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# Gases as Precursory Signals: Experimental Simulations, New Concepts and Models of Magma Degassing

M. Pichavant, N. Le Gall and B. Scaillet

## Abstract

Volatile release during magma ascent in volcanic conduits (magma degassing) forms the basis for using volcanic gases as precursory signals. Recent high temperature high pressure experimental simulations have yielded results that challenge key assumptions related to magma degassing and are important for the interpretation of glass inclusion and gas data and for using volcanic gas as precursory signals. The experimental data show that, for ascent rates expected in natural systems, pure H<sub>2</sub>O basaltic melts will evolve mostly close to equilibrium when decompressed from 200 to 25 MPa. In the same way, degassing of H<sub>2</sub>O–S species evolves at near equilibrium, although this conclusion is limited by the number of S solubility data available for basaltic melts. However, degassing of CO<sub>2</sub> is anomalous in all studies, whether performed on basaltic or rhyolitic melts. CO<sub>2</sub> stays concentrated in the melt at levels far exceeding solubilities. The anomalous behaviour of CO<sub>2</sub>, when associated with near equilibrium H<sub>2</sub>O losses, yields post-decompression glasses with CO<sub>2</sub> concentrations systematically higher than equilibrium degassing curves. Therefore, there is strong

experimental support for disequilibrium degassing during ascent of CO<sub>2</sub>-bearing magmas. The existence of volatile concentration gradients around nucleated gas bubbles suggests that degassing is controlled by the respective mobilities (diffusivities) of volatiles within the melt. The recently formulated diffusive fractionation model reproduces the main characteristics, especially the volatile concentrations, of experimental glasses. The model also shows that the gas phase is more H<sub>2</sub>O-rich than expected at equilibrium because CO<sub>2</sub> transfer toward the gas phase is hampered by its retention within the melt. However, only integrated gas compositions are calculated. Similarly, only bulk experimental fluid compositions are determined in recent experiments. Thus, constraints on the local gas phase are becoming necessary for the application to volcanoes. This stresses the need for the direct analysis of gas bubbles nucleated in decompression experiments. Pre-eruptive changes in volcanic CO<sub>2</sub>/SO<sub>2</sub> and H<sub>2</sub>O/CO<sub>2</sub> gas ratios are interpreted to reflect different pressures of gas-melt segregation in the conduit, an approach that assume gas-melt equilibrium. However, if disequilibrium magma degassing is accepted, the use of volatile saturation codes is no longer possible and caution must be exercised with the application of local equilibrium to volcanic gases. Future developments in the interpretation of gas data require progress

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from both sides, experimental and volcanological. One priority is to reduce the gap in scales between experiments and gas measurements.

### Keywords

Magma ascent • Decompression experiments  
Disequilibrium degassing • Volatile diffusion  
Gas compositions

### Extended Abstract

Volcanic gases are one of the main tools to monitor changes in the activity of volcanoes and forecast their eruption. Magma ascent toward the surface is associated with the exsolution of volatiles initially dissolved in the melt, a process designated as “magma degassing”. Classically, the interpretation of volcanic gases relies on the assumption that degassing takes place at equilibrium. However, several observations (CO<sub>2</sub> contents of basaltic seafloor glasses, H<sub>2</sub>O and CO<sub>2</sub> concentrations in glass inclusions, explosive basaltic volcanism) do not fit easily in such a model. Recently, decompression, ascent and degassing of magmas in volcanic conduits have been simulated by high temperature high pressure experiments. Results from these simulations stress the need to critically reconsider the whole mechanism of degassing in basaltic but also rhyolitic magmas. The new experimental data show that, at the decompression rates tested, pure H<sub>2</sub>O basaltic melts will evolve mostly close to equilibrium when decompressed from 200 to 25 MPa. In the same way, degassing of H<sub>2</sub>O–S species evolves at near equilibrium, although this conclusion is limited by the number of S solubility data available for basaltic melts. Degassing of CO<sub>2</sub> is anomalous in all studies, whether performed on basaltic or rhyolitic melts. CO<sub>2</sub> stays concentrated in the melt at levels far exceeding solubilities. The anomalous behaviour of CO<sub>2</sub>, when associated with near equilibrium H<sub>2</sub>O losses, yields post-decompression glasses with CO<sub>2</sub> concentrations systematically higher than equilibrium degassing curves. Therefore, there is strong experimental support for disequilibrium

degassing during ascent of CO<sub>2</sub>-bearing magmas. The existence of volatile concentration gradients around nucleated gas bubbles suggests that degassing is controlled by the respective mobilities (diffusivities) of volatiles within the melt. The contrasted diffusivities of dissolved volatile species (in particular H<sub>2</sub>O and CO<sub>2</sub>) selectively limit their transfer toward the gas phase for timescales typical for magma ascent. The diffusive fractionation model recently formulated reproduces the main characteristics, especially the volatile concentrations, of experimental glasses. It provides a framework to interpret the new experimental observations and the systematic deviations from equilibrium observed in CO<sub>2</sub>-bearing systems, although coupling between volatile diffusion and vesiculation requires a more elaborate treatment. The model also shows that the gas phase is more H<sub>2</sub>O-rich than expected at equilibrium because CO<sub>2</sub> transfer toward the gas phase is hampered by its retention within the melt. However, only integrated gas compositions are calculated. In the same way, only bulk experimental fluid compositions are determined in recent experiments. Since the gas phase is essential for the application to volcanoes, constraints on the local gas phase are becoming necessary. Compositions of gas bubbles in decompression experiments must be linked not only with pressure but also with volatile concentrations of local melts, specific degassing textures and mechanisms. As a way in this direction, local gas-melt equilibrium assumes that chemical equilibrium persists at the local scale, despite evidence for disequilibrium at larger scales. However, there are alternative ways to constrain the composition of nucleated gas bubbles, thus stressing the need for their direct analysis in decompression experiments. Pre-eruptive changes in CO<sub>2</sub>/SO<sub>2</sub> and, in some cases, H<sub>2</sub>O/CO<sub>2</sub> gas ratios observed at several basaltic volcanoes are generally interpreted to reflect different pressures of gas-melt segregation in the conduit, an approach that assume gas-melt equilibrium. However, if disequilibrium magma degassing is accepted, volatile saturation codes can no longer be directly used. Caution also must be exercised with the application of local gas-melt equilibrium to volcanic gases which are probably

closer to integrated rather than to local compositions. Future developments in the interpretation of gas data require progress from both sides, experimental and volcanological. One priority is to reduce the gap in scales between experiments and gas measurements to refine interpretations of gas compositions as unrest signals.

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## 1 Magma Degassing and Volcanic Gases as Precursory Signals

Volcanic gases are one of the main tools used to monitor changes in the activity of volcanoes and forecast their eruption. This approach is rooted in the strong pressure dependence of the solubility of volatiles (mainly H<sub>2</sub>O, CO<sub>2</sub>, SO<sub>2</sub>, H<sub>2</sub>S, Cl) in silicate melts. Accordingly, magma ascent toward the surface is associated with the exsolution of volatiles initially dissolved in the melt, a process designated as “magma degassing”. The different volatiles have contrasted solubilities in silicate melts and, therefore, are expected to react differently to decompression. This forms the basis for using volcanic gas ratios to infer magma ascent and depth of gas segregation in volcanic conduits. For example, the sudden increase of gas CO<sub>2</sub>/SO<sub>2</sub> ratio has been used as an indication for deep magma recharge at Stromboli (Aiuppa et al. 2010). At Soufriere Hills volcano (Montserrat), a correlation has been noted between gas HCl/SO<sub>2</sub> and the level of shallow activity as marked by the rate of lava extrusion and dome growth (Christopher et al. 2010; Edmonds et al. 2010).

Classically, the interpretation of volcanic gases relies on the assumption that degassing takes place at equilibrium. In the case of basaltic magmas, this assumption is supported by the high temperatures, low viscosities and high volatile diffusivities (Sparks et al. 1994). Vesiculation (i.e., the combined processes of bubble nucleation, growth and coalescence) is thought to be relatively easy in basaltic melts and degassing of basaltic magma is classically viewed as an equilibrium process. However, several observations do not fit easily in

such a model. They include (1) the existence of basaltic seafloor glasses often supersaturated in CO<sub>2</sub> (e.g., Aubaud et al. 2004), (2) the occurrence of glass inclusions with H<sub>2</sub>O and CO<sub>2</sub> concentrations inconsistent with closed system equilibrium degassing (e.g., Metrich et al. 2010) and (3) the occurrence of explosive basaltic volcanism (e.g., Head and Wilson 2003) which implies sudden rather than gradual release of volatiles.

Recently, decompression and ascent of basaltic magmas in volcanic conduits has been simulated by high temperature high pressure petrological experiments. These simulations stress the need to critically reconsider the whole mechanism of degassing in basaltic but also more silicic magmas. In particular, the assumption of equilibrium degassing is now becoming increasingly challenged. This has major implications for the interpretation of glass inclusion and gas data and, more generally, for the use of volcanic gas as precursory signals. In this Chapter, first, the recent experimental simulations are reviewed. We show that they all demonstrate an anomalous behaviour for CO<sub>2</sub> which tends to stay dissolved within the melt at concentrations too high for equilibrium. Second, the diffusive fractionation model which has been proposed to account for the new experimental observations is described and critically discussed. Finally, the implications of disequilibrium degassing for experimental fluid compositions and the interpretation of volcanic gas data as precursory signals are explored.

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## 2 Experimental Simulations

### 2.1 Basaltic Systems

Following early work on systems with only pure CO<sub>2</sub> (Lensky et al. 2006), decompression experiments on hydrous basaltic melts have been carried out recently by Pichavant et al. (2013) at 1150–1180 °C, for initial pressures of 200–250 MPa, final pressures of 100, 50 and 25 MPa and for decompression rates between ~1.5 down to 0.25 m/s. Melts from Stromboli, pre-synthesized to incorporate dissolved H<sub>2</sub>O (2.7–3.8 wt%) and CO<sub>2</sub> (600–1300 ppm), were

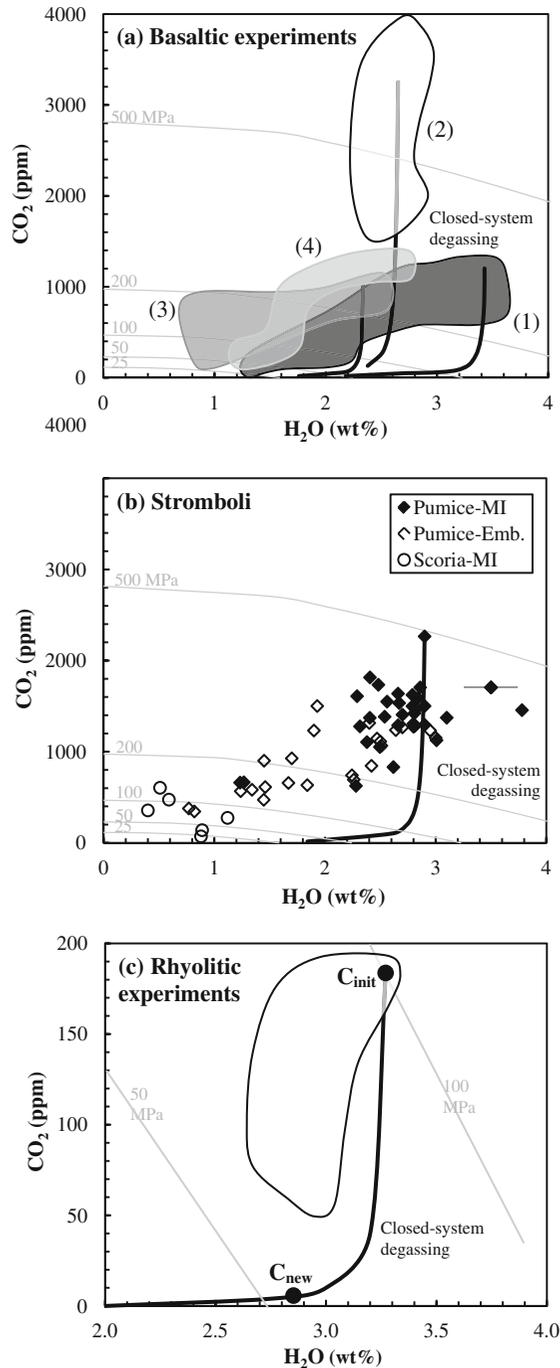
used as starting materials. The experiments were of continuous decompression type, and both constant (one ramp) and variable (two ramps) decompression rates were imposed. Final melt H<sub>2</sub>O concentrations were homogeneous and always close to equilibrium solubility values. In contrast, the rate of vesiculation was found to control the final melt CO<sub>2</sub> concentration. High vesicularity charges had glass CO<sub>2</sub> concentrations that follow theoretical equilibrium degassing paths whereas glasses from low vesicularity charges showed marked deviations from equilibrium, with CO<sub>2</sub> concentrations up to one order of magnitude higher than equilibrium solubilities (Fig. 1a). The experimental results were interpreted in light of the slower diffusivity of CO<sub>2</sub> relative to H<sub>2</sub>O in basaltic melts.

Yoshimura (2015) decompressed a natural evolved basaltic melt containing dissolved H<sub>2</sub>O and CO<sub>2</sub> at 1200 °C and between 1000 and 500 MPa. The short decompression duration of 10 min over this pressure interval simulates a very fast ascent rate (~32 m/s for a rock density of 2650 kg/m<sup>3</sup>). A vesiculated glass was produced and Fourier Transform Infrared Spectroscopy (FTIR) profiles revealed large CO<sub>2</sub> concentration gradients in the melt adjacent to gas bubbles. In contrast, the melt H<sub>2</sub>O content was almost constant throughout the sample. The glass volatile concentration data cover a near vertical trend in the H<sub>2</sub>O–CO<sub>2</sub> diagram (Fig. 1a).

Le Gall and Pichavant (2016a) extended the decompression experiments performed by Pichavant et al. (2013), using essentially the same procedures and materials. Three starting volatile compositions were investigated: series #1 (4.91 wt% H<sub>2</sub>O, no CO<sub>2</sub>), series #2 (2.41 ± 0.04 wt% H<sub>2</sub>O, 973 ± 63 ppm CO<sub>2</sub>) and series #3 (0.98 ± 0.16 wt% H<sub>2</sub>O, 872 ± 45 ppm CO<sub>2</sub>). The volatile-bearing glasses were synthesized at 1200 °C and 200 MPa, then continuously decompressed at a fast decompression rate of 3 m/s in the pressure range 150–25 MPa and then rapidly quenched. Post-decompression glasses were characterized

texturally by X-ray microtomography. Volatile equilibrium was reached or approached during decompression in all series #1 melts with just water. In contrast, disequilibrium degassing occurred systematically in series #2 and #3 melts which retained elevated CO<sub>2</sub> concentrations (Fig. 1a). In similar experiments performed on the same three glass series but at a slower decompression rate of 1.5 m/s, Le Gall and Pichavant (2016b) found that series #1 (CO<sub>2</sub>-free) melts followed equilibrium degassing until 100 MPa final pressure (P<sub>fin</sub>). But at both 60 and 50 MPa P<sub>fin</sub>, a slight H<sub>2</sub>O-supersaturation was recognized, associated with a second bubble nucleation event that occurred at 25 MPa. In comparison, in series #2 and #3 (CO<sub>2</sub>-bearing) melts, disequilibrium degassing was systematic, glasses retaining high non-equilibrium CO<sub>2</sub> concentrations (Fig. 1a).

The behavior of H<sub>2</sub>O-, CO<sub>2</sub>- and S-bearing basaltic melts during decompression was investigated by Le Gall et al. (2015a). Stromboli melts with 2.72 ± 0.02 wt% H<sub>2</sub>O, 1291 ± 85 ppm CO<sub>2</sub> and 1535 ± 369 ppm S were synthesized at 1200 °C and 200 MPa and then decompressed to final pressures (P<sub>fin</sub>) ranging from 150 to 25 MPa, followed by rapid quenching. The continuous decompressions were conducted at rates of 1.5 and 3 m/s. During decompression, S (and H<sub>2</sub>O) were lost slightly more from the melt than expected from equilibrium degassing models, whilst significant CO<sub>2</sub> was retained at elevated concentrations in the melt (Fig. 1a). It was found that the degassing trend recorded by Stromboli glass inclusions could be closely reproduced by the experiments (Fig. 1b; Le Gall et al. 2015a). For andesitic melts, Fiege et al. (2014) observed that the fluid/melt partition coefficient for sulfur increases with the decompression rate. However, the influence of decompression rate on S degassing was marked only for oxidizing conditions, corresponding to sulfate as the only S species, thus making necessary to consider the different behaviour of S<sup>2-</sup> and S<sup>6+</sup> during degassing.



**Fig. 1** H<sub>2</sub>O-CO<sub>2</sub> glass concentration diagrams for **a** basaltic decompression experiments, **b** Stromboli glass inclusions and **c** rhyolitic decompression experiments. In **a**, **b** and **c**, light grey curves are isobars labelled with pressure in MPa. In **a**, fields for post-decompression glasses are distinguished with (1) referring to Pichavant et al. (2013), (2) to Yoshimura (2015), (3) to Le Gall and Pichavant (2016a, b) and (4) to Le Gall et al. (2015a). Black curves are closed-system equilibrium degassing trajectories redrawn from the original figures (Pichavant

et al. 2013; Yoshimura 2015; Le Gall and Pichavant 2016a, b; Le Gall et al. 2015a). In **b**, the glass inclusion data are from Metrich et al. (2010). MI: glass inclusions, Emb: embayments. In **c**, the glass data field and the bold theoretical equilibrium closed-system degassing curve are redrawn from Yoshimura (2015). C<sub>init</sub> and C<sub>new</sub> are the composition of the pre-decompression melt and of the estimated post-decompression melt at the gas-melt interface, respectively (Yoshimura 2015; see also Figs. 5 and 6)

## 2.2 Rhyolitic Systems

Yoshimura (2015) decompressed a natural rhyolitic melt containing dissolved H<sub>2</sub>O and CO<sub>2</sub> from 100 to 50 MPa at 800 °C. The duration of the decompression was 5000 s corresponding to a decompression rate of 0.38 m/s (for a density of 2650 kg/m<sup>3</sup>). FTIR analysis of the vesiculated glass sample showed CO<sub>2</sub> concentration gradients in the melt away from gas bubbles. In contrast, H<sub>2</sub>O was found to be distributed homogeneously within the sample although H<sub>2</sub>O concentrations decreased significantly relative to the pre-decompression melt showing it had re-equilibrated. On the H<sub>2</sub>O–CO<sub>2</sub> diagram, the glass volatile concentrations define a near vertical array located left of the theoretical equilibrium degassing curve (Fig. 1c).

## 2.3 Summary of Experimental Evidence

In all experiments above, melt vesiculation is the result of decompression, in most cases single-step (constant decompression rate) and, more rarely, multi-step (variable decompression rates, Pichavant et al. 2013). Vesiculation leads to the generation of a gas (or fluid) phase. Volatiles partition between melt and gas, and volatile concentrations in post-decompression glasses evolve from those initially dissolved in pre-decompression glasses. The evaluation of equilibrium vs. disequilibrium degassing is performed by comparing volatile concentrations of post-decompression glasses with theoretical closed-system equilibrium degassing trajectories (Fig. 1). For pure H<sub>2</sub>O melts, this equilibrium trajectory is calculated using the experimental solubility data of Lesne et al. (2011). For CO<sub>2</sub>- and S-bearing systems, gas-melt equilibrium thermodynamic models (Newman and Lowenstern 2002; Papale et al. 2006; Burgisser et al. 2015) are used. Results show that pure H<sub>2</sub>O basaltic systems evolve close to equilibrium when decompressed from 200 to 25 MPa with ascent rates of 1.5 and 3 m/s, although small levels of H<sub>2</sub>O supersaturation are observed below 100 MPa (Le Gall and Pichavant 2016a, b). In the same way,

degassing of S species evolves at near equilibrium (Le Gall et al. 2015a) although the reference equilibrium model (Burgisser et al. 2015) is somewhat uncertain due to the limited number of S solubility data for calibration. Degassing of CO<sub>2</sub> is anomalous in all studies, whether performed on basaltic or rhyolitic melts (Pichavant et al. 2013; Yoshimura 2015; Le Gall and Pichavant 2016a, b; Le Gall et al. 2015a). CO<sub>2</sub> stays concentrated in the melt at concentrations far exceeding solubilities (Fig. 1). Except in the very fast basalt decompression experiment of Yoshimura (2015), the anomalous behaviour of CO<sub>2</sub> is associated with significant H<sub>2</sub>O losses which results in post-decompression glasses plotting systematically left of theoretical equilibrium degassing trajectories in Fig. 1. We conclude that recent experimental studies strongly support the possibility of disequilibrium degassing, i.e., that ascending melts can keep volatile concentrations (particularly CO<sub>2</sub>) significantly different from those expected from equilibrium modelling. The questions thus arise of (1) the mechanisms responsible for this disequilibrium behaviour and (2) of the consequences of disequilibrium degassing for the composition of the gas phase.

## 3 Modelling Disequilibrium Degassing

### 3.1 The Diffusive Fractionation Model

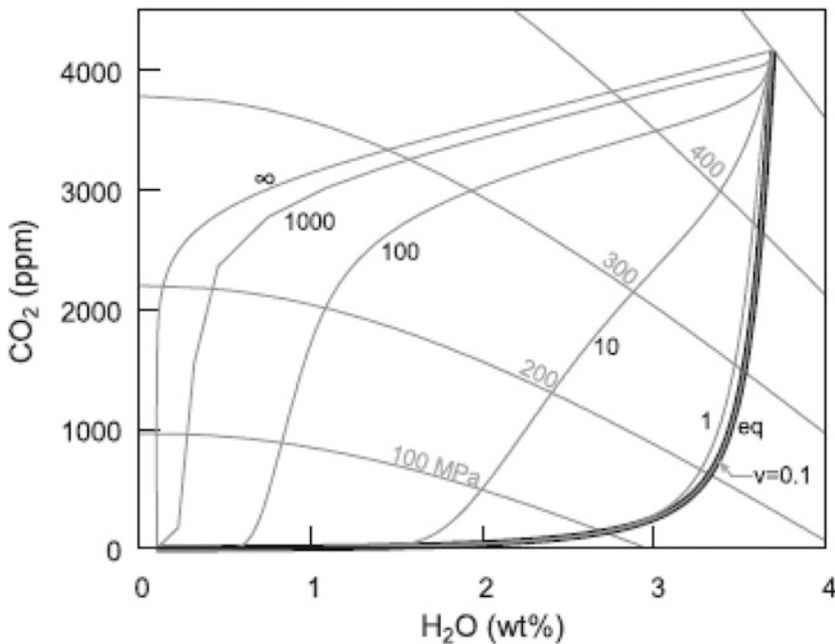
Both Pichavant et al. (2013) and Yoshimura (2015) observed decoupling between the behaviour of H<sub>2</sub>O and CO<sub>2</sub> during experimental decompression and degassing. In both studies, CO<sub>2</sub> concentration gradients were found in post-decompression glasses, either around gas bubbles or near the gas-melt interface. In contrast, no such diffusion profiles were identified for H<sub>2</sub>O, despite concentrations being lower (in most cases) than in pre-decompression glasses (Pichavant et al. 2013; Yoshimura 2015; Le Gall et al. 2015a; Le Gall and Pichavant 2016a, b).

Pichavant et al. (2013) suggested that two characteristic distances, the gas interface distance

(either the distance between two bubbles in the melt or the distance to the gas-melt interface) and the volatile diffusion distance (a function of respective diffusivities of volatiles in the melt) control the degassing process. Yoshimura (2015) quantitatively formulated a diffusive fractionation model to describe the ascent and degassing of volatile-bearing magmas. The reader is referred to this work for details about the calculations. The model is based on a diffusivity of  $\text{CO}_2$  being one log unit lower than for  $\text{H}_2\text{O}$  (e.g., Zhang and Ni 2010). Decompression trajectories computed from the model are shown on Fig. 2 for different ascent rates, from 0.1, 1, 10, 100 to  $\infty$  m/s. Although very high ascent rates (e.g., Peslier et al. 2015) are necessary for degassing trajectories to shift significantly left to the equilibrium reference curve, the modelling results qualitatively reproduce the main characteristics of experimental post-decompression glasses, i.e., the elevated  $\text{CO}_2$  glass concentrations, the significant  $\text{H}_2\text{O}$  losses and the melt concentration trends in  $\text{H}_2\text{O}$ – $\text{CO}_2$  diagrams (Fig. 1).

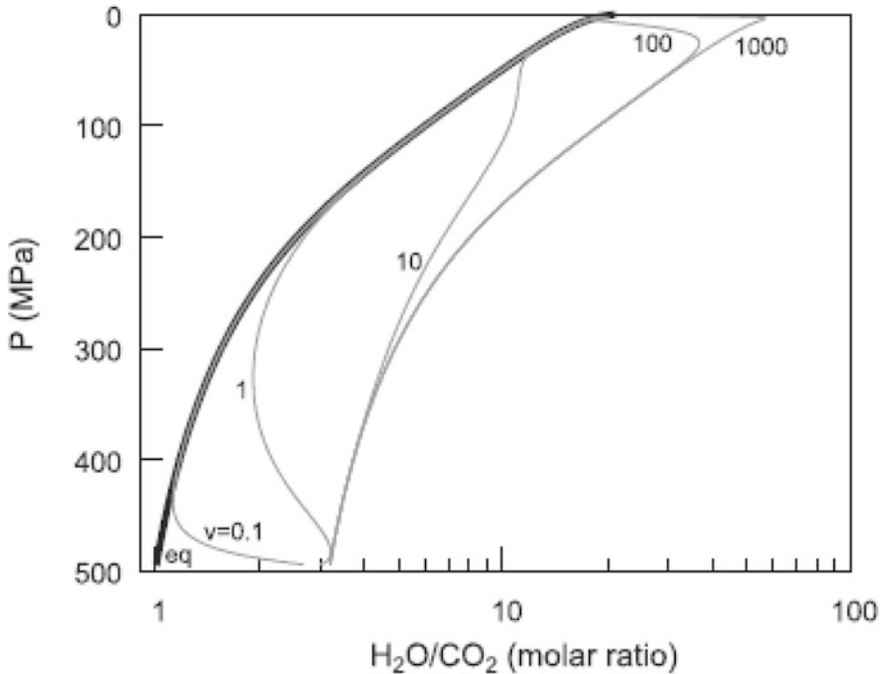
Yoshimura (2015) emphasized the relative simplicity of his model. For example, bubble growth was not considered as in other more elaborated theoretical treatments (e.g., Gonnermann and Manga 2005). Rather than continuously varying boundary (gas-melt) interface volatile concentrations and bubble-bubble distances as in a natural ascending magma, the calculations were performed step-by-step (i.e., at different pressures) along the decompression ramp, with fixed boundary concentrations and bubble-bubble distance (Yoshimura 2015). It is also important to note that the volatile concentrations on Fig. 2 correspond to averages computed by integrating the concentrations in the melt along the diffusion profiles (*distance integrated compositions*).

Gas phase compositions were calculated by mass balance using the initial volatile concentrations and the average volatile concentrations left in the melt after decompression and degassing (Yoshimura 2015). Results are shown on Fig. 3 and they correspond to compositions



**Fig. 2**  $\text{H}_2\text{O}$  and  $\text{CO}_2$  melt volatile concentrations computed with the diffusive fractionation model for different decompression/ascent rates ( $v$  from 0.1 to  $\infty$ , in m/s). Isobars (light curves) are labelled with pressure in MPa.

The heavy curve labelled “eq” is the equilibrium closed-system degassing trajectory as calculated by Yoshimura (2015). Figure redrawn from Yoshimura (2015). See text for details



**Fig. 3** Variations of the gas  $\text{H}_2\text{O}/\text{CO}_2$  molar ratio (integrated compositions) with pressure computed from the diffusive fractionation model using different decompression/ascent rates ( $v$  from 0.1 to 1000, in m/s).

The heavy curve labelled “eq” is the equilibrium closed-system degassing trajectory. Figure redrawn from Yoshimura 2015. See text for details

integrated along decompression (*pressure integrated compositions*). These compositions are more  $\text{H}_2\text{O}$ -rich (higher  $\text{H}_2\text{O}/\text{CO}_2$  ratios) than gases generated under equilibrium degassing.

### 3.2 Coupling Between Diffusion and Vesiculation

Coupling between volatile diffusion and vesiculation is a necessity in diffusive degassing models because vesiculation defines the density of bubbles, their sizes and the distances between them (e.g., Pichavant et al. 2013; Le Gall et al. 2016a, b). This issue was addressed by Yoshimura (2015), although in a relatively simplified manner. The distance between bubbles was defined as being a function of only two variables, the distance between bubbles at the bottom of the decompression column (arbitrary value) and the vesicularity. Vesicularity must change along with decompression and degassing. So, the

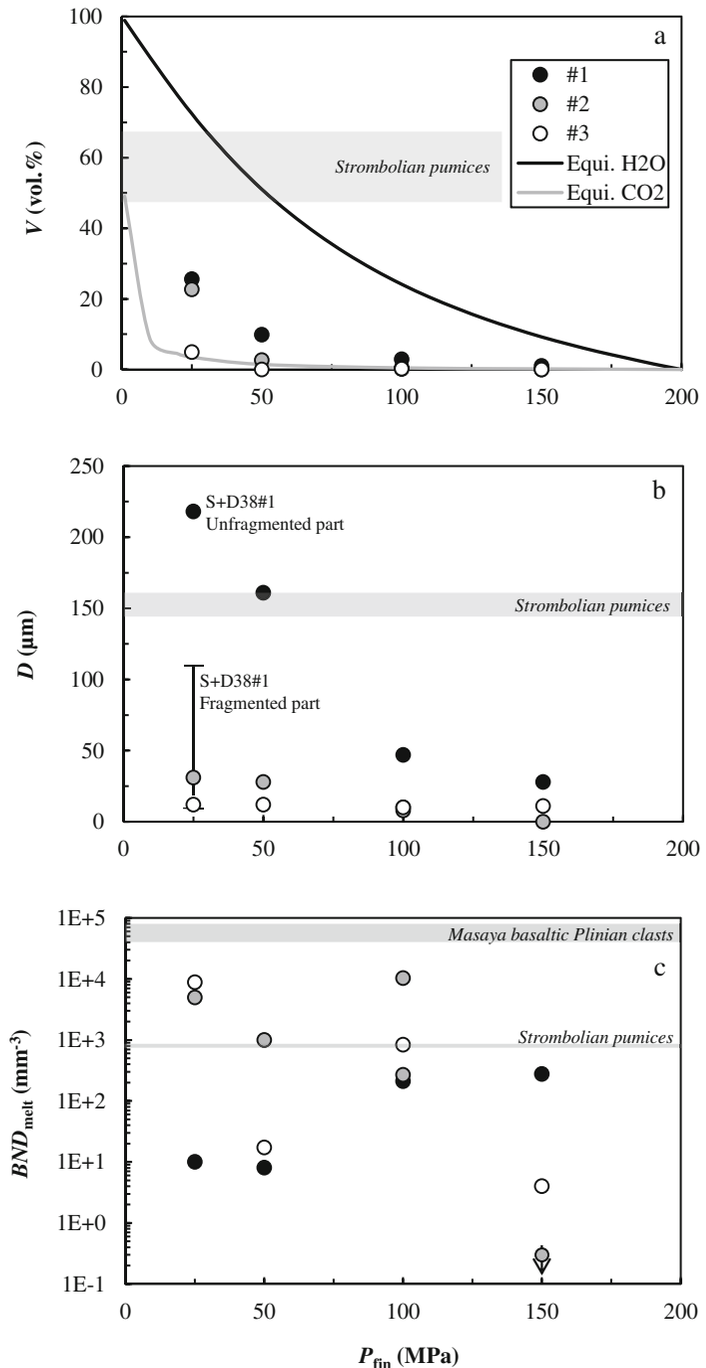
vesicularity term should embody the textural variations associated with magma ascent. In the model of Yoshimura (2015), the vesicularity was computed from the amount of volatiles exsolved upon decompression, using an equation of state for  $\text{H}_2\text{O}-\text{CO}_2$  gas mixtures to calculate the density of the gas and assuming a constant density for the melt. In so doing, it is apparent that only a vesicularity corresponding to equilibrium degassing is considered. Thus, for a given initial bubble-bubble distance, the distance between bubbles in the decompression column depends only on the equilibrium vesicularity. Degassing trajectories (Fig. 2) and integrated gas compositions (Fig. 3) were calculated on this basis.

For comparison, experimental vesicularities, bubble diameters and bubble number densities are shown on Fig. 4 for three series of basaltic melts decompressed from 200 to 25 MPa final pressure ( $P_{\text{fin}}$ ) at 3 m/s (Le Gall and Pichavant 2016a). Systematic variations within the three glass series are observed depending on  $P_{\text{fin}}$ . In

most cases, the vesicularity data plot intermediate between the two equilibrium vesicularity curves, which were computed in a similar way than Yoshimura (2015) but only for two end-member cases corresponding to pure H<sub>2</sub>O and pure CO<sub>2</sub> gas. The vesicularity data for the series #1 melts

(with pure H<sub>2</sub>O) are in general much lower than the theoretical vesicularities calculated for pure H<sub>2</sub>O gas. The data also show large changes in bubble sizes and bubble number densities that do not directly correlate with vesicularity. Le Gall and Pichavant (2016a) emphasized that

**Fig. 4** Textural data for post-decompression experimental glasses plotted as a function of final pressure ( $P_{fin}$ ) and comparison with data for natural basaltic pumices (Stromboli, Masaya). Vesicularities (a), bubble diameters (b) and bubble number densities (c) for three series of basaltic melts decompressed from 200 to 150, 100, 50 and to 25 MPa  $P_{fin}$  at 3 m/s (Le Gall and Pichavant 2016a). Pre-decompression melt concentrations, series #1: 4.91 wt% H<sub>2</sub>O (no CO<sub>2</sub>), series #2: 2.41 ± 0.04 wt% H<sub>2</sub>O, 973 ± 63 ppm CO<sub>2</sub> and series #3: 0.98 ± 0.16 wt% H<sub>2</sub>O, 872 ± 45 ppm CO<sub>2</sub>. Note that, for charge S + S38#1 which was partially fragmented, the vesicularity (a) and BND (c) data concern the unfragmented part. The bubble diameter data (b) are for both the unfragmented (black symbol) and the fragmented (minimum and maximum values) parts. Figure redrawn from Le Gall and Pichavant (2016a)



degassing textures result from several processes including bubble nucleation, growth, coalescence, plus buoyancy-driven bubble migration. We conclude that, although the diffusive fractionation model of Yoshimura (2015) provides a basis for coupling volatile diffusion calculations and vesiculation processes, more work is needed to incorporate the complex textural changes associated with ascent of volatile-bearing melts.

## 4 Implications for Gas Phase Compositions

### 4.1 Available Data and Models

Despite the limitations noted above, the diffusive fractionation model provides a framework to interpret the experimental observations and the systematic deviations from equilibrium degassing observed in CO<sub>2</sub>-bearing systems. However, it should be emphasized that the model uses analytical data (glass volatile concentrations) and physicochemical properties (volatile diffusivities) related only to the melt phase. The question arises of the consequences of disequilibrium degassing for the gas phase composition. It is worth remembering here that the precursory signals come from gas data.

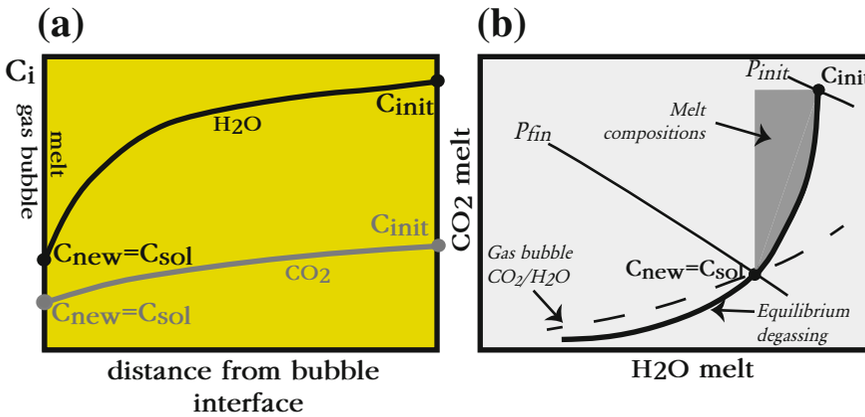
In the decompression experiments summarized above, the gas phase has not been chemically analysed although some mass balance calculations were performed to estimate the composition of the gas phase in the H<sub>2</sub>O-, CO<sub>2</sub>- and S-bearing experiments of Le Gall et al. (2015a). However, it is emphasized that, with this method, only bulk experimental gas compositions are provided (*charge and pressure integrated compositions*). No information is available on the composition of individual bubbles generated during decompression. The gas calculations performed by Yoshimura (2015) also use a similar mass balance approach, i.e., pressure integrated fluid compositions are given. However, the local gas at the gas-melt interface has an equilibrium composition (local gas-melt equilibrium). The differences between the disequilibrium (calculated with the model) and the equilibrium

(calculated assuming equilibrium degassing) gases (Fig. 3) is the consequence of CO<sub>2</sub> degassing being hampered by its retention within the melt. Therefore, disequilibrium is evidenced in the compositions of the pressure integrated fluids.

### 4.2 Composition of Gas Bubbles

The experiments and the diffusive fractionation model show that melt and gas both evolve under disequilibrium during magma ascent and degassing. For the melt, this conclusion is based either on volatile concentration measurements in glass at some distance of the gas/melt interface (Pichavant et al. 2013; Yoshimura 2015; Le Gall and Pichavant 2015a, 2016a, b) or on average concentrations calculated by integration along diffusion profiles (Yoshimura 2015). For the gas, constraints are available only on integrated compositions (Le Gall et al. 2015a; Yoshimura 2015). Since the gas phase is essential for the application to volcanoes, and given the interpretations proposed for the melt phase, constraints on the gas phase composition at smaller scales are becoming necessary. This requires linking compositions of gas bubbles in decompression experiments not only with pressure but also with volatile concentrations of local melts as well as with degassing textures and mechanisms.

As a way toward this direction, local gas-melt equilibrium can be assumed. This implies that chemical equilibrium persists locally between gas and melt, despite evidence for disequilibrium at larger scales. Therefore, the volatile compositions of melt and gas at the interface are defined by equilibrium partitioning of volatiles between these two phases (e.g., Dixon and Stolper 1995). To illustrate this concept, a schematic representation of the gas-melt interface for a H<sub>2</sub>O- and CO<sub>2</sub>-bearing melt decompressed isothermally from an initial ( $P_{\text{init}}$ ) to a final ( $P_{\text{fin}}$ ) pressure is shown on Fig. 5a. Initial volatile concentrations ( $C_{\text{init}}$ ), together with the  $P_{\text{init}}$  and  $P_{\text{fin}}$  isobars and the equilibrium degassing trajectory are shown on the H<sub>2</sub>O-CO<sub>2</sub> diagram of Fig. 5b. If local gas-melt equilibrium is assumed, the interface melt H<sub>2</sub>O and CO<sub>2</sub> concentrations at  $P_{\text{fin}}$  ( $C_{\text{new}}$ )



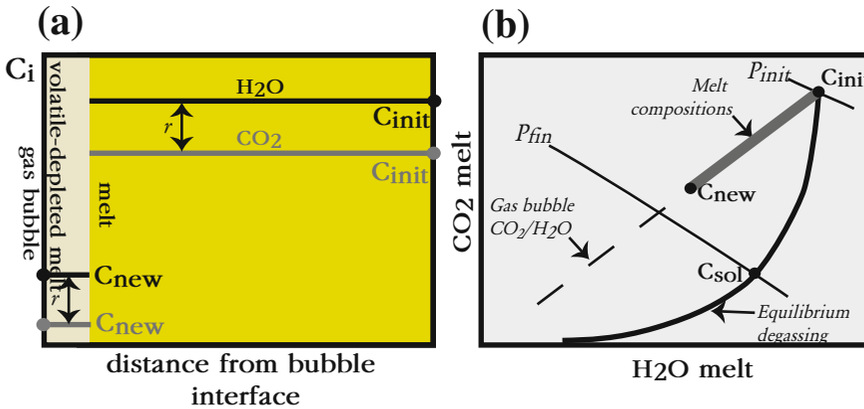
**Fig. 5** Schematic illustration of local gas-melt equilibrium. **a** detail of the gas-melt interface region in a concentration ( $C_i$ ) versus distance diagram where  $C_i$  refers to the volatile concentration in the melt. The gas bubble is on the left. The two curves are melt volatile concentration profiles for H<sub>2</sub>O (black) and CO<sub>2</sub> (grey) respectively, generated as a result of diffusion in the melt during decompression from  $P_{init}$  to  $P_{fin}$ .  $C_{init}$  give volatile concentrations of the pre-decompression melt,  $C_{new}$  gas/melt interface volatile concentrations at  $P_{fin}$  and  $C_{sol}$  volatile solubilities at  $P_{fin}$ . Black lettering is used for

H<sub>2</sub>O and grey for CO<sub>2</sub>. **b** H<sub>2</sub>O–CO<sub>2</sub> diagram illustrating the evolution during decompression and degassing. The black bold curve is the equilibrium degassing trajectory. The two black lines are isobars labelled with initial ( $P_{init}$ ) and final ( $P_{fin}$ ) pressures along the decompression path. The dashed curve is the CO<sub>2</sub>/H<sub>2</sub>O isopleth passing through  $C_{sol}$  and it defines the composition of the gas bubble in local equilibrium with the interface melt. The shaded domain gives the range of possible melt compositions generated upon decompression from  $P_{init}$  down to  $P_{fin}$

are equal to their solubilities ( $C_{sol}$ ) at  $P = P_{fin}$  (intersection of the equilibrium degassing curve with the  $P_{fin}$  isobar, Fig. 5b). Note that diffusive fractionation generates H<sub>2</sub>O and CO<sub>2</sub> concentration gradients within the melt (Fig. 5a), the range of possible melt compositions during decompression being represented by the dark grey region in Fig. 5b. The interface melt is the only melt at equilibrium with the local gas at  $P_{fin}$  which has a CO<sub>2</sub>/H<sub>2</sub>O corresponding to the fluid isopleth on Fig. 5b (e.g., Dixon and Stolper 1995). For a low pressure (e.g., 25 MPa), the local gas (e.g., a gas bubble nucleated at  $P_{fin}$ ) is relatively H<sub>2</sub>O-rich. In comparison, the pressure integrated gas assuming bulk *equilibrium* degassing from  $P_{init}$  to  $P_{fin}$  would be necessarily less H<sub>2</sub>O-rich since most of the CO<sub>2</sub> must have been outgassed from the melt. This gas is less H<sub>2</sub>O-rich than the pressure integrated gas produced by *disequilibrium* degassing from  $P_{init}$  to  $P_{fin}$  (Fig. 3). Thus, individual bubbles nucleated at  $P_{fin}$  can have CO<sub>2</sub>/H<sub>2</sub>O different from the composition of integrated gases generated continuously during decompression.

An alternative way to constrain the composition of gas bubbles is illustrated on Fig. 6. It starts from the observation that bubble nucleation is, from a kinetic point of view, an instantaneous process (e.g., Mourtada-Bonnefoi and Laporte 2002, 2004). Nucleation of a gas bubble draws volatiles from the local melt and the possibility that the initial CO<sub>2</sub>/H<sub>2</sub>O of the gas bubble is the same as the local melt should be considered. According to this hypothesis, represented schematically on Fig. 6, the local melt next to the nucleated bubble ( $C_{new}$ ) is volatile-depleted but its CO<sub>2</sub>/H<sub>2</sub>O ( $r$ , Fig. 6a) is the same than the initial melt ( $C_{init}$ ). Melt and gas bubble compositions are thus both located on a mixing line between  $C_{init}$  and  $C_{new}$  which passes through the origin of the H<sub>2</sub>O–CO<sub>2</sub> diagram (Fig. 6b). The net result is the nucleation of individual gas bubbles more CO<sub>2</sub>-rich than expected from local gas-melt equilibrium (Fig. 5b).

The previous discussion emphasizes the compositional variability of nucleated gas bubbles and the need for their direct analysis in decompression experiments. Comparison between the



**Fig. 6** Hypothetical model for the composition of a gas bubble nucleated during decompression of H<sub>2</sub>O-, CO<sub>2</sub>-bearing basaltic melts. **a** detail of the gas-melt interface region in a concentration ( $C_i$ ) versus distance diagram where  $C_i$  refers to the volatile concentration in the melt. The gas bubble is on the left. The two horizontal lines are melt volatile concentrations for H<sub>2</sub>O (black) and CO<sub>2</sub> (grey) drawn as straight lines because bubble nucleation is an instantaneous event. The narrow zone near the interface is the melt region depleted in volatiles drawn to form the bubble.  $C_{new}$  are volatile concentrations in the depleted melt region. Notice that the H<sub>2</sub>O/CO<sub>2</sub> ratio ( $r$ ) is identical in both the depleted and non-depleted melt regions because during nucleation volatiles are drawn from the local melt and the initial CO<sub>2</sub>/H<sub>2</sub>O of the gas bubble is the same as the local melt.  $C_{init}$  give volatile concentrations of the

pre-decompression melt,  $C_{new}$  gas/melt interface volatile concentrations left after bubble nucleation at  $P_{fin}$ . Black lettering is used for H<sub>2</sub>O and grey for CO<sub>2</sub>. **b** H<sub>2</sub>O–CO<sub>2</sub> diagram illustrating the evolution during decompression and degassing. The black bold curve gives the schematic location of the theoretical equilibrium degassing trajectory. The two black lines are isobars labelled with initial ( $P_{init}$ ) and final ( $P_{fin}$ ) pressures along the decompression path. Melts produced as a result of decompression and bubble nucleation plot on the straight line joining  $C_{init}$  and  $C_{new}$ . This line passes through the origin of the diagram because both  $C_{init}$  and  $C_{new}$  have the same CO<sub>2</sub>/H<sub>2</sub>O ratio. Note that the location of  $C_{new}$  along this line is arbitrary. The CO<sub>2</sub>/H<sub>2</sub>O ratio of the gas bubble (dashed line) is also the same as  $C_{init}$  and  $C_{new}$ . It is higher than the gas bubble controlled by local gas-melt equilibrium (Fig. 5)

CO<sub>2</sub>/H<sub>2</sub>O of nucleated bubbles and results of gas-melt volatile partitioning models (e.g., Dixon and Stolper 1995; Papale et al. 2006) would provide a crucial test of the local gas-melt equilibrium model (Fig. 5). If the nucleated bubbles prove to be CO<sub>2</sub>-rich, then alternative models of control of gas composition would be supported (Fig. 6). One important aspect is that, in CO<sub>2</sub>-bearing systems, bubble nucleation during decompression is continuous, occurring over a large pressure range (Le Gall and Pichavant 2016a, b). This is because CO<sub>2</sub>-bearing melts are volatile-supersaturated (Fig. 1) and, so, the driving force for nucleation of new bubbles is always present. Thus, decompression of CO<sub>2</sub>-bearing melts continuously leads to the nucleation of new bubbles, which increases the relevance of the hypothetical mechanism illustrated in Fig. 6.

## 5 Discussion and Perspectives for Gas Monitoring

### 5.1 Degassing Processes

Experimental simulations show that, for ascent rates expected in natural systems, equilibrium degassing occurs in pure H<sub>2</sub>O melts. In contrast, results for CO<sub>2</sub>-bearing melts conclusively demonstrate that degassing generates melt volatile concentrations out of equilibrium. The experimental database supporting this conclusion has recently expanded. It now includes basaltic and rhyolitic melts, and S-bearing as well as S-free systems. Several of those experimental decompression studies have been scaled to natural systems so that results are realistic and

applicable. The decompression experiments on Stromboli basalt cover ascent rates ranging from 0.25 to 3 m/s (Pichavant et al. 2013; Le Gall and Pichavant 2016a, b; Le Gall et al. 2015a), well in the range of current estimates for basaltic magmas (e.g., Rutherford 2008; Peslier et al. 2015). An ascent rate approximately 10 times faster was used by Yoshimura (2015). Therefore, in the case of basaltic magmas, equilibrium degassing should be viewed more as a reference situation rather than as a general mechanism. This is quite a change in paradigm which has major implications for how gas signals are interpreted.

One remaining issue concerns the role of crystals on bubble nucleation, heterogeneous rather than homogeneous. All decompression studies considered in this paper were performed on very crystal-poor, if not totally crystal-free, melts and bubble nucleation appears to be mostly homogeneous. Crystals present in experimental basaltic products include Fe–Ti oxides (Le Gall and Pichavant 2015b) and rare Fe sulphides (Le Gall et al. 2015a). Le Gall and Pichavant (2015b) have documented heterogeneous nucleation of bubbles on Fe–Ti oxide crystals (and also on Fe sulphides, Le Gall et al. 2015a). Recently, Shea (2017) has stressed the importance of magnetite as a key mineral phase promoting heterogeneous bubble nucleation in natural magmas. However, Fe oxide phenocrysts and sulphides are uncommonly present in amounts exceeding a few vol.% in natural magmas. This led Le Gall and Pichavant (2015b) to conclude that heterogeneous bubble nucleation is not an important mechanism in basaltic melts if driven by Fe oxides. Yet, heterogeneous nucleation on silicate phases is still an open question. For example, olivine, clinopyroxene and plagioclase are typical phenocrysts and microlites in Stromboli basalts (e.g., Pichavant et al. 2011). On the basis of limited textural evidence, Pichavant et al. (2013) ruled out the possibility of heterogeneous nucleation of gas bubbles on clinopyroxene and olivine crystals. However, additional investigations seem warranted to guarantee full applicability of the decompression experiments above.

Disequilibrium degassing, as documented in the experiments, is the consequence of the anomalous behaviour of CO<sub>2</sub>. CO<sub>2</sub>-supersaturated melts are systematically generated during decompression. The interpretation suggested by Pichavant et al. (2013) and quantitatively formulated by Yoshimura (2015) is that, because of its restricted diffusive mobility within the melt, CO<sub>2</sub> has limited access to the gas phase for timescales typical of magma ascent. However, our knowledge of volatile diffusivities in silicate melts is still very fragmentary. There are very few diffusivity data for H<sub>2</sub>O and CO<sub>2</sub> on the same melt. S is another volatile which reputedly has a slow diffusivity in silicate melts. Yet, the behaviour of S during degassing differs from that of CO<sub>2</sub>, although we are still short of S solubility data for basaltic melts (e.g., Lesne et al. 2015). Acquisition of fundamental data (especially volatile diffusivity and solubility data for basaltic melts) is needed for the elaboration of more detailed interpretations of the decompression experiments. Future magma ascent models should also incorporate the textural complexities associated with the vesiculation process.

## 5.2 Gases as Unrest Signals

Pre-eruptive changes in gas ratios have been observed at several basaltic volcanoes such as Stromboli (Burton et al. 2007; Aiuppa et al. 2010), Etna (Aiuppa et al. 2007) and Villarrica (Aiuppa et al. 2017) among others. Transition from passive degassing to more explosive paroxysmal eruption regimes is marked by temporal increases of the CO<sub>2</sub>/SO<sub>2</sub> gas ratio in the volcanic plume. In some cases, the CO<sub>2</sub>/SO<sub>2</sub> variations are correlated with a decrease of the H<sub>2</sub>O/CO<sub>2</sub> gas ratio (e.g., Aiuppa et al. 2017). These variations in volcanic gas ratios have been generally interpreted to reflect different pressures of gas-melt segregation in the conduit, high CO<sub>2</sub>/SO<sub>2</sub> (and low H<sub>2</sub>O/CO<sub>2</sub>) indicating deep conditions and low CO<sub>2</sub>/SO<sub>2</sub> (and high H<sub>2</sub>O/CO<sub>2</sub>) shallow conditions (e.g., Edmonds 2008; Burton

et al. 2007; Allard 2010; Aiuppa et al. 2017). In this approach, the pressure-dependent evolution of the gas phase exsolved upon magma ascent and decompression is calculated by using volatile saturation codes (Newman and Lowenstern 2002; Moretti and Papale 2004; Papale et al. 2006; Burgisser et al. 2015). This implicitly assumes chemical equilibrium between gas and melt, an assumption which, as shown above, is now largely questioned. If disequilibrium magma degassing is accepted, then the consequences for the interpretation of gas signals need to be examined.

Firstly, one might argue that gas-melt equilibrium can persist at local scale, despite disequilibrium at larger scales. Thus, volatile saturation codes could still be used and applied to local gas and melt compositions, for example to model the composition of unconnected bubbles nucleated within the melt. In contrast, volcanic gases necessarily require, to be sampled, that the magma is permeable and, so, that the gas phase is connected. It is quite possible that the gases sampled are mixtures of different components, either integrated from several discrete degassing events along ascent or issued from different parts of the plumbing system. Therefore, volcanic gases are probably more representative of integrated compositions as discussed above than to compositions of local gases. We have shown previously that individual bubbles with compositions defined by local gas-melt equilibrium at a given pressure (Fig. 5) can have  $\text{CO}_2/\text{H}_2\text{O}$  different from integrated gases generated continuously during decompression (Fig. 3). We conclude to the limited applicability of local gas melt equilibrium to interpret volcanic gas ratios.

Secondly, disequilibrium gas-melt degassing due to  $\text{CO}_2$  retention within the melt implies that  $\text{CO}_2/\text{SO}_2$  and  $\text{H}_2\text{O}/\text{CO}_2$  gas ratios can no longer be directly related to pressures of gas-melt segregation. Calculations using the diffusive fractionation model (Fig. 3) show that the pressure integrated gases have a higher  $\text{H}_2\text{O}/\text{CO}_2$  (and also presumably a lower  $\text{CO}_2/\text{SO}_2$  because  $\text{CO}_2$  is retained

within the melt) than the same gases calculated assuming equilibrium with the melt (Fig. 3). This demonstrates the possibility of changing the gas ratios depending on the degassing mechanism (equilibrium vs. disequilibrium). It is worth emphasizing that disequilibrium degassing associated with  $\text{CO}_2$  retention produces integrated fluids that are less, not more,  $\text{CO}_2$ -rich (Fig. 3).

The  $\text{CO}_2$ -rich gases observed on basaltic volcanoes have been generally attributed to deep-seated processes such as fluxing of  $\text{CO}_2$  or arrival of  $\text{CO}_2$ -rich magmas (e.g., Aiuppa et al. 2010, 2017; Allard 2010). In contrast, the degassing mechanism of Fig. 6 (although it needs validation from direct analysis of gas bubbles in decompression experiments) allows  $\text{CO}_2$ -rich gas bubbles to be generated at low pressures. It also provides an example of how gas ratios can be changed at constant pressure depending on the degassing mechanism. The initially  $\text{CO}_2$ -rich bubbles (Fig. 6) will probably shift rapidly with time toward lower  $\text{CO}_2/\text{H}_2\text{O}$  because of preferential diffusion of  $\text{H}_2\text{O}$  from the melt. However, nucleation is a continuous process in  $\text{CO}_2$ -bearing basaltic melts (Le Gall and Pichavant 2016a, b) and reequilibration of previously nucleated bubbles by diffusion will be accompanied by the nucleation of new  $\text{CO}_2$ -rich bubbles.

We conclude that future developments in the interpretation of gas data require progress from both sides, experimental and volcanological. Some crucial experimental information at small scale is still missing such as the composition of individual gas bubbles nucleated in the decompression experiments and the influence of crystals on bubble nucleation. In parallel, at larger scales, the representativity and the significance of the gas phase sampled on active basaltic volcanoes needs to be better demonstrated, for example by combining gas measurements with detailed textural studies of eruption products. It is expected that future work will narrow the gap in scales between experiments and gas measurements to refine interpretations of gas compositions as unrest signals.

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