the end of each experiment, the capsule was cut in half longitudinally using a wire saw, mounted in epoxy, and polished to 0.25 µm on nylon pads with diamond pastes.

Analytical techniques

Experimental run products of the kushiroite reaction were analyzed with a JEOL JSM-6510 scanning electron microscope (SEM) at CRPG using an accelerating voltage of 15 kV, a beam current of 10 nA, and a spot size of ~1 µm. Backscattered electron images of experimental run products of the albite = jadeite + quartz reaction were acquired on the QEMSCAN FEI Quanta 650 F at RWTH Aachen (Germany).

Raman spectroscopy was used to determine the nature of silica polymorphs in all experiments. Raman spectra were acquired using a LabRAM HR spectrometer (Horiba Jobin Yvon) at GeoRessources (Nancy, France). The spectrometer is equipped with a 600 gr/mm grating and an edge filter. The confocal hole aperture is 500 µm and the slit aperture is 100 µm. The excitation beam (wavelength, 514.53 nm; power, 200 mW) was produced by a Stabilite 2017 Ar+ laser (Spectra Physics, Newport Corporation) and focused on the sample using a 50× objective. The acquisition time (2 s) and the number of accumulated spectra (20) were chosen to optimize the signal-to-noise ratio. All spectra were recorded over Raman shifts of 150–1400 cm⁻¹. Quartz and coesite were identified by their respective peaks at 470 and 520 cm⁻¹.

EXPERIMENTAL TEXTURES AND PRODUCTS

Experimental conditions and results of quartz-coesite, kushiroite, and albite reactions are reported in Table 1, and calculated frictions are reported in Table 2. Pictures of typical run products are provided in Online Materials¹ OM2. In quartz-coesite experiments, coesite was always observed with variable amounts of quartz. Coesite mainly occurs as straight veins in the euhedral quartz matrix, with no preferential orientation. In kushiroite reaction experiments, grain growth was observed in all runs, with anhedral grains ranging from 20 to 50 µm. Kushiroite mainly appears near corundum and anorthite, replacing gehlenite. In runs in which kushiroite was observed (V106 and V113), anorthite, gehlenite and corundum were still present after 24 h.

Non-negligible amounts of the low-pressure phases in the three investigated reactions (i.e., quartz, anorthite + gehlenite + corundum, and albite) persisted after the transition reaction occurred. This result is consistent with Hays (1966a, 1966b), who observed low-pressure phases persisting above the transition to kushiroite, indicating a continuous reaction over a pressure interval of a few kilobars. In contrast to our experiments with a transitional pressure interval of ~0.1 GPa, Bose and Ganguly (1995) found only either quartz or coesite in their experiments with a pressure interval of ~0.05 GPa. We could not estimate modal abundances from our experimental textures due to significant sample losses during cold decompression.

Roles of intensive parameters on friction

In the following sections, experiments from Bose and Ganguly (1995) were chosen as the reference pressure (P_{eff}) for the quartz-coesite system. They showed that after 30 h, the friction

TABLE 2. Friction values

Size (")	T (°C)	Duration (h)	P _{app}	ΔP _{app}	P _{eff}	ΔP _{eff}	F (%)	ΔF		
Talc cell (CRPG laboratory)										
Friction as a function of temperature										
½	900	24	3.72	0.10	3.00	0.05	24.1	3.2		
	1000 ^a		3.75	0.10	3.07	0.06	22.0	3.2		
	1100 ^a		3.77	0.10	3.14	0.06	20.1	3.2		
	1200 ^a		3.79	0.10	3.21	0.06	18.3	3.2		
	1300		3.82	0.10	3.28	0.07	16.6	3.2		
Friction as a function of duration										
½	900	2.7	4.01	0.10	3.00	0.05	33.8	3.4		
		4.5	3.92	0.10	3.00	0.05	30.7	3.3		
		6	3.82	0.10	3.00	0.05	27.3	3.3		
		15	3.82	0.10	3.00	0.05	27.3	3.3		
		24	3.72	0.10	3.00	0.05	24.1	3.2		
	1300	6	3.63	0.10	3.00	0.05	20.9	3.2		
		24	3.82	0.10	3.28	0.07	16.6	3.2		
		24	3.82	0.10	3.28	0.07	16.6	3.2		
		Friction as a function of pressure and assembly size								
		½	1300	24	1.50	0.10	1.30	0.03	15.1	4.3
¾			1.51	0.06	1.30	0.03	15.9	4.4		
Friction as a function of assembly parts										
½	900	6	3.82	0.10	3.00	0.05	27.3	3.3 ^b		
			>3.97		3.00	0.05	>32.2			
		24	3.82	0.10	3.00	0.05	27.3	3.3		
			3.72	0.10	3.00	0.05	24.1	3.2 ^c		
			3.82	0.10	3.28	0.07	16.6	3.2		
	1300	24	3.82	0.10	3.28	0.07	16.6	3.2		
			3.82	0.10	3.28	0.07	16.6	3.2		
	NaCl cell (LMV laboratory)									
	½	900	9	3.25	0.10	3.00	0.05	8.4	3.1	
	½	24		3.16	0.10	3.00	0.05	5.4	3.0	
½	1300	24	3.38	0.10	3.28	0.07	4.8	3.0		
BaCO₃ cell (ULG laboratory)										
½	800	24	2.38	0.10	2.16	0.05	10.2	4.3		
½	1000		2.85	0.10	2.69	0.05	6.1	3.4		

Notes: P_{app} and P_{eff} represent the pressure (GPa) applied by the press and the pressure on the sample, respectively. F represents the friction in %. ΔP_{app}, ΔP_{eff} and ΔF are 2σ standard deviations of applied pressure, sample pressure and friction, respectively. **{AU: check corrected footnotes}**

^a Values from the linear fit in Figure 2.

^b Standard calibration (900 °C, 6 h), alumina sleeve, Pb foil.

^c MgO spacers density, standard calibration (1300, 24 h), assembly batch 2.

is null in their CsCl-graphite-NaCl assemblies, based on crossed calibrations using high-pressure vessel experiments. In the quartz-coesite system, our experiments at both 900 and 1300 °C allow us to investigate the role of temperature on friction (Fig. 2). In 24 h experiments in talc cells, the quartz-coesite transition occurred between applied pressures of 3.68 and 3.77 GPa at 900 °C and between 3.77 and 3.87 GPa at 1300 °C. In contrast, Bose and Ganguly (1995) observed the transition at 3.00 and 3.28 GPa at 900 and 1300 °C, respectively, in CsCl cells. Taking their values as the effective pressure on the sample, our results indicate friction values (F) for talc cells of 24.1% at 900 °C and 16.5% at 1300 °C (Table 1). A linear fit to these values (Fig. 2) shows that friction decreases by about 2% when temperature increases by 100 °C in 24 h experiments. In NaCl cells, the quartz-coesite transition occurred between 3.11 and 3.21 GPa at 900 °C (F = 5.4 ± 3.0%) and between 3.41 and 3.46 GPa at 1300 °C (F = 4.8 ± 3.0%), indicating no friction evolution within error. In BaCO₃ cells, the albite = jadeite + quartz reaction occurred between 2.35 and 2.4 GPa at 800 °C and between 2.8 and 2.9 GPa at 1000 °C. Compared to 2.16 and 2.69 GPa at 800 and 1000 °C obtained in NaCl assemblies used by Holland (1980), F = 10.2 ± 4.3 and 6.1 ± 3.4%, respectively, indicating no friction evolution in this range of temperature.

Our experiments on the kushiroite = anorthite + gehlenite + corundum reaction were conducted at 1300 °C to understand the role of pressure on the friction value in talc cells. In ½" as-

semblies, this transition occurred between applied pressures of 1.45 and 1.54 GPa in 24 h experiments. According to the equation of Hays (1966a) based upon experiments using talc assemblies, the reference pressure for this transition is 1.3 GPa, resulting in a friction value of 15.1 ± 4.3%.

This value is identical, within error, to the friction value obtained from the quartz-coesite transition at the same temperature (16.6 ± 3.2%), implying that the friction value is independent of confining pressure at identical temperatures and experimental durations. This result is partially consistent with McDade et al. (2002), who argued that the friction correction is independent of pressure and temperature at 1000–1600 °C and 1.5–3.2 GPa in BaCO₃ cells. However, our results show that friction is temperature dependent in talc cells, decreasing by 2% over a temperature increase of 100 °C between 900 and 1300 °C. This difference can be explained by the slower stress accommodation of talc compared to NaCl and BaCO₃ cells due to their higher strength. It is also possible that friction becomes more strongly temperature dependent at low temperatures (<1000 °C). We speculate that stress accommodation is fast enough above 1000 °C to be undetectable under these experimental conditions.

FRICTION EVOLUTION

We performed experiments on the quartz-coesite reaction at 900 °C and over various run durations between 2.7 and 48 h to investigate the evolution of friction during a single run (Fig. 3). In talc cells, the transition occurred at progressively lower applied pressures with increasing run duration (Table 1); friction thus evolved from 33.8 ± 3.4% after 2.7 h to 30.7 ± 3.3% after 4.5 h, 27.3 ± 3.3% after 6 h, 24.1 ± 3.2% after 24 h, and to 20.9 ± 3.2% after 48 h (Fig. 3). Whereas a linear fit to the data for longer durations (≥6 h) indicates that friction decreases by ~2% every 10 h, a power-law fit better describes the full data set. This may explain the ubiquitous presence of quartz in all experiments conducted in talc cells via the early crystallization of quartz from the amorphous SiO₂ starting material at low effective pressures and its subsequent metastability. In contrast, in

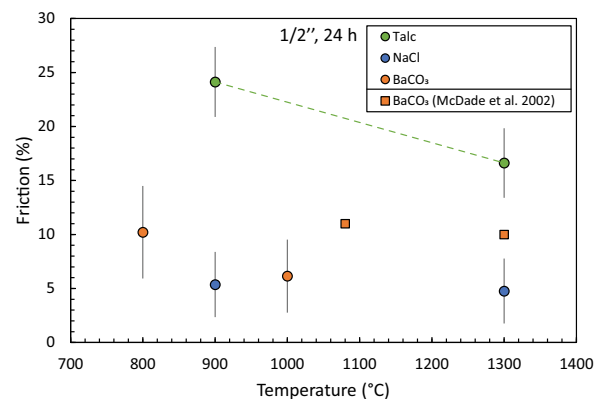


FIGURE 2. Friction evolution as a function of temperature for 24 h experiments in ½" cell assemblies. The duration of experiments from McDade et al. (2002) was 22 h. Linear fits describe the change in friction as a function of temperature (°C): $F = -0.019 T + 40.991$ for talc cells; $F = -0.006 T + 10.370$ for NaCl cells; $F = -0.020 T + 26.463$ for BaCO₃ cells. (Color online.)

talc cell assemblies at 1300 °C, friction did not evolve between 6 and 24 h duration experiments, indicating that a frictional steady state was achieved within <6 h.

In NaCl cells at 900 °C, the apparent pressure of the quartz-coesite transition decreased from between 3.21 and 3.30 GPa after 9 h ($F = 8.4 \pm 3.1\%$) to between 3.11 and 3.21 GPa after 24 h ($F = 5.4 \pm 3.0\%$), indicating no friction evolution within error, contrary to what is reported in talc cells over longer duration experiments. The friction decrease in talc cells is, however, consistent with the data of Bose and Ganguly (1995) in CsCl cell assemblies (both $\frac{1}{2}$ " and $\frac{3}{4}$ "), which show that friction decreased from 6.6% after 2 h to negligible after 35 h, or by $\sim 2\%$ every 10 h (Fig. 3).

THERMOMECHANICAL PROPERTIES OF ASSEMBLY COMPONENTS

Alumina sleeve

Experiments V32 and V42 (Table 1) were performed at 900 °C and applied pressures of 3.87 and 3.97 GPa, respectively, with an alumina sleeve around the capsule instead of a conventional MgO sleeve. In both experiments, only quartz was observed, and the pressure limitation of the carbide core precludes experiments at pressures above ~ 4 GPa. Thus, the minimum friction observed using alumina sleeves is $>32.3\%$ (Fig. 4). However, the main advantage of alumina sleeves is that the sample capsule conserves its cylindrical shape, preventing excessive deformation during quenching and cold depressurization, a net advantage when working with single crystals. McDade et al. (2002) obtained a friction value of 3.6% in NaCl cells by using an alumina sleeve and Alsimag (Al_2O_3) plugs to fill the tapered graphite furnace. Given the high density and hardness of Alsimag, these plugs are better pressure transmitters than the crushable MgO used in our assembly setup. The combination

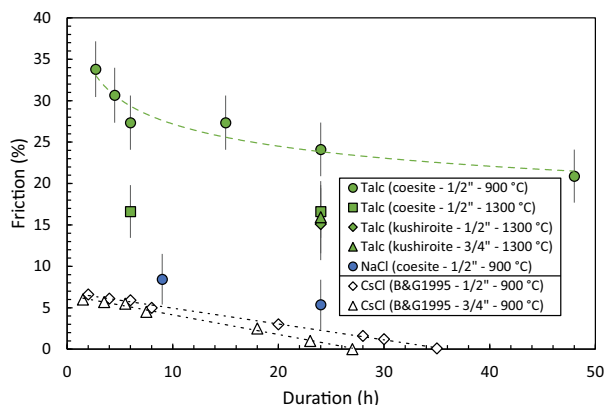


FIGURE 3. Evolution of the friction value as a function of experimental duration. B&G1995 data correspond to the study of Bose and Ganguly (1995), performed at 900 °C with CsCl cells. At 900 °C, the friction value decreases of about 2% every 10 h for experimental durations >6 h. At 1300 °C, no friction decrease is observed between 6 and 24 h. Lower pressure (kushiroite experiments) and sizing of assemblies ($\frac{1}{2}$ " and $\frac{3}{4}$ ") do not seem to have a critical influence on the friction value and its evolution. The slope of the friction decrease observed in NaCl cell assemblies is comparable with the slope observed in Bose and Ganguly (1995). (Color online.)

of an alumina sleeve with MgO spacers thus greatly decreases pressure transmission and should only be used to limit fracturing of the experimental product in cases requiring particular care for the capsule.

Density of MgO spacers

Experiments V109 and V117 made with talc-Pyrex assemblages were conducted 900 °C and applied pressures of 3.68 and 3.77 GPa, respectively, with higher density MgO spacers (2.8 vs. the usual 2.0 g/cm^3). Both coesite and quartz were observed in V117, positioning the apparent quartz-coesite transition at 3.72 GPa. The friction obtained using these MgO spacers ($24.1 \pm 3.2\%$; Fig. 4) is identical within error to that obtained using the standard assembly under the same conditions ($27.3 \pm 3.3\%$). This result demonstrates that the densities of assembly components do not significantly affect the effective sample pressure. This, however, further highlights that the Alsimag spacers used by McDade et al. (2002) are a good pressure medium, as they achieved a friction value of only 3.6%.

Assembly size ($\frac{1}{2}$ " vs. $\frac{3}{4}$ ")

Kushiroite reaction experiments were performed in both $\frac{1}{2}$ " and $\frac{3}{4}$ " assemblies at 1300 °C for 24 h (Table 1). The calculated friction values are identical within errors, i.e., $15.1 \pm 4.3\%$ and $15.9 \pm 4.4\%$ for the $\frac{1}{2}$ " and $\frac{3}{4}$ " assemblies, respectively (Fig. 3), suggesting that the size of assembly parts has a negligible effect on friction compared to assembly materials (and their associated thermomechanical properties).

Experiments V110 and V116 were performed using assemblies with thicker talc cells and thinner Pyrex sleeves (batch 2,

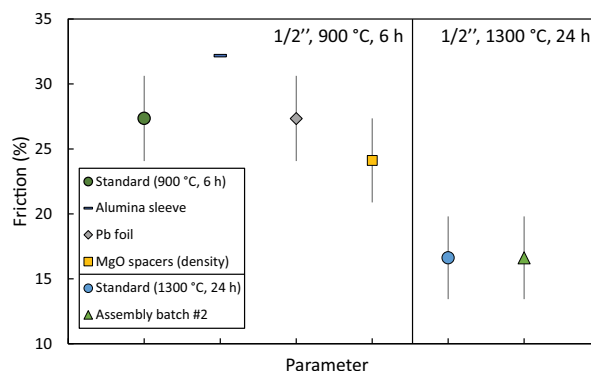


FIGURE 4. Variations in friction with changes in assembly size and components, relative to experiments performed using the assembly shown in Figure 1a (talc cell, MgO sleeve, PTFE film, 2.0 g/cm^3 MgO spacers) at 900 °C for 6 h and 1300 °C for 24 h ($\frac{1}{2}$ " assembly). "Alumina sleeve" indicates that the regular MgO sleeve around the capsule was replaced by an alumina sleeve. "Pb foil" indicates that the cell was wrapped in standard Pb foil instead of the PTFE film used throughout this study. "MgO spacers" indicates that higher density MgO spacers (2.8 g/cm^3) were used within the graphite furnace (including the sleeve around the capsule). "Assembly batch 2" indicates the assembly shown in Figure 1d, using a thicker talc cell and thinner Pyrex sleeve than those in the regular assembly (Fig. 1a). See Online Materials¹ OM1 for details. Note that the friction reported for "Alumina sleeve" is a minimum value because coesite was not observed in this series of experiments due to the pressure limitations of the press and carbide core. (Color online.)

see Online Materials¹ OM1 for size details). The friction determined for this assembly ($16.6 \pm 3.2\%$) is identical to that for the standard assembly ($16.6 \pm 3.2\%$) at 1300 °C for 24 h (Fig. 4), further demonstrating that assembly materials have a stronger influence on friction than their sizes.

Pb foil vs. PTFE foil

Experiments V107 and V39 were performed at 900 °C and applied pressures of 3.77 and 3.87 GPa, respectively, using talc cells and standard Pb foil instead of the PTFE film used elsewhere in this study (excluding ULiège experiments, Table 1). Coesite was only observed in experiment V39, implying that friction ($27.3 \pm 3.3\%$) was identical within errors to that using a PTFE-wrapped assembly ($27.3 \pm 3.3\%$; Fig. 4). As previously mentioned, the key advantage of using PTFE film instead of Pb foil is the preservation of the carbide core of the pressure plates. The steel base plug, surrounded by pyrophyllite, is frequently stuck in the carbide core when removing the assembly, especially after experiments run at pressures above 2.0 GPa. The use of Pb foil precludes covering the base plug to avoid melting the Pb at the contact point with the thermocouple plate, which can electrically short-circuit the experiment. Therefore, the use of PTFE film is recommended to limit the development of fractures within the carbide core, which will both extend the life of the carbide core and enhance pressure reproducibility over that lifetime.

IMPLICATIONS FOR PRESSURE REPRODUCIBILITY DURING PISTON-CYLINDER EXPERIMENTS

In this section, we provide general recommendations to improve pressure and temperature reproducibility during piston-cylinder experiments. These recommendations can be adapted, depending on the capsule material and the scientific purpose, e.g., for oxygen-fugacity buffering or volatile-bearing experiments. These recommendations are depicted in *P-T-t* space in the form of preferential practices in Figure 5.

At temperatures above 1600 °C, use of the NaCl cell is difficult to use because of the melting point of NaCl. Consequently, only talc or BaCO₃ cells are recommended for very high-temperature experiments. Talc cells require a very high-frictional correction factor, regardless of *P-T-t* conditions (Table 2), and should thus be avoided in very high-pressure experiments (>3 GPa) because the carbide core could suffer from excessive applied pressure. To generate pressures above 3 GPa, we thus recommend using NaCl cells below 1600 °C because they exhibit the lowest friction corrections (Figs. 2 and 3) or BaCO₃ cells at higher temperatures. For experimental durations of a few to ~48 h and pressures below 3 GPa, talc cells are as competent as NaCl and BaCO₃ cells because they all show similar frictional evolution with time. However, for short experiments (≤ 6 h), we recommend using NaCl or BaCO₃ cells as the decrease in friction value with time is comparable and because the friction in talc cells is much greater (following a power law) in shorter duration experiments (Fig. 3). As described in the “Friction evolution” section, this could lead to unexpectedly low effective pressures, promoting the crystallization and metastability of low-pressure phases. For experiments longer than 48 h (and below 1600 °C), we recommend using NaCl cells: whereas friction becomes negligible in CsCl cells after ~30 h (Bose and Ganguly 1995), the behavior

of talc cells in experiments longer than 48 h remains unclear. Therefore, for temperatures above 1600 °C and durations longer than 48 h, we recommend using BaCO₃ cells because of their lower frictional correction compared to talc cells.

Whereas the size of assembly parts has a negligible effect on friction, different sleeve and spacer materials (here, MgO vs. alumina) can significantly change the friction factors. Friction decreases by about 2% per 100 °C increase between 900 and 1300 °C at a given duration. Experimental duration also has a strong effect on the friction value at low temperatures, with friction decreasing by about 0.2% per hour at 900 °C; this effect is not observed at 1300 °C. In contrast to CsCl cells, in which friction becomes negligible after around 30–35 h, the friction in talc cells, although decreasing with increasing run duration, is never negligible, especially at low temperatures. Therefore, special care must be taken for low-temperature experiments (roughly <1000 °C) using talc cells, especially at short durations, because the slow stress accommodation of the assembly results in slow frictional evolution during the first several hours. Other key parameters not studied here may also critically affect friction and its evolution. In particular, the quality of the carbide core and the development of fractures are expected to influence the friction generated on the assembly. A more systematic calibration procedure, including the status of the core, is thus necessary to provide a complete model of friction and its evolution during an experiment. This is fundamental for phase relationship experiments or the quest for low-degree melts from mantle lithologies,

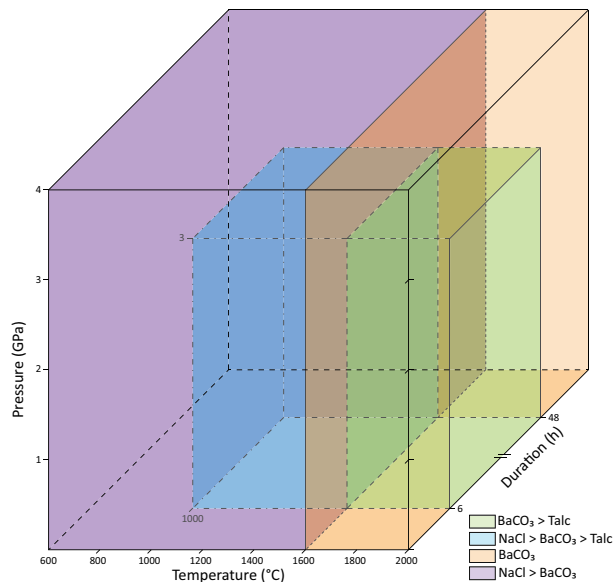


FIGURE 5. Recommended cell types for piston-cylinder experiments as a function of pressure, temperature, and experimental duration. Note that the duration axis is not linear. These general recommendations limit the uncertainty on pressure and its evolution during HP-HT experiments, except for avoiding NaCl cells in experiments at very high temperature (>1600 °C) because the NaCl cell no longer acts as an electrical insulator. See discussion in the “Implications for pressure reproducibility during piston-cylinder experiments” section for further details. Cell types are recommended as “best > intermediate > worst” according to the color scale at bottom right. (Color online.)

for which accurate and reproducible P - T conditions are essential. Furthermore, a dynamic model for friction correction should be developed to ensure pressure reproducibility among HP-HT petrological studies.

ACKNOWLEDGMENTS

The authors thank Fred Davis and an anonymous reviewer for their constructive reviews, as well as Kate Kiseeva and Don Baker for their editorial handling. We thank M.-C. Caumon for technical assistance with Raman analyses and P. Baillot for his technical expertise in the laboratory. We also thank K. Koga and F. Faure for fruitful discussions. P.C. thanks C. McCammon for fundamental advice when building the piston-cylinder laboratory at CRPG.

FUNDING

This study was mainly financed by l'Agence Nationale de la Recherche through grant ANR INDIGO (ANR-14-CE33-0011). C. Dalou and E. Füre were supported by the European Research Council (ERC) under the European Union's Horizon 2020 research and innovation program (grant agreement no. 715028). B. Charlier is a Research Associate of the Belgian Fund for Scientific Research-FNRS. A. Triantafyllou was supported by the FRS-FNRS for the PROBARC project (Grant CR no. 1. B. 414.20F). This is CRPG contribution no. 2747.

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MANUSCRIPT RECEIVED JANUARY 8, 2021

MANUSCRIPT ACCEPTED JULY 29, 2021

MANUSCRIPT HANDLED BY KATE KISEEVA

Endnote:

¹Deposit item AM-22-87958, Online Materials. Deposit items are free to all readers and found on the MSA website, via the specific issue's Table of Contents (go to http://www.minsocam.org/MSA/AmMin/TOC/2022/Aug2022_data/Aug2022_data.html).