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HAL Id: insu-00745334
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Submitted on 25 Oct 2012

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Extremely reducing conditions reached during basaltic intrusion
in organic matter-bearing sediments

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Abstract

Redox conditions in magma are widely interpreted as internally buffered and closely related to that of their mantle source regions. We use thermodynamic calculations to show that high-temperature interaction between magma and organic matter can lead to a dramatic reduction of the magma redox state, and significant departure from that of the original source. Field studies provide direct evidence of the process that we describe, with reported occurrences of graphite and native iron in igneous mafic rocks, implying very reducing conditions that are almost unknown in average terrestrial magmas. We calculate that the addition of 0.6 wt%
organic matter (in the form of CH or CH$_2$) to a standard basalt triggers graphite and native iron crystallisation at depths of few hundred meters.

Interaction with organic matter also profoundly affects the abundance and the redox state of the gases in equilibrium with the magma, which are CO-dominated with H$_2$ as the second most abundant species on a molar basis, H$_2$O and CO$_2$ being minor constituents. The assimilation of only 0.1 wt% organic matter by a basalt causes a decrease in its oxygen fugacity of 2-orders of magnitude. The assimilation of 0.6 wt% organic matter at depths < 500 m implies minimum CO content in the magma of 1 wt%, other gas components being less than 0.1 wt%. In the light of our calculations, we suggest that the production of native iron-bearing lava flows and associated intrusions was most likely accompanied by degassing of CO-rich gases, whose fluxes depended on the magma production rates.

**Key words:** magma-organic matter interaction, graphite native iron saturation, gas emissions, carbon monoxide, Siberian Traps, Disko Island.

1. **Introduction**

Volcanic gases may have critically influenced the evolution of planetary atmospheres and biogeochemical cycles (Holland, 2002; Kump and Barley, 2007; Gaillard et al., 2011). The classic view is that the nature and the abundance of gas emanations reflect redox state and volatile abundances of the magma source region in the mantle (Jambon, 1994; Carmichael, 1991). Volatiles are released to the atmosphere during volcanic degassing and re-introduced into the mantle by subduction (e.g. Hilton et al., 2002; Wallace, 2005). Mantle redox state is considered to have undergone little evolution over much of Earth history (Canil, 2002; Li and Lee, 2005; Trail et al., 2011), resulting in H$_2$O-CO$_2$-(SO$_2$) dominated volcanic gases for most magmas, regardless of their tectonic setting and age (Li and Lee, 2005; Gaillard et al., 2011; Trail et al., 2011). This picture is corroborated by geochemical surveys of the composition of
volcanic gases emitted by active volcanoes (e.g. Symonds et al., 1994; Aiuppa et al., 2007, 2012; Edmonds and Gerlach, 2007; Metrich et al., 2011), showing that volcanic species are mainly composed of H₂O, CO₂ and SO₂. Recent works show, however, that magma-host rock interaction contributes additional gases that can enhance or even overwhelm the mantle-derived contribution (Svensen et al. 2004, 2007, 2009; Iacono-Marziano et al., 2007, 2009; Ganino and Arndt, 2009; Troll et al., 2012), and locally impacts the magma redox state (Ganino et al., 2008). During continental flood magmatism, contact metamorphism of country rocks is proposed to have controlled the nature of gas emissions and their environmental consequences (Svensen et al. 2004, 2007, 2009; Ganino and Arndt, 2009). The high biological impact ascribed to the emplacement of the Siberian Traps and the Emeishan igneous province, for instance, has been attributed to the occurrence of carbonates, evaporites and organic matter-rich rocks in the sedimentary sequences traversed by ascending magmas (Svensen et al. 2004, 2009; Ganino and Arndt, 2009). Previous studies focused on low temperature (<650°C) contact metamorphism, which generates carbon and sulphur gases (CO₂, CH₄, SO₂) and halocarbons (Svensen et al. 2004, 2007, 2009; Ganino and Arndt, 2009; Retallack and Jahren, 2008). The high temperature contribution of magmatic assimilation has been generally neglected, being difficult to quantify (Ganino and Arndt, 2009). Here we study the high temperature (1200°C) assimilation of carbonaceous rocks into basaltic magmas, by simulating the incorporation of organic matter in the form of CH or CH₂, and investigating its effect on the redox state of the magma, its mineralogical assemblage and its volatile content. Crystal-melt-gas equilibria are first explored via thermodynamic calculations, taking into account both magmatic and sedimentary contributions to gas production. The conditions of basalt-organic matter interaction in relevant geological settings will be then presented.

Numerous studies have investigated the composition and the thermodynamics of volcanic gases, paying particular attention to the redox processes governing their composition (e.g.
Giggenbach (1987, 1997; Gerlach, 1993; Moretti et al, 2004). Giggenbach (1987) dealt with C-O-N-H-S gases, taking into account the role of the rock matrix in buffering their redox conditions. Gerlach (1993) investigated C-O-H-S-Cl-F species and modelled their exchanges with the melt in terms of transfer of oxygen from the gas to the lava. Moreover, Giggenbach (1997) explored the role of organic sedimentary material in the production of natural carbon gasses. Compared to these studies, the main originality of our work is that we combine: (i) high temperature magmatic conditions, (ii) gas-melt equilibria, i.e. not only oxygen exchange between gas and melt is taken into account, but also the solubility of some gas species in the melt, (iii) open system conditions, i.e. both magmatic and non-magmatic volatiles are considered, (iii) saturation of graphite and native iron in the melt, as we deal with reduced condition.

2. Methods: thermodynamic calculations

We modelled the composition of volcanic gases resulting from interaction with organic matter and magma degassing. To reach this aim we used gas-melt thermodynamic calculations that take into account S-H-O-C gaseous species at temperatures and pressures in equilibrium with basaltic liquids (Gaillard and Scaillet, 2009; Gaillard et al., 2011). We considered that the melt, prior to interaction with organic compounds, is basaltic in composition with 8 wt% FeO. At the initial conditions (P = 200 MPa, T = 1200°C) the basaltic melt is saturated in CO₂, H₂O and S: 0.08, 0.50 and 0.10 wt% respectively (Mavrogenes and O'Neill, 1999; Iacono-Marziano et al., in press). We investigated the roles of the initial oxidation state and gas content of the magma prior to interaction and tested the effect of coal (CH) or hydrocarbon (CH₂) incorporation. The initial oxygen fugacity (fO₂) of the magma varied between the Fayalite-Magnetite-Quartz (FMQ) and the Nickel-Nickel Oxide (NNO) buffers, as generally observed in basaltic magmas (Li and Lee, 2005), yielding Fe₂O₃/FeO mole ratios between
0.07 and 0.11. The initial (i.e. at 200 MPa) gas content of the magma was between 0.3 and 0.7 
\%
, yielding total volatile contents (dissolved in the melt + bubbles) of 0.50 wt\% \( H_2O \), 0.10 
\text{wt\%}\ S \text{ and } 0.24-0.64 \ \text{wt\%}\ CO_2 \), which are typical of undegassed basalts (Dixon et al., 2008).

We tested initial dissolved water content between 0.2 and 1 \text{wt\%} \ and found it to have a 
negligible influence on our calculations. Starting conditions are listed in Table 1. Calculations 
show also that varying the total FeO content of the basalt between 8 and 13 \text{wt\%} \ changes by 
less than 5\% \ the concentration of the species with mole fractions higher than 0.1.

The gas composition was calculated following Gaillard and Scaillet (2009) and Gaillard et al. 
(2011). This methodological approach is long known as the “Equilibrium constants and mass 
balance method” (Holloway, 1987) to calculate equilibria in the C-O-H system, briefly 
recalled below. The mole fractions of the different gas species were calculated from the 
thermodynamic constants of the following reactions at \( P \) and \( T \) (after Shi and Saxena, 1992).

\[
\begin{align*}
\text{C}^{[\text{graphite}]} + \text{O}_2 &= \text{CO}_2^{[\text{fluid}]} \quad (1) \\
\text{CO}^{[\text{fluid}]} + \frac{1}{2}\text{O}_2 &= \text{CO}_2^{[\text{fluid}]} \quad (2) \\
\text{H}_2^{[\text{fluid}]} + \frac{1}{2}\text{O}_2 &= \text{H}_2\text{O}^{[\text{fluid}]} \quad (3) \\
\text{CH}_4^{[\text{fluid}]} + \text{O}_2 &= \text{CO}_2^{[\text{fluid}]} + 2\text{H}_2\text{O}^{[\text{fluid}]} \quad (4) \\
\frac{1}{2}\text{S}_2^{[\text{fluid}]} + \text{O}_2 &= \text{SO}_2^{[\text{fluid}]} \quad (5) \\
\text{H}_2\text{S}^{[\text{fluid}]} + \frac{1}{2}\text{O}_2 &= \frac{1}{2}\text{S}_2^{[\text{fluid}]} + \text{H}_2\text{O}^{[\text{fluid}]} \quad (6)
\end{align*}
\]

Melt-gas equilibria were also considered:

\[
\begin{align*}
\text{H}_2\text{O}^{[\text{fluid}]} + \text{O}^{2-}^{[\text{melt}]} &= \text{2 OH}^{-}^{[\text{melt}]} \quad (7) \\
\text{CO}_2^{[\text{fluid}]} + \text{O}^{2-}^{[\text{melt}]} &= \text{CO}_3^{2-}^{[\text{melt}]} \quad (8) \\
\text{S}_2^{[\text{fluid}]} + 2\text{O}^{2-}^{[\text{melt}]} &= \text{O}_2 + 2\text{S}^{2-}^{[\text{melt}]} \quad (9) \\
2\text{FeO}^{[\text{melt}]} + \frac{1}{2}\text{O}^{2-}^{[\text{gas}]} &= \text{Fe}_2\text{O}_3^{[\text{melt}]} \quad (10)
\end{align*}
\]

Equilibria (1) to (6) and (10) were simultaneously solved at every given pressure and 
temperature, under the requirement that the amounts of H, S, O and C remain constant in the
system (gas + melt + graphite). Seven redox couples were considered for each calculation, involving sulphur (with 3 different redox states), carbon (with 4 different redox states) and hydrogen (with 2 different redox states), together with iron (2 redox states). Each equilibrium constant for equilibria from (1) to (6) is compiled in Symonds and Reed (1993), while solubility laws for $\text{H}_2\text{O}$, $\text{CO}_2$ and $\text{S}$ (equilibria 7-9) were taken from Iacono-Marziano et al. (in press) and Mavrogenes and O’Neill (1999). The ferric-ferrous equilibrium (10) was computed after Kress and Carmichael (1991). The concentrations of reduced C species (e.g. CO, CH$_4$) in the basaltic melt was neglected in our calculations, as they are known to have very low solubility in a basaltic melt at ~200-300 MPa (Pan et al., 1992; Morizet et al., 2010). In conclusion, our approach combines existing experimental studies performed on specific volatiles into a multi-component model, under the assumption of gas-melt equilibrium. Organic matter was incrementally added up to 0.8 wt%. Two types of organic compounds were taken into account, CH and CH$_2$, which are representative of mature organic matter that is most likely to interact with basaltic magmas (e.g. coal or petroleum). To simplify our calculations, we ignore the possible presence of oxygen in the organic matter, as the oxygen content generally decreases with the maturity of the organic matter (Hetenyi, 1998). Degassing is simulated by decreasing pressure, keeping the masses of the different components (H, S, C, O) constant. Gas compositions are calculated at variable final pressures and shown below for 10 and 0.1 MPa. Figure 1 in the Supplementary Material schematically summarizes the strategy of the numerical calculations.

3. Results

3.1 Fluid phase composition

Assimilation of organic matter has dramatic effects both on the amount and the redox state of the gases in equilibrium with the basalt (Fig.1). When no organic matter is assimilated the
fluid phase consists mainly of H$_2$O and CO$_2$ (respectively 0.52-0.66 and 0.15-0.33 on a molar basis, with 0.05-0.14 SO$_2$, and H$_2$, CO, H$_2$S and S$_2$ <0.05) as typically observed in volcanic gas emissions from basalts worldwide (Symonds et al., 1994; Edmonds and Gerlach, 2007). Figure 1 (see also Figure 2 in the Supplementary Material) shows the influence of the initial conditions (initial gas content of the magma and initial fO$_2$) on gas phase compositions at 0.1 MPa resulting from the assimilation of CH or CH$_2$. For CH addition, we show how gas composition varies for initial fO$_2$ of NNO or FMQ, and for an initial gas proportion of 0.3 or 0.7% of the magma. More oxidizing conditions (initial fO$_2$ of NNO) slightly favours the survival of oxidized species (e.g. H$_2$O, CO$_2$) and S species in general (H$_2$S, S$_2$, SO$_2$) for a given amount of organic matter added to the system. Higher initial gas contents (0.7 wt%) also slightly promote oxidized species relative to reduced ones. The most extreme cases are therefore represented by: (i) fO$_2$ = NNO and gas proportion = 0.7% (Fig.1a), and (ii) fO$_2$ = FMQ and gas proportion = 0.3% (Fig.1b). CH$_2$ addition slightly enhances the production of H$_2$ respect to that of CO. Similarly to CH addition, the two most extreme cases are shown in Fig.1c and d. In all cases, with the assimilation of only 0.25-0.35 wt% CH or CH$_2$, the gas composition changes from CO$_2$-H$_2$O dominated to strongly reduced and CO-H$_2$ dominated. In general, the effects of the initial fO$_2$ and gas fraction are minor, as are differences arising from whether CH or CH$_2$ is added: in all cases the degassing trends are very similar and the amount of the dominant species (i.e. those with mole fractions >0.1) changes by less than 30%, for a given amount of organic matter assimilated (Fig.SM2).

3.2 Graphite and native iron saturation

An important consequence of the addition of organic matter is the crystallization of graphite from the magma (equilibrium 1). Graphite saturation depends only slightly on factors such as the type of organic matter assimilated (CH or CH$_2$), the initial redox conditions, and the initial
gas content of the magma but strongly on pressure and therefore on depth (Fig. 2). At pressures higher than 60 MPa, graphite crystallizes from the magma when low amounts of organic matter are assimilated (0.1-0.3 wt%). Coal and oil ingestion at 1200°C and graphite crystallization can be simplified as:

\[(x) \text{CH} + (y) O^{\text{magma}} \rightarrow (y) \text{CO} + (1/2 x) \text{H}_2 + (x-y) \text{C} \quad (11)\]

\[(x) \text{CH}_2 + (y) O^{\text{magma}} \rightarrow (y) \text{CO} + (x) \text{H}_2 + (x-y) \text{C} \quad (12)\]

where \(x\) and \(y\) are stochiometric coefficients that are essentially affected by pressure. For a given amount of organic matter assimilated \((x)\), high pressure favours graphite crystallization (i.e. low \(y\)), while low pressure enhances CO and \(H_2\) production, by subtracting oxygen to the magma (i.e. high \(y\)). At shallow depths (i.e. low pressure), increasing CH or \(CH_2\) addition therefore causes a dramatic decrease in magma \(fO_2\), whereas at deeper conditions the decrease in \(fO_2\) is less pronounced. Reactions (11) and (12) conveniently summarize the overall process of assimilation of organic matter by a basaltic melt, however our calculations also take into account other C-H-O and S species (i.e. \(CO_2\), \(H_2O\), \(CH_4\), \(S_2\), \(SO_2\), \(H_2S\); see reactions 1 to 6), which have concentrations lower than 1 wt%, and are therefore neglected here for simplifying.

Figure 3 shows the evolution of the redox state of a magma with an initial gas content of 0.3 wt % at 200 MPa as a function of the assimilated organic matter. The \(fO_2\) of the magma rapidly decreases with the addition of organic matter from the FMQ buffer (initially imposed conditions). For only 0.1 wt% CH assimilated, \(fO_2\) decreases by more than 2 log-units. For more than 0.5 wt% CH assimilated, \(fO_2\) decreases by ~6 log units (FMQ-6). When the \(fO_2\) of the magma reaches FMQ-6, native iron crystallizes (Gaillard and Scaillet, 2009) and buffers magma redox state, until all FeO is exhausted. The amount of organic matter needed to reach graphite and native iron saturation due to the variable initial conditions (i.e. type of organic matter assimilated, initial gas fraction and initial \(fO_2\) of the magma), varies by less than 30%
at a given pressure (Fig. 2). At graphite saturation, native iron crystallization occurs with decreasing pressure by graphite volatilization, as expressed by the reaction:

$$C_{\text{graphite}} + \text{FeO}^{\text{magma}} \rightarrow \text{CO}^{\text{gas}} + \text{Fe}^{\text{native}}$$

Reaction (13) implies that native iron formation consumes graphite and produces CO, in addition to that supplied by reactions (11) and (12).

We calculate that the critical pressure for native iron saturation to occur is 10 MPa. At higher pressures, the redox state of the magma is not reduced enough for native iron to be stable. This critical pressure mainly depends on magma temperature and to a lesser extent on its total iron content. A hotter and Fe richer magma would crystallize native iron at slightly higher depths (e.g. a magma with 13 wt% total FeO at 1200°C is saturated in native iron at 12 MPa).

Reactions (11-13) can help illustrating what happens to a basaltic magmas ascending through sedimentary sequences containing organic matter-rich layers. When a magma is contaminated deep in a basin with oil or shale, graphite crystallizes for very low amounts of organic matter assimilated (Fig.2). This graphite is then consumed during ascent to degas carbon monoxide, and to produce, at depths of few hundred meters (varying as a function of the density of the sedimentary host rocks), native iron if significant amounts (more than 0.5 wt%) of organic matter are assimilated (Fig.2). If the ascending magma also flows through coal-bearing sediments at shallow depths, coal assimilation is even more likely to furnish the amount of organic matter necessary to reach iron saturation. When the magma becomes saturated in native iron, further addition of organic matter does not increase CO degassing, unless pressure is decreased.

A probable side effect of native iron separation that is not taken into account by our model is sulphide precipitation. Native iron separation decreases the total iron content of the melt and therefore also its FeO content, consequently decreasing the maximum amount of sulphur that it can dissolve (Gaillard and Scaillet, 2009): sulphides are therefore likely to precipitate in
response to native iron precipitation. However, this process would not produce large amounts of sulphides, given the relatively low S content of the magma, and would probably happen after the precipitation of relatively large amounts of native iron.

3.3 Effect of temperature on gas composition

The effect of temperature on the composition of the gas phase was calculated between 1200 and 300°C. The gas phase produced at 1200°C at graphite and iron saturation (0.6 wt% CH added) was then re-equilibrated at lower temperatures: Figure 4 shows its composition at two different final pressures (0.1 and 10 MPa, Fig. 4a and b, respectively). While at magmatic temperatures (>1000 °C) the gas phase is CO-H2-dominated, it is mainly constituted of H2O, CH4 and CO2 when the temperature decreases below 500°C. Such low temperature gases are close in composition to those resulting from thermal metamorphism of country rocks (Svensen et al., 2009).

4. Field evidence of magma-organic matter interactions

The presence of igneous graphite in mafic rocks has been interpreted as an evidence of magma-organic matter interaction in different geological contexts, e.g. at the Karoo Basin, South Africa (Henning et al., 1997), or at Borrowdale in Cumbria, United Kingdom (Ortega et al., 2010). However, the occurrence of native iron in basaltic extrusive or intrusive rocks is probably the clearest evidence of extremely reducing conditions reached during magmatic assimilation of carbonaceous rocks (crystallization of native iron implying oxygen fugacities lower than FMQ-6); three unambiguous cases are presented below.

4.1 Disko Island, Greenland
The first occurrences of native iron in basaltic magmas were found at the island of Disko in central West Greenland (Nordenskjöld, 1872). Since then, disseminated native iron has been described in several basalts at Disko, associated with wustite, carbides and sulphides (Steenstrup, 1884; Pedersen, 1979a,b 1985; Ulff-Møller, 1985). Numerous graphite rich xenoliths have been also found throughout the iron bearing basalts and interpreted as thermally metamorphosed fragments from the underlying bituminous shales on the basis of their mineral assemblages and sedimentary structures (Melson and Switzer, 1966; Pedersen, 1978, 1979b).

Disko lavas belong to the Paleocene volcanic succession related to the North Atlantic mantle plume (Pedersen and Larsen, 2006). They were emplaced in the Nuussuaq Basin (part of the West Greenland Basin), whose Cretaceous and Tertiary sedimentary sequences comprise abundant carbonaceous sediments including coal seams up to 2 m thick (Medenbach and ElGoresy, 1982), which severely contaminated picritic magmas (see Pedersen and Larsen, 2006 for a review). The crustally contaminated rocks occur as dykes, necks, lava flows and tuffs and contain variable amounts of graphite and native iron. At the classic locality of Asuk, a ~100 m thick sequence of sediment-contaminated lavas, mostly iron-bearing, occurs in which baked, party glassy, shale xenoliths are frequent, some still preserving sedimentary bedding (Pedersen, 1979a). Pedersen and Larsen (2006) proposed that the abundance of graphite and native iron in the Disko and Nuussuaq rocks depends on the pressure at which they equilibrated: using thermodynamic estimations of the equilibrium between graphite and a C-H-O gas phase at high temperature and pressure (French and Eugster, 1965; French, 1966), they predicted an equilibration pressure of 0.1 MPa for Disko lavas and 50-100 MPa for the native iron free graphite andesite tuffs from Nuussuaq. This is in a very good agreement with what our calculations predict.

4.2 Bühl, Germany
Native iron-bearing xenoliths have also been observed in Miocene alkali basalts at Bühl near Kassel, in West Germany (Medenbach and ElGoresy, 1982). The xenoliths have been found in a basalt pipe that intruded Tertiary coal seams and have been interpreted as the products of a reduction under near surface conditions (<10 MPa) of manganese-bearing iron minerals contained in Tertiary sediments and coal seams cut by the erupting basalts (Medenbach and ElGoresy, 1982). Carbon from coal has been considered as the reducing agent (Medenbach and ElGoresy, 1982).

4.3 Siberian Platform, Russia

The highest concentrations of native iron in magmatic rocks have been observed in the North and North-west of the Siberian Platform, in several trap intrusions hosted by the Middle Carboniferous-Permian coal-bearing terrigenous deposits (Ryabov and Lapkovsky, 2010). The occurrence and composition of native iron in the different intrusions (Maimecha, Khininda, Khungtukun intrusions, Dzhaltul igneous complex and Ozernaya Mountain intrusion) are very similar. Two types of native iron have been recognized: fine disseminations (0.2-0.5 mm) in dyke-like bodies and nodules (up to 200 kg in weight) concentrated in the upper zone of the intrusions (or of the first intrusive phase in the Dzhaltul complex). Dyke-like bodies are 0.3-3.9 m-thick and contain fragments of sedimentary rocks. The 2-40 m thick nodule-bearing horizons are located within a 40-110 m-thick olivine gabbro zone where they are associated with graphite, carbides and finely dispersed carbonaceous matter. The quantity and size of nodule increase upwards in the ore-bearing horizons. The current depths of native iron mineralizations are between 40 and 600 m. Subsidence of several hundred meters throughout the Mesozoic and Cenozoic has been, however, estimated for the West Siberian Basin (Saunders et al., 2005).
Several lines of evidence suggest that the temperature of ore formation was 750-1200 °C and that at least some of the native iron formed before the crystallization of silicates, e.g. native iron inclusions occur in plagioclase, olivine and pyroxene (Ryabov and Lapkovsky, 2010). Although sulphides are commonly associated with the native iron, the total sulphur content of the mineralization is < 0.2 wt%. Carbon isotopic compositions of graphite and carbides are similar to that of the organic matter (Ryabov and Lapkovsky, 2010). More generally, the occurrence of varying amounts of finely dispersed carbonaceous matter and its carbon isotopic composition suggest that nearly all ore-bearing intrusions in the northwest of the Siberian Platform have been contaminated by carbonaceous matter to different extents (Ryabov et al., 2006; Ryabov and Lapkovsky, 2010). Carbonaceous matter occurs in the form of amygdales in dolerite of explosive breccias, and also as small rounded inclusions in plagioclase (Ryabov et al., 2006; Ryabov and Lapkovsky, 2010). Abundant bitumen-filled amygdules (schungite) in volcanic units of the Noril’sk region and massive reduction of sulphate to sulphide during the formation of the major Ni sulphide deposits of the Norilsk-Talnakh region (Arndt et al., 2005) provide additional evidence of interaction with hydrocarbons.

5. Gas emissions produced by assimilation of organic matter

Native iron-bearing intrusions and lava flows described in the previous section are hosted in, or associated with, coaliferous strata and bituminous shales, suggesting that graphite and native iron saturation was triggered by assimilation of organic matter (mainly coal). Our calculations restrict native iron crystallization to pressures lower than 10 MPa (Fig. 2), i.e. 350-500 m, depending on the density of the overlying sedimentary rocks, however the ingestion of organic matter could have happened also at deeper levels (e.g. where petroleum accumulations occur). The occurrence of native iron constrains the minimum amount of
organic matter assimilated (either CH or CH\textsubscript{2}) to be 0.5 wt % (Fig.2). Moreover, the coexistence of graphite and native iron also constrains the composition of the fluid phase. At 0.1 MPa, the fluid phase in equilibrium with graphite and native iron is CO-dominated, regardless the initial conditions (Fig. 1, Fig. SM1, and Table 2). Figure 5a presents the same results as Figure 1d, with the difference that the composition of the fluid phase is expressed in wt% of the magma. We calculated that graphite and native iron-saturated lavas like those observed at Disko and Bühl degassed 1.3 wt% CO, the other gas components representing less than 0.1 wt% (Table 2). These estimations take into account only the contribution of reactions (11) and (12), without considering the additional production of CO associated to native iron formation (reaction 13) and are hence minima. We also recall that an oxygen-free organic matter was considered in our calculations; the presence of oxygen in the assimilated organic matter would result in a higher degree of assimilation and therefore in a higher CO production. Fluid bubbles hosted in the groundmass glass of native iron-bearing Disko basalts were investigated by heating and cooling microscopic techniques, revealing strongly reduced volatile molecules most likely formed during the magmatic stage (Solovova et al., 2002). The estimated fluid composition at ~1180°C and pressures lower than 10 MPa is CO-H\textsubscript{2} dominated (Solovova et al., 2002).

As native iron was observed not only in basaltic flows, but also in subvolcanic intrusions that were probably emplaced a few hundred meters deep (e.g. in North Siberia, see previous section), we also calculated the composition of the fluid phase in equilibrium with graphite and native iron at 10 MPa (i.e. the maximum depth for native iron crystallization calculated at 1200°C). At 10 MPa, saturation in graphite and native iron is reached for ~0.5 wt% CH or CH\textsubscript{2} addition. Figure 5b and Table 2 show that, although the coexisting fluid phase is still largely CO-dominated, calculated CO contents of the magma are slightly lower (~1.0 wt%), and those of the other components slightly higher (but always <0.1 wt%) than at 0.1 MPa.
Again, these estimations consider only reactions (11) and (12) and not (13). In the absence of robust constraints on the degree of metallisation of iron in the native iron-bearing basalts we cannot correctly estimate the amount of CO produced by reaction (13), which is nevertheless potentially significant. A maximum estimate can be done by considering that all FeO initially present in the basalt (8 wt%) has been transformed into native iron: this would have produced additional 3.1 wt% CO, which then totalizes ~4.2 wt% of the magma (Table 2). Even if this additional contribution is not considered, both Figure 5 and Table 2 clearly show how the ingestion of only 0.6 wt% of organic matter by a basalt strongly enhances magma degassing: the total volatile content of the magma increases from ~0.9 wt% in the absence of assimilation to ~1.5 wt% at 0.1 MPa (Fig. 5a, Table 2), and from ~0.5 wt% to ~1.2 wt% at 10 MPa (Fig. 5b, Table 2).

If the temperature of magmatic gases decreases during ascent after separation from the magma, their composition would change from CO toward H₂O-CH₄-CO₂-dominated (Fig.4b). For instance, at 300°C the calculated gas composition is ~ 46 wt% H₂O, 28 wt% CH₄ and 26 wt% CO₂.

The enhanced magmatic degassing of C species (including CO at high temperatures and CH₄-CO₂ at low temperature) is likely to have environmental implications when the associated magma production rate is high. The emplacement of Disko and Nuussuaq lavas that belong to the North Atlantic Large igneous Province (Storey et al., 1998) could have caused environmental damages, due to the large volumes of magma that were contaminated by organic matter-rich sediments (Pedersen and Larsen, 2006). Lethal hazard for animal life has been invoked at a local scale due to intense CO and CO₂ emanations (Pedersen and Larsen, 2006). A global impact of these gas emissions can also be speculated: magmatic activity in Western Greenland precedes the Paleocene-Eocene hyperthermal (rapid and transient global
warming events) and several authors have suggested that the two events can be related (e.g. Kennett and Stott, 1991; Wignall, 2001).

Siberian magmatism/volcanism is another case in point: it has often been proposed to have caused the end-Permian crisis, and the strong perturbation of the Earth’s carbon cycle, globally marked by a negative carbon isotope excursion (e.g. Wignall, 2001). In particular, the interaction between magmas and carbonaceous sediments has been proposed as the major cause of the environmental impact of the Siberian Trap emplacement (e.g. Svensen et al., 2009). In a companion paper (Iacono-Marziano et al., submitted to EPSL) we quantify the gas production potential of a single intrusive event in coal-bearing sediments at the Siberian Traps and investigate the dispersion of these gases in the atmosphere. We also discuss the potential impact of these gas emissions on the environment, distinguishing short term implications at a local scale from long term ones at the global scale.

The process described and quantified here could concern not only LIP sized events, but also much smaller intrusions/eruptions. Virtually any active volcanic area whose sedimentary substrata comprise layers of organic-rich material can produce important amounts of CO, with potentially dangerous consequences for nearby human beings.

6. Conclusions

The long term volatile cycle is generally described as the exchange between two major reservoirs, the mantle and the atmosphere, where volcanism has the role of conveying volatiles from the mantle to the surface (e.g. Hilton et al., 2002; Wallace, 2005). Recent studies have however revealed that magma-host rock interaction may contribute additional gases, in particular when host rocks are composed of carbonates, evaporites or organic matter (Svensen et al., 2004, 2007, 2009; Iacono-Marziano et al., 2007, 2009; Ganino and Arndt, 2009). In this paper, we show that another crucial effect of magma-host rock interactions is
that they can impact severely on the redox state of the magma and thus that of volcanic gases. Very minor amounts of organic matter (<0.5 wt%) assimilated by the magma decrease its redox state by several log-units. The assimilation of only 0.6wt% of organic matter almost doubles its total volatile content and produces unusual CO-dominated gases. We therefore conclude that while the long-term deep volatile cycle controls the background level of volcanic degassing, occasional interaction between magma and sedimentary host rocks may yield peak values of abnormal gas compositions, representing a potentially major perturbation of the volatile cycles.

References


Acknowledgements

G.I.M. was founded by the European Community’s Seventh Framework Programme (grant agreement n° PIEF-GA-2008-220926). F.G. was supported by the ERC grant number 279097. This work benefited from conversations with Jean-Robert Disnar and Jérémy Jacob. We thank B. Marty for the careful handling of the manuscript, H. Svensen and an anonymous reviewer for their constructive comments that greatly improved the paper.
Table 1. Numerically tested configurations of organic matter assimilations by basalts.

<table>
<thead>
<tr>
<th>N°</th>
<th>Redox conditions a</th>
<th>Gas content b</th>
<th>Volatile contents c</th>
<th>Organic matter d</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>FMQ</td>
<td>0.3 %</td>
<td>0.5 H₂O (wt%)</td>
<td>CH</td>
</tr>
<tr>
<td>2</td>
<td>FMQ</td>
<td>0.3 %</td>
<td>0.2 S (wt%)</td>
<td>CH</td>
</tr>
<tr>
<td>3</td>
<td>FMQ</td>
<td>0.7 %</td>
<td>0.5 CO₂ (wt%)</td>
<td>CH</td>
</tr>
<tr>
<td>4</td>
<td>FMQ</td>
<td>2.1 %</td>
<td>0.2 H₂O (wt%)</td>
<td>CH</td>
</tr>
<tr>
<td>5</td>
<td>FMQ</td>
<td>0.7 %</td>
<td>0.1 S (wt%)</td>
<td>CH</td>
</tr>
<tr>
<td>6</td>
<td>NNO</td>
<td>0.3 %</td>
<td>0.5 CO₂ (wt%)</td>
<td>CH</td>
</tr>
<tr>
<td>7</td>
<td>NNO</td>
<td>0.7 %</td>
<td>0.1 H₂O (wt%)</td>
<td>CH</td>
</tr>
<tr>
<td>8</td>
<td>NNO</td>
<td>0.7 %</td>
<td>0.2 S (wt%)</td>
<td>CH</td>
</tr>
<tr>
<td>9</td>
<td>NNO</td>
<td>0.7 %</td>
<td>1.0 CO₂ (wt%)</td>
<td>CH</td>
</tr>
<tr>
<td>10</td>
<td>FMQ</td>
<td>0.3 %</td>
<td>0.5 H₂O (wt%)</td>
<td>CH₂</td>
</tr>
<tr>
<td>11</td>
<td>FMQ</td>
<td>0.7 %</td>
<td>0.1 S (wt%)</td>
<td>CH₂</td>
</tr>
<tr>
<td>12</td>
<td>NNO</td>
<td>0.3 %</td>
<td>0.5 CO₂ (wt%)</td>
<td>CH₂</td>
</tr>
<tr>
<td>13</td>
<td>NNO</td>
<td>0.7 %</td>
<td>0.1 H₂O (wt%)</td>
<td>CH₂</td>
</tr>
</tbody>
</table>

All calculations have been conducted at 1200°C, with pressure varying from 200 to 0.1 MPa.

a) Initial redox conditions of the basaltic magma at 200 MPa and 1200°C
b) Initial gas content of the basaltic magma at 200 MPa and 1200°C
c) Initial volatile contents (dissolved in the melt + in the form of bubbles) of the basaltic magma at 200 MPa and 1200°C
d) Type of organic matter assimilated

Table 2. Fluid phase composition in equilibrium with graphite and native iron.

<table>
<thead>
<tr>
<th>Emitted species</th>
<th>10 MPa (wt% a)</th>
<th>0.1 MPa (wt% a)</th>
<th>10 MPa without assimil. (wt% b)</th>
<th>0.1 MPa without assimil. (wt% b)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CO*</td>
<td>1.00-1.09</td>
<td>1.46-1.50</td>
<td>0.02</td>
<td>0.03</td>
</tr>
<tr>
<td>CO</td>
<td>3.11</td>
<td>2.22</td>
<td></td>
<td></td>
</tr>
<tr>
<td>CO₂</td>
<td>0.08</td>
<td>0.00</td>
<td>0.35</td>
<td>0.31</td>
</tr>
<tr>
<td>H₂</td>
<td>0.04</td>
<td>0.04-0.07</td>
<td>0.00</td>
<td>0.00</td>
</tr>
<tr>
<td>H₂O</td>
<td>0.03-0.04</td>
<td>0.00</td>
<td>0.07</td>
<td>0.40</td>
</tr>
<tr>
<td>H₂S</td>
<td>0.02-0.03</td>
<td>0.00</td>
<td>0.02</td>
<td>0.03</td>
</tr>
<tr>
<td>CH₄</td>
<td>0.02</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
</tr>
<tr>
<td>S₂</td>
<td>0.00</td>
<td>0.00</td>
<td>0.02</td>
<td>0.04</td>
</tr>
<tr>
<td>SO₂</td>
<td>0.00</td>
<td>0.00</td>
<td>0.03</td>
<td>0.14</td>
</tr>
<tr>
<td>Tot</td>
<td>1.19-4.41</td>
<td>1.50-3.79</td>
<td>0.51</td>
<td>0.95</td>
</tr>
</tbody>
</table>

a) Gas abundances (in wt% of the magma) produced by high temperature assimilation of organic matter. Given ranges account for the variations in initial conditions (initial fO₂ and gas fraction of the magma), type of organic matter assimilated, i.e. CH or CH₂ and amount of organic matter necessary to reach graphite-native iron saturation (see Section 3.1 for explanation).
b) Gas species produced by the degassing of a basaltic magma (initial H₂O, CO₂ and S contents of 0.50, 0.08 and 0.10 wt%, respectively) without assimilation.
* carbon monoxide produced by reaction 13 in the text.
Figure Captions

**Fig. 1**
Gas compositions at 0.1 MPa expressed in mole fraction as a function of the amount of organic matter assimilated. The effects of different initial conditions of oxygen fugacity (NNO or FMQ), gas proportion in the magma (0.3 or 0.7 %) and type of organic matter assimilated (CH or CH$_2$) are shown. Four different configurations listed in Table 1 are shown: #7 (a), #1 (b), #13 (c), #10 (d). Black and red lines indicate the amount of organic matter necessary to reach graphite and native iron saturation, respectively.

**Fig. 2**
Graphite and iron saturation as a function of pressure (and roughly of depth) for CH and CH$_2$ addition. Calculations for CH assimilation were performed using initial fO$_2$ of FMQ and NNO and initial gas fraction of 0.3 and 0.7%. Only the two extreme cases (fO$_2$ = FMQ, gas fraction = 0.3% and fO$_2$ = NNO, gas fraction = 0.7%) are shown for CH$_2$. Variations between the different cases are always within 30%. The presence of graphite alone in a magmatic rock does not clearly indicate the amount of organic matter assimilated by the basaltic magma, because of the strong dependence on the depth of the intrusion emplacement. On the contrary, the coexistence of graphite and native iron precisely constraints the minimum amount of CH or CH$_2$ (0.5-0.8 wt%) assimilated at depths lower than 350-500 m (depending on the density of the sedimentary host rocks).

**Fig. 3**
Variation of the oxygen fugacity of the magma (expressed as logarithmic units relative to the FMQ buffer) as a function of the amount of coal assimilated. Calculations were performed at 1200°C and 10 MPa using pre-assimilation fO$_2$ of FMQ and volatile contents of 0.50 wt% of H$_2$O, 0.10 wt% of S and 0.24 wt% CO$_2$ (corresponding to a gas fraction of 0.3 % at 200 MPa). Red and black lines indicate native iron and graphite saturation, respectively.

**Fig. 4**
Calculated gas compositions as a function of temperature for ~0.6 wt% CH assimilated (required for both graphite and iron saturation). Two different final pressures are shown: 0.1 MPa (a) and 10 MPa (b).
Fig. 5
Gas contents of the magma (expressed in wt%) at 0.1 MPa (a) and 10 MPa (b) as a function of the amount of organic matter assimilated. Calculations were performed using initial fO$_2$ of FMQ and initial gas fraction of 0.3 %. The lines indicate the amount of organic matter necessary to reach graphite and iron saturation and the composition of the coexisting gas phase.
Fig. 1
wt% CH or CH$_2$ added

Graphite saturation

CH addition
- QFM 0.3-0.7% gas
- NNO 0.3-0.7% gas

CH$_2$ addition
- QFM - 0.3% gas
- NNO - 0.7% gas

Native iron saturation

Fig. 2
Fig. 3
Fig. 4
Fig. 5