

OPEN The emissions of CO₂ and other volatiles from the world's subaerial volcanoes

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Volcanoes are the main pathway to the surface for volatiles that are stored within the Earth. Carbon dioxide (CO₂) is of particular interest because of its potential for climate forcing. Understanding the balance of CO₂ that is transferred from the Earth's surface to the Earth's interior, hinges on accurate quantification of the long-term emissions of volcanic CO₂ to the atmosphere. Here we present an updated evaluation of the world's volcanic CO₂ emissions that takes advantage of recent improvements in satellite-based monitoring of sulfur dioxide, the establishment of ground-based networks for semi-continuous CO2-SO2 gas sensing and a new approach to estimate key volcanic gas parameters based on magma compositions. Our results reveal a global volcanic CO₂ flux of 51.3 \pm 5.7Tg CO₂/y $(11.7 \times 10^{11} \, \text{mol CO}_2/\text{y})$ for non-eruptive degassing and $1.8 \pm 0.9 \, \text{Tg/y}$ for eruptive degassing during the period from 2005 to 2015. While lower than recent estimates, this global volcanic flux implies that a significant proportion of the surface-derived CO₂ subducted into the Earth's mantle is either stored below the arc crust, is efficiently consumed by microbial activity before entering the deeper parts of the subduction system, or becomes recycled into the deep mantle to potentially form diamonds.

Volcanism is the main pathway for the transfer of carbon and other volatiles stored in the deep Earth to the surface^{1,2}. Volcanic carbon dioxide (CO₂) is a key non-anthropogenic regulator of atmospheric CO₂ levels and has, over geologic time scales, affected the evolution of Earth's climate³⁻⁵. Quantifying the volcanic CO₂ flux is an ongoing challenge for the volcano science community and was the primary driver for launching the Deep Earth Carbon Degassing (DCO-DECADE) initiative of the Deep Carbon Observatory. DCO-DECADE was first established in 2011 during IAVCEI Commission of the Chemistry of Volcanic Gases workshop in Kamchatka and has since provided unprecedented global coverage of volcanic gas emission and composition measurements through continuous monitoring and campaign efforts (https://en.wikipedia.org/wiki/ Deep_Earth_Carbon_Degassing_Project).

Here we present the results of this initiative to date, providing a comprehensive assessment of the present-day (2005–2015) global volcanic CO₂ flux to the atmosphere. We combine measured volcanic fluxes of sulfur dioxide (SO₂) with measured or estimated C/S molar ratios to derive volcanic CO₂ fluxes, as initially done by Allard et al.⁶ and Williams, et al. These data are then extrapolated to volcanoes world-wide. Our present update greatly benefits from recent advances in satellite data analysis and a global compilation of annual emissions measured by the Ozone Monitoring Instrument, OMI8, which provides SO₂ flux data for numerous volcanoes simultaneously. This space-based record is complemented by and compared to ground-based SO₂ flux data, particularly from the Network for Observation of Volcanic and Atmospheric Change NOVAC9, to obtain annual budgets of global volcanic SO₂ emissions. At the same time, coverage of C/S ratios from degassing volcanoes has significantly

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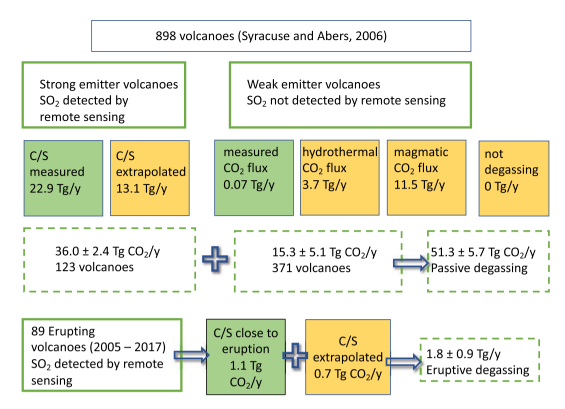


Figure 1. Schematic diagram of how fluxes were calculated and estimated. Green fields indicate measured values, yellow fields indicate estimated or extrapolated values. Numbers are the results as discussed in text with final uncertainties.

improved with the ubiquitous deployment of Multicomponent Gas Analysis Systems (MultiGAS) providing such data over wide temporal and spatial scales¹⁰. We focus primarily on degassing from subaerial volcanic vents but we also include a preliminary evaluation of diffuse soil degassing from the volcanoes' flanks or from active tectonic regions, which has been compiled in the Magmatic Degassing (MaGa) database (www.magadb.net) and just reported in Werner, *et al.*¹¹.

Data and Method

Emissions from subaerial volcanoes with plumes were compiled and evaluated by the DCO-DECADE synthesis group to produce an updated and complete estimation of global volcanic SO_2 and CO_2 fluxes. Our compilation applies to approximately 900 volcanoes, listed by Syracuse and Abers¹². Most of them are located in subduction zones and are documented for subduction-related geophysical parameters, which thus offers the opportunity to compare our results for volcanic output to subduction forcing functions. We also consider volcanoes not listed in Syracuse and Abers but documented for SO_2 or CO_2 fluxes, these being mainly from hot spot and continental rift localities.

The presented dataset includes volcanic emissions from both persistent strong emitters and discrete eruptions, but also provides new estimates for the emissions of weak contributors. Strong emitters are defined as volcanoes that release greater than $0.014~TgSO_2/y$ as detected by satellite. Emissions from erupting volcanoes are identified by using the Global Volcanism Program catalogue for eruptive activity between 2005 and 2017. Weak contributors have SO_2 emissions below OMI's detection limit $(0.014~TgSO_2/y^8)$. The extrapolations to unmeasured volcanoes utilize new data on C/S ratios and a classification into magmatic and hydrothermal categories based on visual observations, volcano databases (predominantly from Smithsonian Institution's Global Volcanism Program, 2013), field reports, and observations made by the authors. Note that while we report and discuss molar C/S ratios, we use mass ratios to compute CO_2 mass fluxes from SO_2 mass fluxes. The summary methodology of computing and extrapolating the fluxes is shown in Fig. 1.

 SO_2 flux from strong emitters and during eruptions. Our compilation of SO_2 flux data covers 11 years from 2005 to 2015 and uses information from long-term monitoring from space and ground, as well as short-term campaign data and reports from the literature. When contemporaneous satellite- and ground-based (long-term or campaign) data exist for the same volcano, we made a critical case-by-case selection of the best and most representative estimate of the annual emission, considering the coverage of each method, specific conditions of measurement and, in certain cases, the separation of the contributions from two neighboring volcanoes that cannot be separated by satellite observations alone. In 28 out 125 cases, ground-based measurements are the only measurements available and are used for the calculations. Therefore, in order to be internally consistent, we used satellite-based measurements where possible and ground-based measurements only when satellite observations

were not available. From the OMI survey the global SO_2 flux from persistent volcanic degassing in the period 2005-2015 is estimated as 23.2 ± 2 Tg SO_2/y (±1 s.d.) for ~90 measured volcanoes (Table S1 and ref. 8). This includes SO_2 from large emitters such as Anatahan (1335 t SO_2/d or 0.5 Tg/y), Bagana (3775 t SO_2/d or 1.4 Tg/y), Aoba (2870 t SO_2/d or 1 Tg/y) and Manam (1484 t SO_2/d or 0.5 Tg/y). Here we complement this satellite-based estimate of decadal emission data with data from the ground-based NOVAC SO_2 flux network 9,13 and from campaign-style measurements for the same 2005-2015 period. The SO_2 flux obtained from ground-based measurements is 6.9 ± 0.8 Tg SO_2/y for 66 volcanoes, some of which were simultaneously observed by OMI. Using the combination approach described above, we account for 125 individual volcanoes in persistent degassing which, altogether, produced 24.9 ± 2.3 Tg SO_3/y in 2005-2015 (Table S1).

The SO_2 flux released during explosive and effusive eruptions is also compiled in Table S1. The time period considered for eruptions extends from 2005 to 2017 and is based on the SO_2 climatology produced by NASA (https://so2.gsfc.nasa.gov/index.html). During this 13-year time period eruptions from 89 volcanoes were observed and emitted a total of 33.3 Tg SO_2 . This corresponds to an average global eruptive flux of 2.6 Tg SO_2 /y, \sim 10 times lower than the annual flux from global persistent degassing. A similar proportion was previously assessed 14,15, despite differences in absolute figures. Therefore, our best estimate for strong volcanic emitters that have been measured either by satellite or by ground-based techniques during both persistent and eruptive degassing is 27 ± 2 Tg SO_2 /y.

CO₂ flux from Strong Emitters. Our study builds on previous work by Werner, et al. 11 and incorporates additional volcanoes with SO₂ flux constraints by OMI and by ground-based techniques. Here we consider the 125 volcanoes which, as described above, had their SO₂ flux measured from space and/or from the ground. Among these, 67 have measured C/S ratios (Table S1). As shown by 16, in assessing volcanic CO₂ fluxes it is critical to use C/S ratios that are representative for the magmatic gas component and not or negligibly affected by sulfur removal during low-temperature hydrothermal processes. In addition to the C/S ratios reported and scrutinized in 10,17, we here report C/S ratios for 10 more volcanoes. These ratios were measured with MultiGAS in crater plumes, thereby minimizing low temperature hydrothermal influences and meeting the requirements proposed by¹⁰. While C/S ratios have been shown to vary with volcanic activity, long-term average ratios correlate with petrologic indicators and therefore are thought to represent magmatic source compositions. Following the approach of 10 , we thus averaged the available gas data for each volcano to obtain a mean \pm 1 SD (1 standard deviation) CO2/SO2 ratios and effectively integrate the "time-averaged" degassing of each volcano. This procedure minimizes the weight of transient phases of either CO₂enriched or, instead, CO₂ depleted volcanic gases that are typically observed either prior to or after eruption, owing to the contrasted solubilities of CO2 and S in magmas (e.g. 18,19) and the earlier degassing of CO₂ during magma ascent and decompression. We note, moreover, that such sequential degassing steps average out in the long-term, as magmas are convectively transported from their deep (CO₂-degassing favored) to near-surface (S-degassing favored) storage zones (eg.²⁰⁻²²). Mafic to intermediate volcanoes typically erupt more than once in a decade. Over timescales of years/decades, therefore, we stress that time-averaged volcanic gas compositions will most closely approach the CO₂/S proportions in the primary undegassed magma and that of the magma source^{23,24}. The error analyses of our results are discussed separately below. For the 67 volcanoes with measured SO₂ fluxes and representative magmatic C/S ratios we obtain a total flux of 22.9 Tg CO₂/y. An additional 8 volcanoes have measured C/S ratios but no SO₂ flux constraints. In order to estimate the CO₂ flux from the remaining volcanoes that have measured SO₂ fluxes (among a total of 125), we need an indirect approach to assess unmeasured magmatic C/S ratios. Aiuppa, et al. 17 have shown that magma's geochemical signatures can be used to predict magmatic C/S ratios for arc volcanoes that are located in a same arc segment. The 35 predicted ratios directly taken from Aiuppa, et al.¹⁷ are shown in Table S1. For 33 of these volcanoes with measured SO₂ flux we infer a cumulative CO₂ flux of 11.1 Tg CO₂/y. To estimate C/S ratios of the remaining 24 volcanoes we again adopt the approach of 17 where unmeasured volcanoes are categorized in Groups 1, 2 and 3 based on their tectonic location. Group 1 and 2 volcanoes have respective average C/S ratios of 1.2 ± 0.5 and 2.4 ± 0.7 (range: 2.1 ± 0.7 to 2.7 ± 0.7) that primarily reflect the C input from subducted slab-derived fluids, whereas Group 3 volcanoes receive substantial additions of carbon from the overlying crust and display higher C/S ratios $(5 \pm 1.5 \text{ on average})^{17}$. Here we distribute volcanoes undocumented for C/S in Groups 1, 2 and 3 according to their tectonic setting and we assign them the above average ratio for each Group. Using this approach, we are able to assign C/S ratios to 61 volcanoes in Table S1. Among these, 23 have a known SO₂ flux and hence provide a total flux of 2.0 Tg CO₂/y. Using this approach, we thus obtain CO₂ fluxes for a total of 123 volcanoes with either measured or estimated C/S ratios. There are only two arc volcanoes, namely Poas in Costa Rica and Iwo-Jima in Japan, with directly reported CO₂ flux (0.038 Tg CO₂/y and 0.1 Tg CO₂/y, respectively)¹¹. Table S1 also includes 4 volcanoes with known SO2 flux but whose C/S ratio cannot be inferred from the above method since they occur in hot spot or continental rift settings. The hot spot volcano, Kilauea, is an important emitter: its ground-based measured CO₂ flux averaged 8,587 ± 7,161 t/day (3.1 ± 2.6 Tg/y) in the period 2005- 2017^{11} , excluding the 2008-2010 phase of heightened eruptive activity. A lower flux of $3,174 \pm 1708$ t CO₂/day $(1.2\pm0.6\ Tg\ CO_2/y)$ was estimated from solely OMI SO_2 fluxes and C/S ratios in 2005–2018. The reported flux error is based on a SD of 0.3 for the C/S ratio. As shown by Werner et al.¹¹, in general there is a quite good agreement between CO₂ fluxes indirectly derived from ground-based and satellite measurements at those volcanoes covered by both methodologies, with overall only 20% higher values for the ground-based measurements. In the case of Kilauea, the flux difference merely results from the use of a constant C/S ratio of 0.92 combined with the entire OMI SO₂ flux record, whereas ground-based sensing actually captured a period of anomalously high CO₂ degassing (high C/S ratio) at the summit in 2005–2007, prior to the 2008 eruption²⁵. In order to be consistent with our approach using OMI and C/S ratios for high emitters, we consider here the OMI-based conservative CO₂ flux of 1.2 Tg CO₂/y at Kilauea.

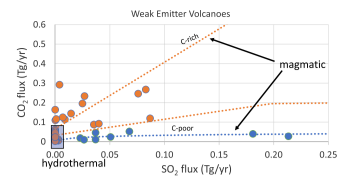


Figure 2. Weak emitter volcanoes SO₂ detected by ground-based campaigns, CO₂ either directly measured or determined using ground-based SO₂ flux measurements and C/S ratios (Table S2).

In summary, we assess a best estimate of 13.1 Tg CO_2/y for the cumulative CO_2 flux from volcanoes whose SO_2 plume emissions are strong enough to be quantified either from space or from the ground, but whose C/S ratios were not measured. Therefore, the overall persistent (non-eruptive) CO_2 flux from the 125 volcanoes discussed above is 36.0 Tg CO_2/y or 8.2×10^{11} mol CO_2/y .

CO₂ flux from explosively erupting volcanoes. In order to estimate the CO₂ flux from explosively erupting volcanoes, it is necessary to know the C/S ratio during such eruptions. But, due to obvious challenges, C/S determinations during such events are scarce (Table S1). One example is the Cotopaxi 2015 eruption, in Ecuador, during which a C/S range of 0.6 to 2.1 was measured²⁶. We note that the inferred ratio of 1.8 we used in Table S1 to compute the eruptive CO₂ flux from this volcano plots well in this range. Although more determinations of eruptive gas ratios are badly needed, we show below that explosive eruptions contribute modestly to global volcanic CO₂ emissions and, so, that even a factor 2 uncertainty in C/S ratio during an explosive event has a relatively minor bearing. For a small number of volcanoes the C/S ratio has been measured just before or after an explosive eruption. Assuming that this ratio is representative for the eruption, we can estimate a CO₂ flux for these volcanoes. 26 out of the 89 volcanoes that erupted between 2005 and 2017 have this datum available (Table S1). For these volcanoes we estimate a total eruptive release of 14.5 Tg CO₂ during the eruptions from 2005–2017, i.e. a mean annual flux of 1.1 Tg CO₂. If we simply extrapolate the average flux per erupting volcano to all (arc and non-arc) of the 89 volcanoes that erupted in 2005-2017, we obtain a total eruptive release of 49.5 Tg CO₂ or a flux of 3.8 Tg CO₂/y. If we instead use Aiuppa et al.'s approach¹⁷ to predict C/S ratios at the 52 undocumented arc volcanoes that erupted in 2005-2017, based on their grouping category, we can estimate an arc eruptive CO₂ flux of 0.7 Tg CO₂/y (9.0 Tg CO₂ in total). The remaining 11 erupting volcanoes are in hot spots or rifts, where we do not know C/S ratios (Table S1). Therefore, our best estimate for eruptive CO₂ emissions in the period 2005–2017 is 23.5 Tg CO₂ or about 1.8 Tg CO₂/y $(4.1 \times 10^{10} \text{ mol CO}_2/\text{y})$. This is nearly identical to the estimate of 1.6 Tg CO_2/y for explosive eruptions and 1.9 Tg CO_2/y for both explosive and effusive eruptions presented in Werner et al. 11. These figures represent only about 5% of the total CO₂ flux from persistently degassing volcanoes, therefore reiterating the early discovery that quiescent volcanic degassing contributes the bulk of volcanic emissions globally^{7,14,15}.

SO₂ and CO₂ fluxes from weak emitters. Out of all approximately 900 volcanoes listed in Syracuse and Abers¹², there are 125 strong volcanic emitters documented by either OMI or ground-based SO₂ flux measurements (Table S1). There are also 19 weaker emitting volcanoes, undetected by OMI, but whose CO₂ flux was determined from either ground-based sensing of SO₂ flux and C/S ratio (Table S1) or direct CO₂ measurement. These 19 volcanoes, not listed in Table S1, emit 1.50 TgCO₂/y. Apart from these 144 volcanoes, there remain 756 volcanoes in Syracuse and Abers's¹² inventory for which we have essentially no data and that require extrapolation in order to constrain the global volcanic volatile fluxes (Fig. 1). All these volcanoes either are not degassing at all or are weak emitters of SO₂ and, therefore, we expect their overall contribution to the global CO₂ budget to be small.

In addition to the 19 weak emitters mentioned above, we select a number of volcanoes in Table S2 that have low SO_2 fluxes ($<0.1\ Tg/y$) but well characterized C/S ratios. 38 of these have both measured CO_2 and SO_2 fluxes, two volcanoes displaying relatively high SO_2 flux (Ebeko, 0.18 Tg/y, and Satsuma Iwojima, 0.21 Tg/y). A plot of CO_2 flux versus SO_2 flux data for the 38 volcanoes (Fig. 2) reveals two main populations of degassing. The first population, which we define as "hydrothermal", is characterized by very low SO_2 flux ($<0.003\ Tg\ SO_2/y$ or $<8\ t\ SO_2/day$) and CO_2 flux up to $<0.1\ Tg/y$ ($<2.75\ t\ CO_2/day$). SO_2 fluxes below the limit of $8\ t\ SO_2/day$ are exceedingly difficult to measure, even with ground-based techniques, and point to a very low magmatic gas supply, which justifies that the corresponding volcanoes be categorized as non-magmatic or hydrothermal degassers. The 275 t CO_2/day value (the highest observed) reflects the fact that low-temperature hydrothermal gas emissions can have exceedingly high C/S ratios 10 . The second population, defined as "magmatic", has an SO_2 flux $>0.003\ Tg/y$ and is composed of two distinct sub-groups (C-rich and C-poor) in terms of CO_2 flux. The low-C group (blue line) has CO_2 fluxes $<0.05\ TgCO_2/y$ irrespective of the SO_2 flux. The high-C group has up to about 3 times as much CO_2 than SO_2 . Two fitting lines are shown for the high-C group, one is a linear fit, the other is an exponential fit. The low-C group is fitted by an exponential fitting curve.

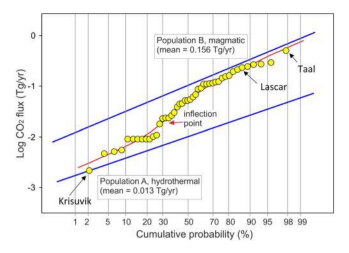


Figure 3. CO₂ GSA method commonly applied to partitioning complex distribution of soil CO₂ flux data in different log normal populations²⁷ applied to weak emitter volcanoes using data from Table S2. Also shown are several volcanoes that span a range of CO₂ fluxes. Two populations are identified: population (**A**) hydrothermal with mean CO₂ flux of 0.013 Tg CO₂/y and (**B**) magmatic with mean CO₂ flux of 0.156 Tg CO₂/y. These populations are used to extrapolate to other weak emitter volcanoes for which no data is available.

While the above evaluation gives us the general understanding that volcanic CO_2 emissions fall into two groups (hydrothermal and magmatic), we can use the following statistical approach to compute the probability that a non-measured volcano has a CO_2 flux value below a determined value that is intrinsic to the population of the weak emitter volcanoes. In order to do this, we use the measured CO_2 flux of all 45 volcanoes in Table S2 to extrapolate to the remaining volcanoes that have no measurements, in order to obtain the most probable total global volcanic CO_2 flux.

For our extrapolation to non-measured weak emitters, we utilize the GSA method commonly applied to partitioning complex distribution of soil CO_2 flux data in different log normal populations²⁷. In this treatment, the first step is to plot the log CO_2 flux values in a logarithmic probability plot (Fig. 3) where a log normal population plots along a straight line while the combined distribution resulting from the overlapping of n log normal populations plot along a curve characterised by n-1 inflection points. The 45 measured log CO_2 flux values (yellow circles in Fig. 3) plot along a curve (red line in Fig. 3) with an inflection point at a cumulative probability of 35% indicating the overlapping of the two log normal populations A (fraction 0.35, mean = -2, $\sigma = 0.33$) and B (fraction 0.65, mean = -0.99, $\sigma = 0.4$). In terms of n log data, the low flux population A (hydrothermal) has a mean of 0.013 Tg/yr (with confidence intervals of 5% and 95% of 0.009 and 0.019 Tg/yr, respectively) while the high flux population B (magmatic) has a mean of 0.156 Tg/yr (with confidence intervals of 5% and 95% of 0.11 and 0.216 Tg/yr, respectively). These results show that hydrothermal fluxes are likely 0.013 Tg CO_2 /y, rather than the 0.1 Tg CO_2 /y estimated visually in Fig. 2, while the magmatic CO_2 flux is likely 0.16 Tg CO_2 /y (431 t CO_2 /day) which cannot be estimated from the data plotted in Fig. 2. We emphasize that with more data for weak emitters, this characterization will improve in certainty.

The next step is to determine which of the 756 volcanoes for which there are no data available are either not emitting gas, or belong to the hydrothermal or magmatic CO₂ flux populations. In order to assess this, we utilized the Global Volcanism Program (GVP) data base, examined recent photographs of these volcanoes to better characterize activity and relied on our combined experiences. The approach that we take is to generally evaluate visually whether an unmeasured volcano is likely to exhibit 'magmatic' or 'hydrothermal' characteristics. A magmatic gas signature would be recognized by a visible fumarolic plume and/or recent (2005–2017) eruptive activity. A hydrothermal gas signature is assigned for volcanoes that have warm, potentially steaming ground, degassing through mud-pools or water, no coherent plume and no large fumaroles. A volcano that is not active, i.e. not degassing at all, lacks all of the above characteristics.

While arguably subjective, this method provides an estimation of the number of degassing volcanoes world-wide that do not have a detectable SO_2 emission and allows for evaluating the type of degassing. We classified 404 volcanoes as not degassing, 278 volcanoes that are degassing hydrothermal gas and 74 volcanoes that are degassing magmatic gas (756 total volcanoes), thus giving a total of 371 degassing volcanoes not detected by OMI or in the NOVAC network (Fig. 1).

Ascribing the mean estimated hydrothermal and magmatic flux values to these sets of volcanoes results in 3.7 Tg CO₂/y degassing from hydrothermal volcanoes and 11.5 Tg CO₂/y from magmatic volcanoes for a total of 15.2 Tg CO₂/y for these unmeasured volcanoes globally. Including the mean of 0.07 Tg CO₂/y of the 19 measured weak emitters (Table S2) results in a total of 15.2 TgCO₂/y or 3.5×10^{11} mol CO₂/y for all weak emitters (Table 1, Fig. 1).

Error assessment. An important consideration when evaluating the fluxes of volcanoes globally is the assessment of the error. Carn *et al.*⁸ provide estimates of the uncertainty in annual mean SO_2 fluxes measured

	Number of	Global CO2 flux Tg/yr				
	volcanoes		5%	95%		
hydrothermal (A)	278	3.7	2.5	5.2		
magmatic (B)	74	11.5	8.1	16.0		
non degassing	404	0	0	0		
TOTAL		15.2	10.6	21.2		

Table 1. CO₂ fluxes from weak emitters (SO₂ not detected by OMI) globally.

From 2005 to 2017	number	CO ₂ flux Tg CO ₂ /y		
Strong emitter volcanoes (SO ₂ flux measured by OMI or ground-based)				
with C/S measured	67	22.9		
with C/S extrapolated based on petrology (Aiuppa et al., in rev)	33	11.1		
with C/S extrapolated no petrology available	23	2.0		
Total flux from strong emitter volcanoes	123	36.0±2.4		
Weak emitter volcanoes				
measured CO ₂	19	0.07		
ascribed hydrothermal	278	3.7		
ascribed magmatic	74	11.5		
non degassing	404	0		
Total flux from weak emitters	775	15.3 ± 5.1		
Erupting volcanoes (2005 to 2017)				
with C/S data available close to eruption	26	1.1		
with C/S extrapolated	52	0.7		
Total flux from erupting volcanoes	78	1.8 ± 0.9		
Total passive degassing		51.3 ± 5.7		
Total CO ₂ flux from the world's subaerial volcanoes		53.1 ± 5.8		

Table 2. Results of CO₂ flux estimates from subaerial volcanoes in the period from 2005 to 2017.

by OMI (~55%), which includes errors associated with retrieved SO₂ columns and with variable plume altitude, wind speed, and SO₂ lifetime. The uncertainty for the ground-based SO₂ fluxes measured by the NOVAC network and selected for this compilation is below 30%. Table S1 provides the fractional uncertainty of the SO₂ flux for each volcano. This uncertainty is estimated at 55% and 30% for space- and ground-based measurements, respectively^{8,9}. This reported uncertainty of \sim 30% for ground-based SO₂ flux measurements with scanning-DOAS follows from the analyses presented in 9,13,28 . The annual means in SO₂ fluxes and their standard errors reported here were calculated from the daily means of SO₂ flux measurements, which in turn were calculated from the individual measurements of flux, when at least five measurements of 'good quality' were available in a given day. An individual SO₂ flux measurement was considered 'good quality' when several criteria were met, namely: low spectral fit error in the derivation of SO₂ column densities, distance to the plume less than 5 km, complete coverage of the scanned plume, use of values calculated by triangulation for plume height, and use of the (then) best available information for plume speed (ECMWF ERA-interim). With these criteria, we consider that our measurements were taken under 'good conditions', as defined in⁹. This means relative uncertainties of 10% for spectroscopy, 10% for measurement geometry, 20% for wind speed, and 20% for atmospheric scattering, all added as independent variables in quadrature. The value of 20% for scattering (or radiative transfer effects), is slightly higher than 'good', to account for possible effects not accounted for in this analysis. Table S1 provides the fractional uncertainty of the SO₂ flux for each volcano. The error of CO₂ flux is strongly dependant on the methods that are used. In our compilation total error estimates for volcanoes with measurements are composed of the errors in the SO₂ flux measured by OMI or by ground-based techniques and the variability of this flux over the 13-year time period, as well as the errors in the C/S ratios used and their variability over the 13-year time period. SO₂ flux variability, C/S variability and CO₂ flux variabilities over the 13-year time period are expressed as a standard deviation in Tables S1 and S2 (one sigma). The errors of our estimates become significantly larger for non-measured volcanoes where extrapolated values are used for C/S ratios based on tectonic setting or when we need to visually categorize volcanoes as degassing or not degassing, and hydrothermal vs. magmatic (Table 2, Fig. 1). For the predicted C/S ratios based on the volcano group and petrologic method, we ascribe a SD of 1.5 for group 3 and 0.8 for Groups 1 and 2, consistent with the SD ascribed in 17. The summary of all volcanic CO₂ fluxes is shown in Table 2 and we report the cumulative SD obtained from the SD of the flux SO₂ flux measurements and the C/S ratios. In order to assess the error of the measured and predicted CO₂ flux from strong emitters, we apply the Monte Carlo method to the summary of the strong emitter volcanoes CO₂ data set. In the simulation, the CO₂ flux for each volcano is set to vary randomly within its mean \pm SD value and the resulting CO₂ fluxes are summed together. This procedure is repeated 100 times, resulting in 100 randomly-generated sums. The total values reported in Table 2 are the

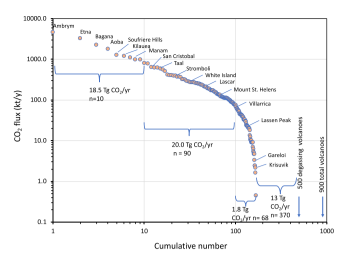


Figure 4. Cumulative number of degassing volcanoes. Values indicate the measured or estimated CO_2 fluxes of the total 500 degassing volcanoes and the total 900 volcanoes. Data show that the top 100 volcanoes emit about 40 Tg CO_2/y . The remaining 68 volcanoes for which we have estimates emit only 1.8 Tg CO_2/y .

ranges (mean \pm 1 SD) of 70% of the random generated sums. Using the Monte Carlo approach, the total CO₂ flux from passively degassing volcanoes then becomes 51.3 \pm 5.7 Tg CO₂/y.

The assessment of errors of the eruptive CO_2 flux requires additional information that is not currently available. For the majority of eruptive SO_2 measurements, no errors are provided²⁹ and the C/S ratios used for computing the CO_2 emissions are not measured during the eruptions but only prior to the eruptions. Given these uncertainties, we define the error on the eruptive CO_2 flux as $\pm 50\%$, emphasizing that it is poorly constrained with present data.

Preliminary considerations for diffuse CO₂ emissions. A preliminary estimate of the volcanic-hydrothermal CO₂ emitted by diffuse emission from soil and lakes is attempted using the published data that are reported in the Magmatic Degassing (MaGa) database at www.magadb.net. The relatively low number of observations with respect to the probable large global number of diffuse degassing structures, hampers the approach of simply summing the catalogued emissions. For this reason, our approach consists of (i) defining the Typical Diffuse CO₂ Emission (TDCE) from diffuse degassing and (ii) estimating a reasonable number of the volcanoes hosting diffuse degassing structures. We note that many papers treating CO₂ degassing from volcanoes also include a non-quantified (but possibly significant) fraction of CO₂ from background-biogenic sources present. For this reason, we limit the computation of a typical diffuse degassing environment to those articles where the deep-volcanic contribution is clearly differentiated from the biogenic source. Since the published total CO₂ fluxes catalogued in MaGa include also the contribution from biogenic sources, TDCE is estimated starting from the 73 cases (Table S3) where the 'volcanic' CO₂ output is explicitly separated from the biogenic one. The resulting CO₂ fluxes, expressed in TgCO₂/y, show a lognormal distribution with a mean of 1.99 and standard deviation of 0.78. Applying a Monte Carlo approach we estimate that the mean of the data (i.e. our best estimation of TDCE) is 0.18 TgCO₂/y (0.09-0.33 TgCO₂/y, 95% confidence interval). Assuming that all the 487 degassing volcanoes host a diffuse degassing structure the total emission from diffuse degassing processes would result in 93 Tg/y (47 Tg/y-174 Tg/y, 95% confidence interval), or 21.22×10^{11} mol/y.

Results and Discussion

Table 2 summarizes the results of the CO_2 flux calculations. Including both strong and weak emitters, the total global flux of CO_2 from passively degassing volcanoes is 51.3 Tg CO_2 /y or 11.7×10^{11} mol CO_2 /y. Our value of 15.3 Tg CO_2 /y from weakly emitting volcanoes represents about 30% of the total CO_2 flux from passively degassing (51.3 Tg CO_2 /y) volcanoes. Notably, our preliminary estimate of diffuse degassing (93 Tg CO_2 /y) is almost twice the total volcanic emission estimate. We stress, however, that this preliminary result could change in the future as the databases of diffuse degassing will include a larger number of data.

Distribution of CO₂ fluxes from global emitters and global fluxes of other volatiles. Figure 4 shows the rank-order distribution of CO₂ emitters globally. The compilation of 12 has approximately 900 volcanoes in total of which about 400 are not degassing, leaving about 500 that have some type of degassing activity. As previously recognized for SO₂^{8,30} and CO₂^{11,17,31}, a few large emitters contribute the vast majority of emissions to the global volcanic gas flux. Based on our compilation, the top ten volcanoes contribute approximately 18.5 Tg CO₂/y or 50% of the 40 Tg CO₂/y emitted by the top 100 volcanoes. This finding is in agreement with the recent estimate of 38.7 ± 2.9 Tg CO₂/y for the 91 most actively degassing volcanoes presented by 17 . This leaves about 400 volcanoes that have some type of degassing and of these we have emission estimates for 68 volcanoes that emit 1.8 Tg CO₂/y. The remaining ~370 volcanoes therefore emit about 13 Tg CO₂/y, considering our total estimate of 51.3 Tg CO₂ for all passively degassing volcanoes. The results also show that, taken together the 80 small emitters

Arc SUM	CO ₂ (Tg/y) strong emitter	CO ₂ (Tg/y) weak emitter	CO ₂ (Tg/y) ascribed mag/hydro	CO ₂ total Tg/y	SO ₂ (Tg/y)	CO ₂ total 10 ⁹ mol/yr	SO ₂ 10 ⁹ mol/yr	H ₂ O (10 ⁹ mol/yr)	HCl (10 ⁹ mol/yr)	H ₂ O/ CO ₂	HCl/ CO ₂
South America	3.16	0.44	2.83	6.44	7.81	146	122	1680	7	11	0.05
CentAm + Mex	3.39	0.52	0.24	4.14	2.05	94	32	3759	14	40	0.15
Alaska + Aleut	0.65	0.64	0.37	1.66	0.72	38	11	1407	17	37	0.44
Kam + Kuriles	1.89	0.28	2.11	4.28	2.18	97	34	5117	25	53	0.26
Japan	1.30	0.22	1.27	2.79	1.53	63	24	18243	72	288	1.14
IBM	0.80	0.00	0.28	1.07	1.07	24	17				
PNG	5.15	0.00	0.25	5.40	3.01	123	47	1958	13	16	0.11
Indonesia	4.11	0.20	3.24	7.55	2.56	172	40	2739	19	16	0.11
Philippines	0.36	0.51	0.24	1.10	0.27	25	4	400	3	16	0.11
Lesser Antilles	1.30	0.00	0.13	1.43	0.47	33	7				
New Zealand	0.38	0.48	0.01	0.87	0.15	20	2				
N and S Vanuatu	7.39	0.00	0.38	7.77	4.50	177	70				
Scotia	0.12	0.00	0.67	0.79	0.15	18	2				
Italy	3.65	0.12	0.00	3.76	0.81	86	13	619	3	7	0.04
Total Arc	34	3	12	49	27	1115	426	35923	173		
Rift & Plume SU	М			•							-
Congo	1.00			1.00	1.29	22.7	20.2	60.56	0.22	2.67	0.01
Tanzania		0.29		0.29		6.6					
Yemen	0.16			0.16	0.04	3.6	0.6				
Ethiopia					0.02					8.04	0.07
Antarctica					0.02						
Total Rift	1.2	0.3		1.4	1.4	32.9	20.7	60.6	0.2		
Plume SUM											
Iceland											
Galapagos		0.14		0.14	0.01	3.3					
Hawaii	1.16			1.16	1.83	26.3	28.6	17.09	0.23	14.75	0.13
Reunion	0.04			0.04	0.09	0.8	1.3				
Total Plume	3.5	0.7		4.2	4.7	96.3	71.4	17.1	0.2		

Table 3. GLobal arc, continental rift and plume passive degassing volatile fluxes based on revised CO₂ fluxes. Notes: Gas ratios are from high temperature arc-by-arc, rift and hot spot fumaroles compilation³². Congo gas ratios are from³⁶; PNG and Philippine ratios are assumed the same as for Indonesia because no high T data is available.

(releasing <100 kt CO₂/y) for which we have data emit a combined 3.1 Tg CO₂/y or almost twice the amount of the erupting volcanoes (1.8 Tg CO₂/y). Therefore, for the most accurate estimates of global emissions, the weak emitters remain at least as significant as the erupting volcanoes. Importantly, weak emitters may become more active and become strong emitters and vice versa, necessitating continued efforts for quantifying volcanic gas emissions from satellite- and ground-based observations.

The new updated volcanic CO_2 flux compilation also requires an updated estimation of the flux of other volatiles using previously published other compositional ratios (i.e. H_2O/CO_2 , HCl/CO_2) of high temperature gases³². Future work should consider updating these high temperature gas data, especially for H_2O but this is beyond the scope of this work. The data compiled on an arc-by-arc basis is shown in Table 3. Note that in Table 3 we include the estimate of the not measured volcanoes. Following the approach described above, we sum up the number of volcanoes for each arc that are inferred to have hydrothermal degassing and assign 3.7 Tg CO_2 /y to these volcanoes (Table 1). Likewise, we sum the volcanoes that are inferred to have magmatic degassing and assign 11.5 Tg CO_2 /y. This amount is then added to the amounts measured for the weak and strong emitter volcanoes for each arc. The data compiled on an arc-by-arc basis is shown in Table 3. For some arcs, we do not have any available gas ratios and cannot calculate H_2O and HCl fluxes.

Hot spot and continental rift passive degassing fluxes are also shown in Table 3. We recognize that there is a potentially large quantity of diffuse CO_2 degassing from the East African Rift^{33,34} and from caldera-hosted hydrothermal systems³⁵, however, the total continental rift volcanic flux is dominated by Nyiragongo and Nyamuragira in the Congo for which gas ratios are also available measured by FTIR in the plume³⁶. Likewise, there is significant hot spot CO_2 flux from low temperature hydrothermal systems³⁷, however, the hot spot volcanic flux is dominated by Kilauea. In the future a more rigorous treatment of hot spot and rift CO_2 degassing, with particular attention to low temperature hydrothermal systems is needed. Overall, our estimates of volcanic degassing from hot spots and rifts are orders of magnitude smaller than volcanic degassing from arcs, entirely consistent with previous assessments^{31,38,39}.

Comparison with other recent volcanic gas flux compilations. Our value of 51.3 ± 5.7 Tg CO₂/y is at the lower end of the recent evaluation of Werner *et al.*¹¹ who estimated 88 ± 21 Tg CO₂/y for persistently passive

degassing volcanoes. Werner *et al.*¹¹ calculated and compiled CO_2 fluxes for a set of 102 volcanoes with direct measurements and 55 OMI-detected volcanoes that emit a total CO_2 flux of 44 and 27 Tg CO_2 /y, respectively. The difference between our estimate and the Werner *et al.*¹¹ estimate is the result of the new treatment of the volcanoes not emitting large quantities of SO_2 . Our new approach considers the type of degassing at each type of volcano and its tectonic setting and based on these parameters, the likely magmatic C/S ratio.

Compared to other recent compilations, our total arc CO_2 flux of 11×10^{11} mol/y is half of Kagoshima *et al.*³⁸ who report $22 \pm 5 \times 10^{11}$ mol/y based on arc ³He fluxes and about 60% of that reported by Fischer³² who reports 19×10^{11} mol/y based on previously reported arc sulfur fluxes and the same high temperature gas ratios. It is also significantly lower than the global arc CO_2 flux of Shinohara⁴⁰ who report 120×10^{11} mol/y based on detailed studies of the Japan arc and extrapolation to global arcs. However, that study also includes diffuse soil degassing, degassing from springs and emissions during eruptions. If we combine our total flux estimate of 11×10^{11} mol/y with our diffuse flux estimate of 21×10^{11} mol/y, we obtain 32×10^{11} mol/y, still a factor of almost 4 lower than the estimate of Shinohara⁴⁰.

Our maximum estimate of 32×10^{11} mol/y for combined diffuse and volcanic CO_2 degassing includes emission from volcanoes in all tectonic settings. Our low arc fluxes have significant implications for evaluating the global deep carbon cycle. The recent compilation of Kelemen and Manning⁴¹ suggest significant recycling or storage of C into the deep mantle or below the arc crust, respectively. Our low volcanic CO_2 emission rates support this idea. Even with the estimated diffuse degassing flux, the total arc CO_2 flux is on the lower end of most previous estimates suggesting that a significant portion of the incoming C delivered by the subducting plates either gets recycled into the deep mantle^{41–43}, is added to the arc crust⁴¹ and may eventually end up below cratons⁴⁴ or is consumed by microbes and/or trapped in precipitated calcite in the forearc before reaching the zones of magma generation⁴⁵. Compared to the most recent CO_2 flux estimates from mid ocean ridges (MOR) of $1.32\times10^{12}\,\mathrm{mol}\,C/y^{46}$, our total flux (volcanic craters and diffuse) of $3.2\times10^{12}\,\mathrm{mol}/y$ is about a factor of 2.5 higher , establishing that subaerial volcanoes are the most significant emitters of volcanic CO_2 to the Earth's surface.

Data availability

All data are publically available in the EarthChem Library data repository at https://doi.org/10.1594/IEDA/111445. All data are also available in the Tables of the manuscript and in the Supplemental Tables S1–S3.

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References

- 1. Dasgupta, R. Ingassing, storage, and outgassing of terrestrial carbon through geologic time. *Reviews in Mineralogy and Geochemistry* 75, 183–220 (2013).
- 2. Dasgupta, R. & Hirschmann, M. The deep carbon cycle and melting in Earth's interior. Earth & Planetary Science Letters 298, 1-13 (2010)
- 3. Berner, R. A. The Phanerozoic Carbon Cycle. (Oxford University Press, 2004).
- Lee, C.-T. A. et al. Continental arc-island arc fluctuations, growth of crustal carbonates, and long-term climate change. Geosphere 9, 21–36, https://doi.org/10.1130/GES00822.00821 (2013).
- 5. Brune, S., Williams, S. E. & Müller, D. Potential links between continental rifting, CO2 degassing and climate change through time. *Nature Geoscience*, https://doi.org/10.1038/s41561-017-0003-6 (2017).
- 6. Allard, P. Eruptive and diffusive emissions of CO2 from Mt Etna. nature 351, 387-391 (1991).
- 7. Williams, S. N., Schaefer, S. J., Calvache, V. M. L. & Lopez, D. Global carbon dioxide emission to the atmosphere by volcanoes. *Geochim. Cosmochim. Acta* **56**, 1765–1770 (1992).
- 8. Carn, S. A., Fioletov, V. E., McLinden, C. A., Li, C. & Krotkov, N. A. A decade of global volcanic SO2 emissions measured from space. Scientific Reports 7, 44095, https://doi.org/10.1038/srep44095 (2017).
- Galle, B. et al. Network for Observation of Volcanic and Atmospheric Change (NOVAC) A global network for volcanic gas monitoring: Network layout and instrument description. J Geophys Res 115, D05304, https://doi.org/10.1029/2009JD011823 (2010).
- 10. Aiuppa, A., Fischer, T. P., Plank, T., Robidoux, P. & Di Napoli, R. Along-arc, inter-arc and act-to-arc variations in volcanic gas CO2/ST ratios reveal dual source of carbon in arc volcanism. *Earth-Science Reviews* 168, 24–47 (2017).
- 11. Werner, *C. et al.* Carbon Dioxide Emissions from Subaerial Volcanic Regions. In *Deep Carbon: Past to Present*. Edited by Beth N. Orcutt, Isabelle Daniel, Rajdeep Dasgupta. Cambridge University Press, https://doi.org/10.1017/9781108677950 (2019).
- 12. Syracuse, E. M. & Abers, G. A. Global compilation of variations in slab depth beneath arc volcanoes and implications. *Geochem. Geophys. Geosyst*, **1219**(7), https://doi.org/10.1029/2005GC001045 (2006).
- 13. Arellano, S. et al. Synoptic analysis of a decade of daily measurements of SO2 emission in the troposphere from volcanoes of the global ground-based Network for Observation of Volcanic and Atmospheric Change (Submitted).
- Andres, R. J. & Kasgnoc, A. D. A time-averaged inventory of subaerial volcanic sulfur emissions. J. Geophys. Res. 103, 25,251–225,261 (1998).
- 15. Berresheim, H. & Jaeschke, W. The contributions of volcanoes to the global atmospheric sulfur budget. *J. Geophys. Res.* **88**, 3732–3740 (1983)
- Aiuppa, A. et al. A CO2-gas precursor to the March 2015 Villarrica volcano eruption. Geochem. Geophys. Geosyst, 18, https://doi. org/10.1002/2017GC006892. (2017).
- 17. Aiuppa, A., Fischer, T. P., Plank, T. & Bani, P. CO2 flux emissions from the Earth's most actively degassing volcanoes, 2005–2015. Scientific Reports 9, 5442, https://doi.org/10.1038/s41598-019-41901-y (2019).
- 18. Moretti, R., Papale, P. & Ottonello, G. A model for the saturation of C-O-H-S fluids in silicate melts. Volcanic degassing. *Geol. Soc. Lond. Spec. Publ.* 213, 81–101 (2003).
- 19. Edmonds, M. New geochemical insights into volcanic degassing. *Philos. Trans. R.Soc. A Math. Phys. Eng. Sci.* **366**(1885), 4559–4579 (2008).
- 20. Aiuppa, A. et al. Forecasting Etna eruptions by real-time observation of volcanic gas composition. Geology 35, 1115-1118 (2007).
- Shinohara, H. Excess degassing from volcanoes and its role on eruptive and intrusive activity. Rev. Geophys. 46, RG4005, https://doi. org/10.1029/2007RG000244 (2008).
- 22. Blundy, J. D., Cashman, K. V., Rust, A. C. & Witham, F. A case for CO2-rich arc magmas. Earth Planet. Sci. Lett. 290, 289-301 (2010).
- 23. Allard, P., Carbonnelle, J., Metrich, N., Loyer, H. & Zettwoog, P. Sulphur output and magma degassing budget of Stromboli volcano. *nature* 368, 326–330 (1994).

- Allard, P. A CO2-rich gas trigger of explosive paroxysms at Stromboli basaltic volcano, Italy. J. Volcanol. Geotherm. Res. 189, 363–374 (2010).
- 25. Poland, M., Miklius, A., Sutton, A. J. & Thornber, C. R. A mantle-driven surge in magma supply to Kīlauea Volcano during 2003–2007. *Nature Geoscience* 5, 295–300, https://doi.org/10.1038/ngeo1426 (2012).
- 26. Hildalgo, S. et al. Evolution of the 2015 Cotopaxi Eruption Revealed by Combined Geochemical and Seismic Observations. Geochem. Geophys. Geosys. 19, 2087–2108. 2010.1029/2018GC007514 (2018).
- 27. Chiodini, G., Cioni, R., Guidi, M., Raco, B. & Marini, L. Soil CO2 flux measurements in volcanic and geothermal areas. Appl. Geochem. 13(5), 543-552 (1998).
- 28. Kern, C. et al. Radiative transfer corrections for accurate spectroscopic measurements of volcanic gas emissions. Bull Volcanol 72, 233–247, 210.1007/s00445-00009-00313-00447 (2010).
- 29. Carn, S. A., Clarisse, L. & Prata, A. J. Multi-decadal satellite measurements of global volcanic degassing. *J. Volcanol. Geotherm. Res.* 311, 99–134 (2015).
- 30. Mori, T. et al. Time averaged SO2 fluxes of subductionzone volcanoes: Example of a 32-year exhaustive survey for Japanese volcanoes. *Journal of Geophysical Research: Atmospheres* 118, 8662–8674 (2013).
- 31. Burton, M. R., Sawyer, G. M. & Granieri, D. Deep carbon emissions from volcanoes. Reviews in Mineralogy and Geochemistry: Carbon in Earth 75, 323–355 (2013).
- 32. Fischer, T. P. Volatile fluxes (H2O, CO2, N2, HCl, HF) from arc volcanoes. Geochemical J. 42, 21-38 (2008).
- 33. Lee, H. et al. Incipient rifting accompanied by the release of subcontinental lithospheric mantle volatiles in the Magadi and Natron basin, East Africa. J. Volcanol. Geotherm. Res. 346, 118–133 (2017).
- 34. Hunt, J. A., Zafu, A., Mather, T. A., Pyle, D. M. & Barry, P. H. Spatially variable CO2 degassing in the Main Ethiopian Rift: Implications for magma storage, volatile transport and rift-related emissions. *G-cubed*, https://doi.org/10.1002/2017GC006975 (2017).
- 35. Robertson, E. et al. Diffuse degassing through continental rift volcanoes: a soil CO2 survey at Longonot Volcano, Kenya. J. Volcanol. Geotherm. Res. Available on-line June 2016 (2016).
- 36. Sawyer, G. M., Carn, S. A., Tsanev, V. I., Oppenheimer, C. & Burton, M. R. Investigation into magma degassing at Nyiragongo volcano, Democratic Republic of the Congo. *Geochem. Geophys. Geosyst.* 9, https://doi.org/10.1029/2007GC001829 (2008).
- 37. Viveiros, F. et al. Soil CO2 emissions at Furnas volcano, São Miguel Island, Azores archipelago: Volcano monitoring perspectives, geomorphologic studies, and land use planning application. J. Geophys. Res. 115, B12208, B12208, 12210.11029/12010JB007555 (2010)
- 38. Kagoshima, T. et al. Sulfur geodynamic cycle. Scientific Reports 5, 8330, https://doi.org/10.1038/srep08330 (2015).
- 39. Marty, B. & Tolstikhin, I. N. CO2 fluxes from mid-ocean ridges, arcs and plumes. Chem. Geol. 145, 233-248 (1998).
- 40. Shinohara, H. Volatile fluxes from subduction zone volcanoes: insights from a detailed evaluation of the fluxes from volcanoes in Japan. J. Volcanol. Geotherm. Res. 268, 46–63 (2013).
- 41. Kelemen, P. B. & Manning, C. R. Reevaluating carbon fluxes in subduction zones, what goes down, mostly comes up. *Proc. Nat. Acad. Sci.* 112, E3997–E4006 (2015).
- 42. Hilton, D. R., Fischer, T. P. & Marty, B. Noble gases in subduction zones and volatile recycling. In: Porcelli, D., Ballentine, C. & Wieler, R. (eds), MSA Special Volume: Noble Gases in Geochemistry and Cosmochemistry. 47, 319–362 (2002).
- 43. Shaw, A. M., Hilton, D. R., Fischer, T. P., Walker, J. A. & Álvarado, G. Contrasting He-C relationships in Nicaragua and Costa Rica: insights into C cycling through subduction zones. *Earth Planet. Sci. Let.* 214, 499–513 (2003).
- 44. Foley, S. F. & Fischer, T. P. An essential role for continental rifts and lithosphere in the deep carbon cycle. *Nature Geoscience* 10, 897–902, https://doi.org/810.1038/s41561-41017-40002-41567, https://doi.org/10.1038/s41561-017-0002-7 (2017).
- 45. Barry, P. H. et al. Forearc carbon sink reduces long-term volatile recycling into the mantle. nature 568, 487-492 (2019).
- Hauri, E. et al. Carbon in the Convecting Mantle. In Deep Carbon: Past to Present. Edited by Beth N. Orcutt, Isabelle Daniel, Rajdeep Dasgupta. Cambridge University Press, https://doi.org/10.1017/9781108677950 (2019).

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Author contributions

All authors worked on compilation and interpretation of the presented data. T. Fischer wrote the first draft and lead the writing of the paper. S. Arellano, A. Aiuppa, T. Lopez, H. Shinohara, C. Werner, C. Cardellini, P. Kelly and P. Allard contributed to writing of the manuscript. S. Arellano and B. Galle provided NOVAC SO₂ flux data and interpretations, S. Carn provided unpublished satellite data and provided insights into OMI data, P. Allard compiled and interpreted the eruption data, G. Chiodini and C. Cardellini provided interpretation of the diffuse degassing data and compiled the diffuse degassing data, G. Chiodini wrote the section on diffuse degassing.

Competing interests

The authors declare no competing interests.

Additional information

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