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Abstract

To understand deep groundwater flow systems and their interaction with CO₂ emanated from magma at depth in a volcanic edifice, deep groundwater samples were collected from hot spring wells in the Aso volcanic area for hydrogen, oxygen and carbon isotope analyses and measurements of the stable carbon isotope ratios and concentrations of dissolved inorganic carbon (DIC). Relations between the stable carbon isotope ratio ($\delta^{13}$C$_{\text{DIC}}$) and DIC concentrations of the sampled waters show that magma-derived CO₂ mixed into the deep groundwater. Furthermore, groundwaters of deeper areas, except samples from fumarolic areas, show higher $\delta^{13}$C$_{\text{DIC}}$ values. The waters’ stable hydrogen and oxygen isotope ratios ($\delta$D and $\delta^{18}$O) reflect the meteoric-water origin of that region’s deep groundwater. A negative correlation was found between the altitude of the well bottom and the altitude of groundwater recharge as calculated using the equation of the recharge water line and $\delta$D value. This applies especially in the Aso-dani area, where deeper groundwater correlates with higher recharge. Groundwater recharged at high altitude has higher $\delta^{13}$C$_{\text{DIC}}$ of than groundwater recharged at low altitude, strongly suggesting that magmatic CO₂ is present to a much greater degree in deeper groundwater. These results indicate that magmatic CO₂ mixes into deeper groundwater flowing nearer the magma conduit or chamber.
Keywords:

Groundwater flow, magmatic CO$_2$, Dissolved inorganic carbon, Stable isotope, Aso volcano,
Mixing of magmatic CO$_2$ into volcano groundwater flow at Aso volcano assessed combining carbon and water stable isotopes

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1. Introduction

Stable carbon isotope studies of groundwater systems at some volcanoes have revealed that deeply derived CO$_2$ mixes into groundwater that contains CO$_2$ derived from soils (Chiodini et al., 2000; Ohsawa et al., 2002; Evans et al., 2002; Caliro et al., 2005). These studies, using datasets of stable carbon isotope ratios and concentrations of dissolved inorganic carbon (DIC), present a robust approaches to determine whether or not magmatic CO$_2$ is dissolved by groundwater. In addition, it is possible to determine recharge elevations of groundwaters using stable hydrogen and oxygen isotopes based on isotopic altitude effects of precipitation (Yasuhara et al., 1993; Kazahaya et al., 1999; Nakamura et al., 2002; Yasuhara et al., 2002). Using this technique, it is possible to investigate groundwater movement in volcanic edifices. Both of these isotopic methods are powerful tools for investigating the groundwater hydrology of volcanoes.

It is expected that the combined use of these two methods, as first demonstrated at Kuju volcano by Yamada et al. (2005), will be effective to deepen our understanding of the input of magmatic CO$_2$ into volcano groundwater flow systems. Although Yamada et al. (2005) studied shallow groundwater flow systems forming cold carbonic springs and bicarbonate-type hot spring, deep groundwater flow systems and their interaction with CO$_2$ from deep-seated magma requires analysis of groundwater from deep wells drilled into the volcanic edifice. As new hot springs for bathing have been developed by deep
drilling in Japan, including at Aso volcano, we have investigated the interaction between
magma CO$_2$ and deep groundwater using hot spring wells at Aso volcano.

In this paper, we first present results of the isotope and DIC analyses of hot spring
waters, and then assess these results to determine the importance of magmatic CO$_2$ in
deep groundwater in the vicinity of Aso volcano. Results suggest a latent mixing process
of magmatic CO$_2$ into the deep groundwater flow system at Aso volcano.
2. Site description and Methods

Aso volcano is an active volcano in Kyushu Island, southwest Japan. This volcano is characterized by a large caldera, extending 18 km east-west and 25 km north-south, where a central cone is located (Fig. 1). The caldera floor is divided into north and south parts called Aso-dani (Aso Valley) and Nango-dani (Nango Valley) respectively. As Aso-dani is covered with lake sediment (Tanaka, 2000), the Aso-dani landform is much flatter than that of Nango-dani, implying a simple stratified structure of the groundwater flow system at the Aso-dani area. Two large rivers, called Kurokawa and Shirakawa, respectively flow on the lowest part of caldera floors at Aso-dani and Nango-dani. Those rivers, which meet within the caldera, flow out from the caldera at the western rim of the caldera.

Surface geothermal activities of Aso volcano are only observed on the central cones. One type of activity is characterized by volcanic gas discharge from a hot, highly acidic crater lake at the first crater of Mt. Naka-dake (e.g. Saito et al., 2008); the other is characterized by fumarolic gas discharges originated from hydrothermal systems at Yunoto and Tarutama areas on the western slope of Mt. Eboshi-dake (NEDO, 1989) (Fig. 1). Many natural hot springs in and around those areas are recognized as steam-heated hot springs and have been used from antiquity. Recently, many “hot springs”
aside from those in the Uchinomaki area (AHL02–AHL05 in Fig. 1) have been developed for bathing by drilling to about 1000 m deep. Although proprietary chemical data of major dissolved constituents of hot spring waters from Onsen-bunsekisho (data tables of hot springs’ water quality that must be kept by Japanese law), formation processes of the hot spring waters, including their relation to magmatic emanation, have not been investigated at Aso volcano.

Hot spring water samples from 23 wells of variable depth were collected to produce datasets of concentrations and stable carbon isotope compositions of dissolved inorganic carbon (DIC and δ^{13}C, respectively), stable isotopic composition of water (δD and δ^{18}O) and of concentrations major ions (Na, K, Mg, Ca, Cl, SO_4, and HCO_3). Cold spring waters were also collected to determine the relation between δD and recharge elevation of hot spring waters following the method described by Kazahaya and Yasuhara (1994). The samples for water isotope analysis were collected in a glass vial to avoid water evaporation. The samples collected in a CO_2 gas-tight bottle, were stored in a refrigerator at 5°C until chemical and isotopic analyses were undertaken. Water temperature and pH were measured in the field. All sampling locations are presented in Fig. 1.

Major ions (Na, K, Mg, Ca, Cl and SO_4) were determined by ion-chromatography (DX-120, Dionex) and HCO_3 by acid titration. The concentrations of DIC were
determined using a CO$_2$-gas electrode (CE-2041; DKK–TOA Corp.) with an ion meter (IM-22P; DKK–TOA Corp.) after all the carbonate species in the sample water (9 ml) were converted into CO$_2$(aq) by the addition of 1 ml of 10% sulfuric acid. The water samples for measurement of $\delta^{13}$C were injected into a glass vial containing phosphoric acid and filled with helium gas. The generated CO$_2$ gas in the vial was transferred to a mass spectrometer (Delta-Plus; Thermo Finnigan) through pre-treatment equipment of carbon stable isotope of DIC (Gas Bench II; Thermo Finnigan). The obtained $\delta^{13}$C is shown using $\delta$ notation as $\%$-deviation from the value of Vienna-PeeDee Belemnite (V-PDB). The oxygen and the hydrogen isotope ratios ($\delta D$ and $\delta^{18}O$) of the water samples were determined using a mass spectrometer (Geo 20-20 model installed at the Stable Isotope Laboratory of IGNS, New Zealand) with the zinc reduction method and CO$_2$ equilibration method. The obtained $\delta D$ and $\delta^{18}O$ are shown using $\delta$ notation as $\%$-deviation from the value of SMOW. The analytical precisions are $\pm 0.5$mg/l for major ions, $\pm 0.3$mmol/l for DIC, $\pm 0.2\%$ for $\delta^{13}$C, $\pm 1.0\%$ for $\delta D$ and $\pm 0.1\%$ for $\delta^{18}O$, respectively. All analytical results are presented in Table 1.
3. Results and Discussion

3.1. Dissolved inorganic carbon

The relationship between $\delta^{13}$C and DIC are shown in Fig. 2. The two lines in Fig. 2 show theoretical mixing curves calculated considering the dissolution of magmatic CO$_2$ by a groundwater which initially contained only biogenic soil CO$_2$. Although the actual mixing process could be more complex, we estimated the simple theoretical mixing curves because the influence of the isotopic fractionation is so small that we can neglect it because of the following reasons: (1) the all saturation indexes for the calcite were less than 1 (Table 1; SI is calculated by use of the chemical and physical data of the water), implying no precipitation of calcite, or precipitation at a very low rate when the saturation index exceeds the value of 1 (Dandurand et al., 1982; Alessandro et al., 2007); (2) we were careful to sample waters degassing. The equation used to compute these theoretical curves is the following:

$$\delta^{13}C_g = \frac{C_0 \times (\delta^{13}C_0 - \delta^{13}C_m)}{C_g} + \delta^{13}C_m$$  \hspace{1cm} (1)$$

where $C_0$ and $C_g$ represent DIC concentrations of the initial value and the value after mixing respectively, and where $\delta^{13}C_0$, $\delta^{13}C_m$, and $\delta^{13}C_g$ denote the stable carbon isotope
Here, the $\delta^{13}\text{C}$ value of a fumarolic CO$_2$ discharged from Tarutama area (-5‰; NEDO, 1989) was used for $\delta^{13}\text{C}_{\text{in}}$ as magmatic gas, and for $\delta^{13}\text{C}_0$ and $C_0$ consistent with groundwater values in equilibrium with soil CO$_2$, (-30‰, 0.1 mmol/L) and (-30‰, 1 mmol/L) were used. The -30‰ is the lowest value of cold spring water in this area (CS01). Almost all data points are shown on these mixing lines or the enclosed area between the two lines. Ohsawa et al. (2002) suggested using the same kind of $\delta^{13}\text{C}_{\text{DIC}}$ versus DIC concentration diagram that magmatic CO$_2$ mixes into soil CO$_2$ dissolved groundwaters having variable DIC concentrations in Unzen volcano, Japan. Chiodini et al. (2000) presented a similar interpretation using a $\delta^{13}\text{C}_{\text{DIC}}$ versus DIC diagram for central Apennine Italy. Consequently, the results obtained in this study demonstrate that all hot spring waters in Aso volcano are contaminated by magmatic CO$_2$ to varying degrees.

The mixing relation showed on Fig. 2 provides an effective index for expressing the degree of magmatic CO$_2$ mixing because $\delta^{13}\text{C}_{\text{DIC}}$ value increases with the amounts of mixed magmatic CO$_2$. In other words, $\delta^{13}\text{C}_{\text{DIC}}$ of hot spring water will be higher if it is contaminated by more magmatic CO$_2$. Figure 3 presents relations between $\delta^{13}\text{C}_{\text{DIC}}$ of hot spring waters and altitudes of hot spring well bottoms, for the Aso-dani area (A), the Nango-dani area (B), and the whole area (C). The data plot of Aso-dani area (Fig. 3A)
portrays a clear increase in $\delta^{13}C_{\text{DIC}}$ value with decreasing altitude of the well bottom, although the data plots of Nango-dani area (Fig. 3B) show no such clear tendency. However, as depicted in Fig. 3C, some plots of the Nango-dani area agree in the tendency of that of Aso-dani area. This tendency suggests a groundwater system forming in which more magmatic CO$_2$ mixes in the deeper groundwater (Fig. 3C). Five samples, hatched in fig 3c. (NHC01, NHC02, NHC06, NHL02, and NHL04 in Fig. 1), show a different behavior. With the exception of NHL02, topographic catchment areas of these hot spring waters include Yunotani and Tarutama fumarolic areas. As described above, these areas are active geothermal areas with fumarolic gas discharges and natural hot springs of steam-heated type. The fumarolic gases in these areas include a magmatic gas component (Ohsawa et al., 1997). It can therefore be inferred that magmatic CO$_2$ rises near the surface around these areas. Hence hot spring waters that do not show the typical correlation between $\delta^{13}C$ and the altitude of the well bottom, as described above, are produced by near surface mixing of ascending magmatic CO$_2$ and shallow groundwater. This “exceptional” mixing process is linked with the well known formation mechanism of volcanic hot spring water of bicarbonate type. This study has shed light on a formation mechanism of hot spring water in the volcanic edifice different from well known processes. Regarding the formation process of the NHL02 hot spring water, we cannot
reach a definite answer because of insufficient data.

3.2. Water isotopes

The δD and δ¹⁸O values of groundwater samples vary from -62.2‰ to -44.2‰ and -9.2‰ to -7.2‰, respectively. Results are shown on the δD versus δ¹⁸O diagram with the field of andesitic magmatic steam (Giggenbach, 1992) and the meteoric water line (δD = 8δ¹⁸O + 10; Craig, 1961) (Fig. 4). All data of groundwater samples are lie along the meteoric water line, indicating that hot spring waters are derived almost entirely from meteoric water, and that magmatic steam does not mix with groundwater.

In the case where groundwater originates from meteoric water, the recharge altitude of that groundwater can be estimated using the recharge water line: a relation between δD values and mean recharge altitude in the studied area (e.g., Yasuhara et al., 1993; Kazahaya and Yasuhara, 1994; Nakamura et al., 2002; Yasuhara et al., 2002). In this study, four cold spring water samples (CS01–CS04) were used to determine the local recharge-water line. The recharge-water line of Aso volcano is determined as H = -56.4 δD -2063, where H expresses the recharge altitude. The obtained relation is presented in Fig. 5. The vertical bars associated with data points show the topographic recharge altitude of each cold spring.
The relation between the δD data of hot spring waters and altitudes of hot spring well bottoms is presented in Fig. 5. A slight positive tendency of the relation is apparent, which suggests that hot spring water recharged at high altitude flows deeper than that at low altitude. To clarify this, we examined the relation between the altitudes of well bottoms and the recharge altitudes of hot spring waters, dividing them into four groups according to their location: central cones at Aso-dani side, central cones at Nago-dani side, Aso-dani to caldera rim, and Nango-dani to caldera rim. As depicted in Fig. 6, negative correlations were found between the altitudes of well bottoms and the recharge altitudes of hot spring waters in every area, although correlation levels differed among areas. Furthermore, it can be inferred from the correlation line’s slopes that the groundwater flow systems forming hot spring waters in Aso-dani areas (central cones at Aso-dani side and Asodani to caldera rim) are more strongly layered than those in Nango-dani areas (central cones at Nago-dani side and Nango-dani to caldera rim). The systematic relations of recharge altitude and flow depth of hot spring waters in Aso-dani areas are consistent with the general conceptualization of deep groundwater flow systems (Ward and Robinson, 1990): the higher elevation at which the groundwater is recharged, the deeper the groundwater flows. In contrast, in Nango-dani areas, the deep groundwaters that are recharged at several altitudes tend to come together into an almost
identical aquifer.

\[ \text{4. Conclusions} \]

In section 3.1 we have shown that at Aso volcano, deeper groundwater has a stronger interaction with magmatic CO\textsubscript{2}, except in the vicinity of the Yunotani and Tarutama areas. On the other hand, in section 3.2, it was shown that deeper groundwater was recharged at higher altitude. This relation is evident in Aso-dani areas of the northern part of Aso caldera, which is not inconsistent with the groundwater circulation in the stratified volcanic deposits (see section 2.). Combining the results obtained from the inspection of carbon and water isotopes are evident that the $\delta^{13}\text{C}_{\text{DIC}}$ of groundwater recharged at high altitude is more positive than that at low altitude (Fig. 7). Figure 7 presents the good relations between recharge altitude and flow depth of hot spring waters for the Aso-dani areas (central cones at Aso-dani side and Asodani to caldera rim). As clearly shown in Fig. 7, the $\delta^{13}\text{C}_{\text{DIC}}$ of groundwater recharged at high altitude is more positive than that infiltrating at low altitude, suggesting a significant contribution of magmatic CO\textsubscript{2} for such deeper groundwater.

From a simple geometrical consideration of a layered groundwater flow system in volcanic edifice, it is readily understood that groundwater recharged at the highest
altitude flows much nearer the magma conduit or chamber than those recharged at low altitudes (Fig. 8). Consequently, the much deeper groundwaters in Aso-dani areas are thought to be affected strongly by mixing of magmatic CO$_2$. For that reason, results of our study suggest that magmatic CO$_2$ was mixed into the volcano groundwater flow system at Aso volcano. This mechanism differs from the well known mixing process on the formation of bicarbonate type hot spring waters observed in the vicinity of the Yunotani and Tarutama areas (see section 3.1.).

It is reasonable to believe that deep groundwaters are influenced by the magmatic emanation at the central cone side of Aso-dani area, although it is not easy to understand why such a system is apparent at Uchinomaki area at the northern end of Asodani to the caldera rim far from the central cones (AHL02–AHL05 in Fig. 1). For clarification of this point, further geophysical explorations in and around the central cones (e.g., Tsutsui and Sudo, 2004; Hase et al., 2005; Kanda et al., 2008) are necessary.
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**Figure and Table legends**

**Figure 1**

Map showing sampling locations at Aso volcano, Japan. Sampling points of hot springs are shown as diamond, triangle, square, and circle symbols, respectively, on the central cone side of Kurokawa River, caldera limb side of Kurokawa River, central cone side of Shirakawa River, and caldera limb side of Shirakawa River. Hexagonal symbols denote cold spring sampling sites.

**Figure 2**

Concentrations of dissolved inorganic carbon (DIC) versus $\delta^{13}C_{DIC}$. Two lines represent theoretical curves calculated by adding magmatic CO$_2$ to initial groundwater dissolved soil CO$_2$. For details, see the text.

**Figure 3**

Relation between $\delta^{13}C_{DIC}$ values and altitude of the well bottom of hot springs. (A), (B), and (C) respectively show data of the Aso-dani area, the Nango-dani area, and the whole area.

**Figure 4**

The $\delta D$-$\delta^{18}O$ plots of study-area groundwater. The diagonally shaded box shows ranges of $\delta D$ and $\delta^{18}O$ values of andesitic magmatic steam (Giggenbach, 1992). The solid line
expresses the meteoric water line: $\delta D = 8 \delta^{18}O + 10$ (Craig, 1961).

Figure 5

The $\delta D$ value versus altitude of the well bottom. The solid line shows the recharge-water line ($H = -56.4\delta D - 2063$, where $H$ expresses the recharge elevation) as estimated using data of four cold springs.

Figure 6

Relations between the well bottom altitude and recharge altitude as calculated using the equation of recharge-water line and $\delta D$ value. Regression lines and correlation factors are shown on each diagram.

Figure 7

Relation between $\delta^{13}C_{\text{DIC}}$ values and recharge altitude as calculated using the equation of recharge-water line and $\delta D$ value in Aso-dani. Solid and broken lines respectively show the regression lines of the central cone side samples and caldera limb side samples.

Figure 8

Conceptual image of groundwater flow and inflow of magmatic CO$_2$ for groundwaters in Aso volcano at the Aso-dani central cone side. Magmatic CO$_2$ escaped from the magma conduit or chamber to volcanic edifice mixes into groundwater recharged at the high altitude area of the central volcanic cone and flowing through in the proximity of the
magma conduit.
Figure 2

The graph illustrates the relationship between δ¹³C (‰) and DIC (mmol/l) for various locations.

- **Magmatic CO₂**
- **Soil CO₂**

Locations and their corresponding δ¹³C values:

- **Hot Spring**
  - Central Cone Side of caldera floor river
  - Caldera Limb Side of caldera floor river

- **Aso-dani Nango-dani**

- **Cold Spring**
Andestic Magmatic Steam

Global Meteoric Water Line

($\delta_D = 8\delta^{18}O + 10$)

Aso-dani Nango-dani
Central Cone Side of caldera floor river
Caldera Limb Side of caldera floor river

Hot Spring
Aso-dani Nango-dani
Cold Spring
Altitude of well bottom (m)

δD (‰)

Recharge water line

$H = -56.48D - 2063$

$|r| = 0.96$

Central Cone Side of caldera floor river

Caldera Limb Side of caldera floor river

Hot Spring

Aso-dani

Nango-dani

Cold Spring
y = -0.79x + 707
|r| = 0.90

y = -0.47x + 1120
|r| = 0.95

y = -0.10x + 1043
|r| = 0.65

y = -0.25x + 933
|r| = 0.86
Figure 7

Recharge Altitude (m) vs. δ¹³C(‰)

- Contribution of magmatic CO₂
- Groundwater movement

- Deep / Shallow
- Altitude

- Triangle: Hot Spring
- Diamond: Aso-dani
- Central Cone Side of caldera floor river
- Caldera Limb Side of caldera floor river

| r | = 0.98
| r | = 0.80
Figure 8

Rain

Groundwater Flow

Magma conduit

Magma chamber

Central Cone of Aso volcano

“Aso-dani side”

Contribution of magmatic CO$_2$

None, Small, Large