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# Decarbonation of subducting carbonate-bearing sediments and basalts of altered oceanic crust: Insights into recycling of CO<sub>2</sub> through volcanic arcs



Fabio Arzilli<sup>a,b,\*</sup>, Mike Burton<sup>b</sup>, Giuseppe La Spina<sup>c</sup>, Colin G. Macpherson<sup>d</sup>, Peter E. van Keken<sup>e</sup>, Jamie McCann<sup>b</sup>

<sup>a</sup> School of Science and Technology, Geology Division, University of Camerino, Camerino, Italy

<sup>b</sup> Department of Earth and Environmental Sciences, University of Manchester, Manchester, UK

<sup>c</sup> Istituto Nazionale di Geofisica e Vulcanologia-Osservatorio Etneo, Sezione di Catania, Catania, Italy

<sup>d</sup> Department of Earth Sciences, University of Durham, Durham, UK

<sup>e</sup> Earth and Planets Laboratory, Carnegie Institution for Science, Washington, D.C., USA

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# ABSTRACT

The geological carbon cycle has played a key role in controlling climate throughout Earth's history. For the last  $\sim$ 3 billion years plate tectonics has driven subduction. Subducted slabs have transported CO<sub>2</sub> from the lithosphere, hydrosphere, and atmosphere into the Earth, from where it may be released back to the surface through processes such as arc volcanism or can be stored in the deep interior over geological time. Carbonate-bearing sediments and basalts of altered oceanic crust are the primary media by which carbon is subducted. Therefore, quantifying the depth and amount of CO<sub>2</sub> released from different carbonate-bearing lithologies during subduction is fundamental to understanding whether CO<sub>2</sub> is recycled through arc volcanism or buried in the mantle. The magnitude of CO<sub>2</sub> released from subducting slabs at fore- and sub-arc depths is controlled by processes including ocean crust alteration (i.e., carbonation), metamorphic decarbonation, carbonate dissolution and slab-melting. However, the relative contribution of these processes to overall slab decarbonation is still debated, and will be complex given the variety of sedimentary lithologies and subduction geodynamics. Here, we present a global arc-by-arc lithology-specific analysis of the magnitude of slab CO2 released purely by metamorphic decarbonation of carbonate-bearing sediment and basalt during subduction of altered oceanic crust, using a thermodynamically rigorous model. We find that metamorphic decarbonation is highly efficient in low carbonate sediments, such as carbonated clay, and in carbonated basalts of altered oceanic crust, causing all of their CO<sub>2</sub> to be removed. Sediments with medium and higher carbonate contents, such as chalk and limestone, are only partially decarbonated, but the combination of metamorphic decarbonation and carbonate dissolution promotes efficient carbon loss. Together they can explain observed magmatic CO2 emissions in carbonate-rich arcs. Warm slabs, such as Mexico and Cascadia, produce complete metamorphic decarbonation of carbonate minerals beneath fore-arcs. Under more common cold and intermediate thermal regimes metamorphic decarbonation of carbonate minerals occurs at depths between  $\sim$ 80 and 170 km ( $\sim$ 2.3 to 5.5 GPa) promoting CO<sub>2</sub> input into the mantle sources of volcanic arcs. Overall, our results demonstrate that sub-arc decarbonation is typically considered an important potential source of slab-derived CO<sub>2</sub>, which needs to be considered together with carbonate dissolution to explain observed volcanic CO<sub>2</sub> emissions. In many arcs the modelled CO<sub>2</sub> flux from sediment and basalts of altered oceanic crust into the wedge exceeds the observed CO<sub>2</sub> output suggesting that the mantle wedge and arc lithosphere may sequester some CO<sub>2</sub>.

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### 1. Introduction

E-mail address: fabio.arzilli@unicam.it (F. Arzilli).

The geological carbon cycle has played a critical role over Earth's history in controlling climate. The current anthropogenic flux of carbon is two orders of magnitude higher than that of

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<sup>\*</sup> Corresponding author at: School of Science and Technology, Geology Division, University of Camerino, Camerino, Italy.

volcanic output (Friedlingstein et al., 2010; Burton et al., 2013; Aiuppa et al., 2019), however, over most of Earth's history the carbon flux to the atmosphere has been dominated by volcanic CO<sub>2</sub> emissions (Wong et al., 2019). Recent studies (e.g., Aiuppa et al., 2019; Werner et al., 2019) show that arc volcanoes are important CO<sub>2</sub> emitters, but the origin of volcanic arc CO<sub>2</sub> is still debated. For these reasons, the deep carbon cycle linked to subduction zones and arc volcanism has been intensely investigated during the past 20 years (e.g., Kerrick and Connolly, 2001a; Dasgupta and Hirschmann, 2010; Johnston et al., 2011; Dasgupta, 2013; Ague and Nicolescu, 2014; Kelemen and Manning, 2015; Aiuppa et al., 2017; Plank and Manning, 2019; Stewart and Ague, 2020; Farsang et al., 2021). Subducting slabs transport carbon from the Earth's surface towards the mantle, and multiple processes can mobilise carbon during subduction, providing a source of volatiles to the fore-arc and mantle wedge, or it may be added to the mantle at greater depths, for example by forming diamond. Subducted inorganic carbon (e.g., CaCO<sub>3</sub>) and reduced organic carbon reside within subducted sediments (e.g., Clift, 2017), altered oceanic crust (AOC), and serpentinized mantle of the slab. Of these, inorganic carbon is globally the predominant carbon-bearing material in subducting sediments (Clift, 2017; Dutkiewicz et al., 2019; Plank and Manning, 2019) and in the upper basaltic portion of the AOC (Kelemen and Manning, 2015; Gorce et al., 2019; Merdith et al., 2019; Li et al., 2019). The dominant form of carbon in the basalts of the AOC is CaCO<sub>3</sub> precipitated in veins as calcite and aragonite. In rare cases carbonated serpentinite can form when mantle peridotite is exposed to seawater (Kelemen et al., 2011), but the overall contribution of carbonated peridotite in most global flux estimates is low (Alt et al., 2013; Plank and Manning, 2019). Therefore, the present understanding is that subducting sediments and AOC basalts comprise the majority of the subducted inorganic carbon (e.g., Kelemen and Manning, 2015; Clift, 2017; Dutkiewicz et al., 2019; Plank and Manning, 2019), that dominates global input fluxes in the mantle wedge. Since subducted carbon can return to the atmosphere as CO<sub>2</sub> through arc volcanism, the fate of sedimentary and AOC carbon during subduction is a key factor in several global geochemical cycles including CO2 emissions from arc volcanoes, atmospheric CO<sub>2</sub> concentrations in Earth's history, and climate (Fischer et al., 2019; Hilton et al., 2002). However, the balance between the amount of subducted CO<sub>2</sub> returned to the surface through arc magmatism and CO<sub>2</sub> returned to the deep mantle remains a matter of debate (e.g., Kelemen and Manning, 2015; Plank and Manning, 2019).

Several processes control the mobilization of carbon out of subducting slabs towards the mantle wedge at fore and sub-arc depths, including: metamorphic decarbonation (e.g., Gorman et al., 2006; Cook-Kollars et al., 2014), carbonate dissolution (e.g., Frezzotti et al., 2011; Ague and Nicolescu, 2014; Kelemen and Manning, 2015; Farsang et al., 2021), partial melting of the AOC (e.g., Poli, 2015; Martin and Hermann, 2018), and diapirism of sediments intruding into the mantle wedge (Marschall and Schumacher, 2012; Kelemen and Manning, 2015; Chen et al., 2021). Prior molecular modelling studies have suggested that only modest amounts of carbon can be released from subducting slabs through metamorphic decarbonation reactions at depths <70 km (i.e. below fore-arcs; Connolly, 2005; Gorman et al., 2006; Kerrick and Connolly, 2001a,b). These findings concur with experiments indicating that carbonate minerals subducted within AOC basalts can be transported to pressures >5.5 GPa (mantle depths of >170 km) without decarbonation (Poli et al., 2009), in turn suggesting only a minor role for the recycling of CO<sub>2</sub> through arc volcanism. However, this conclusion is difficult to reconcile with voluminous CO<sub>2</sub> released from arc volcanoes (Marty and Tolstikhin, 1998; Sano and Williams, 1996; Wallace, 2005; Burton et al., 2013; Aiuppa et al., 2017, 2019; Wong et al., 2019; Werner et al., 2019) and observations that infer CO<sub>2</sub> contents are commonly elevated in arc magma (Blundy et al., 2010). Recently, several studies suggest that arc magmatism transports substantial volumes of carbon out of mantle wedges (Kelemen and Manning, 2015; Aiuppa et al., 2017; Plank and Manning, 2019). However, the absolute flux of carbon through any arc will depend on the quantity and lithology of subducted sediment, the original amount of CO<sub>2</sub> present in the mantle wedge, and also any potential addition of carbon to the magma during its ascent (Hilton et al., 2002; Macpherson et al., 2010; Troll et al., 2012; Aiuppa et al., 2017; Mason et al., 2017; Aiuppa et al., 2019). The potential involvement of arc crust in the latter case further complicates efforts to determine the location and extent to which subducted carbon is expelled from the slab (Kelemen and Manning, 2015), in turn creating large uncertainties in our understanding of the carbon cycle on Earth.

Thermodynamic modelling of mineral stability improves our understanding of the fate of carbon and other volatiles over a range of conditions within subducting slabs that are difficult to observe directly. Previous models have introduced increasing complexity and drawn on an ever-expanding evidence base regarding carbon behaviour in natural and experimental systems (Kerrick and Connolly, 2001a,b; Connolly, 2005; Gorman et al., 2006; Galvez et al., 2016; Tian et al., 2019a,b). Closed-system models predict negligible metamorphic decarbonation under relatively water-poor conditions (Kerrick and Connolly, 2001a,b). Open-system models do indicate modest metamorphic decarbonation of carbonate minerals when H<sub>2</sub>O-rich fluids, derived from elsewhere in the slab, promote release of CO<sub>2</sub> into both fore-arc and mantle wedge while retaining substantial carbonate, that is transported into the mantle beyond sub-arc depths. For example, Gorman et al. (2006) modelled infiltration of subducted sediment by a slab-derived H<sub>2</sub>O flux that decreased from the fore-arc to low levels beneath the arc (80-170 km;  $\sim$ 2.3–5.5 GPa). The open-system model, developed by Connolly (2005), used an intermediate thermal regime and showed that dehydration commences at 96 km depth, producing a voluminous source of water-rich fluid from the slab between 136 and 156 km depth. Over this depth range the CO<sub>2</sub> concentration in fluids increased, favouring partial decarbonation at sub-arc depths such that 40-90% of the total carbonate was retained beyond sub-arc depths. A recent thermodynamic model predicts that the solubility of carbon in fluids increases with temperature at depths between  $\sim$ 70 and 100 km (Galvez et al., 2016). The extent to which opensystem behaviour couples with the evolving metamorphic assemblage to encourage or inhibit slab decarbonation remains an open and important question, which we aim to address with this work.

Globally, slabs encounter strongly varying thermal conditions during subduction, which should cause the efficiency of carbon remobilization in subduction zones to vary greatly. To date, there has not been a global systematic study of metamorphic decarbonation during subduction that considers the composition of the subducting carbonate-bearing lithologies (Plank and Langmuir, 1998; Plank, 2014; Clift, 2017) within the context of the unique thermal structure and H<sub>2</sub>O fluxes released from each slab (Syracuse et al., 2010; van Keken et al., 2011). We have developed a Perple\_X (Connolly, 2005) based approach to simulate metamorphic reactions to investigate metamorphic decarbonation efficiency in subducting lithologies (sediments and basalts) along the P-T path of 33 different arc sectors (Fig. 1), that represent the majority of subduction zones on Earth. A range of natural carbonate-bearing lithologies was investigated in order to represent the potential subducting carbonate-bearing sediments and AOC basalts (upper basaltic portion of the oceanic crust) in the modern subduction zones (Fig. 1; Supplementary Table 1). We compare the CO<sub>2</sub> flux produced by metamorphic decarbonation of carbonate-bearing sediments and AOC basalts for each subduction zone with CO<sub>2</sub> fluxes emitted from arc volcanism (Aiuppa et al., 2019; Werner et al., 2019) in



Fig. 1. Global map illustrating the location of the subduction zones considered in this study which have the most active degassing arc volcanoes. The names of the subduction zones have two colours that indicate the presence of carbonate-poor lithologies (cyan) and carbonate-rich lithologies (yellow). CASZ refers to the Central American subduction zones, which includes Guatemala/El Salvador, Nicaragua and Costa Rica. (For interpretation of the colours in the figure(s), the reader is referred to the web version of this article.)

order to quantify the potential contribution of metamorphic decarbonation to the deep carbon cycle on Earth. We also compare the  $CO_2$  flux produced by metamorphic decarbonation with that produced from carbonate dissolution (Farsang et al., 2021).

# 2. Methods

### 2.1. Metamorphic decarbonation and fluid flow

Carbonate mineral stability is influenced by the presence of  $H_2O$  and  $CO_2$  in subducting rock (Connolly, 2005; Kerrick and Connolly, 2001a). We combine Perple\_X (Connolly, 2005) software with a custom-designed algorithm to model incongruent metamorphic decarbonation reactions that lead to the stabilization of calcsilicates, in which only  $CO_2$  is transferred to the fluid and minerals are insoluble, as in a model reaction such as:

# aragonite + quartz = wollastonite + $CO_2$

The metamorphic decarbonation of subducting carbonate-bearing sediments and carbonated AOC basalts (upper basaltic portion of the oceanic crust) is investigated through an open-system model, in which the volatiles (H<sub>2</sub>O and CO<sub>2</sub>) are considered as thermodynamic components. We simulate pervasive water infiltration, while allowing removal of CO<sub>2</sub> generated by metamorphic reactions. In this model, we consider the H<sub>2</sub>O released from underlying oceanic crust and hydrated mantle lithosphere, in agreement with evidence from previous studies (Ulmer and Trommsdorff, 1995; Schmidt and Poli, 1998). The CO<sub>2</sub> fluxes derived from sediments and AOC basalts through metamorphic decarbonation are calculated separately and then combined. We therefore use two separate models for each margin. Our model simulates a scenario of H<sub>2</sub>O infiltration in which grain boundary-fluid interaction dominates simulating pervasive water flow (Fig. 2). The flux of H<sub>2</sub>O released at each pressure step of 5 MPa from the serpentinized mantle and the hydrated igneous oceanic crust ( $\sim$ 5 km of gabbros and  $\sim$ 1.5 km of basaltic dikes), calculated from van Keken et al. (2011), is transferred to the AOC basalts with a transport rate of 0.25 m yr<sup>-1</sup> (Wilson et al., 2014). At each pressure step, CO<sub>2</sub> is removed from the system if metamorphic decarbonation occurs and equilibrium is reached (Fig. 2). When stabilization of carbonate minerals is achieved, the quantity of CO<sub>2</sub> in the bulk composition is transferred at the next P-T point (Fig. 2). Metamorphic decarbonation in sediments is calculated with the same approach used for AOC basalts (Fig. 2). The flux of H<sub>2</sub>O released at each pressure step of 5 MPa from the serpentinized mantle, hydrated igneous oceanic crust and hydrated sediments, calculated from van Keken et al. (2011), is transferred to the carbonate-bearing sediments considered in this study for each slab with a transport rate of 0.25 m  $yr^{-1}$  (Wilson et al., 2014). We use the amount of water predicted by van Keken et al. (2011) for two different purposes. This approach is valid only if a) the decarbonation at lower levels does not affect the H<sub>2</sub>O release itself and b) the deep decarbonation does not significantly affect the water composition. Since the main H<sub>2</sub>O release in most subduction zones is from the serpentinite and the H<sub>2</sub>O does not interact significantly with the carbon-poor gabbros, we satisfy a). The scenario b) was tested and only minor differences were found.

# 2.2. Subducted sediment and basaltic compositions

The sampling of sedimentary covers and upper oceanic crust offers a direct window into what rocks may eventually subduct and degas (Plank and Langmuir, 1998; Plank, 2014; Clift, 2017).



**Fig. 2.** Schematic depiction of modelling approach. Subduction is modelled by the stepwise variation of pressure and temperature along a path prescribed by a selected thermal model (Syracuse et al., 2010; van Keken et al., 2011). Metamorphic decarbonation is investigated within AOC basalts and carbonate-bearing sediments. Two separate models are run for each margin, therefore, the CO<sub>2</sub> fluxes derived from sediment and AOC basalt through metamorphic decarbonation are calculated separately and summed after. H<sub>2</sub>O infiltration between grains is simulated in our model. The flux of H<sub>2</sub>O released at each pressure step of 5 MPa from the serpentinized mantle and the oceanic crust (van Keken et al., 2011), is transferred to the AOC basalts with a transport rate of 0.25 m yr<sup>-1</sup> (Wilson et al., 2014). CO<sub>2</sub> is removed from the system when metamorphic decarbonation occurs at each iteration. After carbonate minerals are stabilized, the CO<sub>2</sub> of the bulk composition is transferred at the next P-T condition. The flux of H<sub>2</sub>O released from the AOC basalts overy 5 MPa is transferred to the sediments. Metamorphic decarbonation is investigated within the sediments every 5 MPa using the P-T paths of each subduction zone (Syracuse et al., 2010); van Keken et al., 2011).

The amount of inorganic carbon can be heterogeneous and vary widely in composition within subducting sediments (Plank and Langmuir, 1998; Plank, 2014; Clift, 2017; Dutkiewicz et al., 2019). The sedimentary veneer may also have variable thickness along the trench causing heterogeneity in the amount of carbon that can be transported to depth. We modelled the subduction of the most representative carbonate-bearing lithology observed at the trench of each subduction zone investigated (Plank and Langmuir, 1998; Plank, 2014; Clift, 2017) (Supplementary Table 1). The carbonatepoor lithologies are typically represented by siliceous ooze and carbonated clay ( $< 5 \text{ wt.}\% \text{ CO}_2$  in the bulk composition), whereas carbonate-rich lithologies are represented by calcareous nannofossil ooze, chalk, and limestones with 12-34 wt.% CO<sub>2</sub> in the bulk composition (Plank and Langmuir, 1998; Plank, 2014; Clift, 2017). Terrigenous sediments are excluded in this model, as they can contain little carbonate with more abundant organic carbon. Furthermore, terrigenous sediments are mostly scraped off the slab in the accretionary prism (House et al., 2019).

AOC inorganic carbon is predominant in the upper basaltic portion (300 m of thickness) of the altered oceanic crust (Kelemen and Manning, 2015; Gorce et al., 2019; Merdith et al., 2019; Li et al., 2019). We modelled stable mineral assemblages and metamorphic decarbonation along each subduction geotherm, using a typical carbonated AOC basalt (Staudigel et al., 1989) (Supplementary Table 1).

#### 2.3. Subduction thermal regimes and computational method

In this study we use two-dimensional thermal modelling developed by Syracuse et al. (2010) and van Keken et al. (2011), and specifically their D80 case, in order to investigate the effect of the thermal regime of 33 slabs on metamorphic reactions in subducting lithologies. These arc regions were selected as they have wellconstrained quantifications of magmatic gas emissions (Aiuppa et al., 2019; Werner et al., 2019). In the D80 case, the slab couples to the overriding mantle wedge at 80 km which leads to formation of a cold corner and a reduction of convection in the mantle wedge directly above the shallow slab (Abers et al., 2017, 2020). We focused specifically on the P-T paths of the sedimentary and AOC basaltic layers as used by van Keken et al. (2011).

Thermodynamic modelling was computed employing the Gibbs free-energy minimization approach using the software Perple\_X (version 6.7.5; Connolly, 2005, 2009) and the thermodynamic database reported from Holland and Powell (2011). The Pitzer & Sterner fluid equation of state (PSEoS) (Pitzer and Sterner, 1994; Holland and Powell, 2003) was used to model the thermodynamic behaviour of H<sub>2</sub>O-CO<sub>2</sub> fluids. The solid solution models include carbonate minerals (Do; Holland and Powell, 1998), magnesite (Mag; Holland and Powell, 1998) clinopyroxene (Cpx; Holland and Powell, 1996), garnet (Grt; Holland and Powell, 1998), white mica (Phe; Auzanneau et al., 2010; Coggon and Holland, 2002), amphibole (Amp; Dale et al., 2005), ternary feldspar (Fuhram and Lindsley et al., 1988), K-feldspars (Kfs; Waldbaum and Thompson, 1968), biotite (Bio; Tajcmanová et al., 2009), and chlorite (Chl; Holland et al., 1998).

The numerical solutions have been computed using the Perple\_X program MEEMUM (Connolly, 2005, 2009), which calculates free energy minimization as a function of specific environmental and compositional parameters and allows us to calculate stable assemblage at particular P-T conditions. This program has been used in combination with a custom-designed algorithm to model an open system.

# 2.4. CO<sub>2</sub> flux estimates

Estimates of CO<sub>2</sub> fluxes were obtained using the thickness  $(t_s)$  (Plank and Langmuir, 1998; Plank, 2014; Clift, 2017) and density  $(\rho_s)$  of sediment sequences entering each trench, and the speed of subduction  $(u_s)$  (van Keken et al., 2011). In our simulations, we consider a pressure step  $(\Delta P)$  of 5 MPa, therefore, assuming a lithostatic pressure, the volume of sediment  $(V_s)$  within that variation of pressure, considering a unit metre length of arc, is:

$$V_{s} = \left(\frac{\Delta P}{\rho_{s}g} \cdot \frac{1}{\sin(\alpha)}\right) t_{s} \tag{1}$$

where  $\alpha$  is the dip of the slab at each pressure step. The mass of the sediment ( $M_s$ ) is the product of sediment volume and density that are assumed constant. We assume 3000 kg m<sup>-3</sup> as representative density of sediments and AOC basalts at sub-arc depths. This assumption may promote a minor overestimation of CO<sub>2</sub> fluxes at shallower depths.

Assuming a speed of subduction  $u_{s}$ , the sediments remain within a 5 MPa range for a time  $(\tau_{\Delta P})$ :

$$\tau_{\Delta P} = \left(\frac{\Delta P}{\rho_s g} \cdot \frac{1}{\sin\left(\alpha\right)}\right) \cdot \frac{1}{u_s} \tag{2}$$

From our numerical results, we have derived the weight percent of CO<sub>2</sub> lost at each pressure step, and with our estimates of  $M_s$  and  $\tau_{\Delta P}$  we can compute the CO<sub>2</sub> flux emitted every 5 MPa. Finally, to constrain the CO<sub>2</sub> flux from 1 to 5.5 GPa we quantify the total CO<sub>2</sub> flux emitted at fore-arc (1 to 2.3 GPa) and sub-arc (2.3 to 5.5 GPa)

#### Table 1

The estimates of the CO<sub>2</sub> flux emitted from subducting slabs and the measured CO<sub>2</sub> fluxes from arc volcanoes.

Subduction Trench	Metamorphic decarnonation of AOC basalts			Metamorphic decarbonation of sediments			Metamorphic decarbonation - Total	CO <sub>2</sub> flux from arc-volcanoes (a,b,c)
	Subducted inorganic CO <sub>2</sub> (Mt/vr)	CO <sub>2</sub> flux 1-2.3 Gpa (Mt/vr)	CO <sub>2</sub> flux 2.3-5.5 Gpa (Mt/vr)	Subducted inorganic CO <sub>2</sub> (Mt/vr)	CO <sub>2</sub> flux 1-2.3 Gpa (Mt/vr)	CO <sub>2</sub> flux 2.3-5.5 Gpa (Mt/yr)	CO <sub>2</sub> flux 2.3-5.5 Gpa	(Mt/vr)
Camboursto noon lithelesioo	(, 91)	(1111/91)	(	(	(1112/31)	(1112/91)	(	(
Carbonale-poor innoiogies	1.06	0.00	0.80	0.52	0.00	0.52	1 /2	155
Kurilo (Kur)	1.00	0.00	0.89	0.05	0.00	0.55	0.95	0.71
Kuille (Kui) Hokkoido (Hok)	1.05	0.00	0.60	0.05	0.00	0.05	0.85	0.71
Honshy Japan (Hon)	116	0.00	0.08	0.17	0.00	0.17	0.85	0.05
Kuuchu (Kuu)	1.10	0.00	1.05	0.22	0.00	0.22	1.00	0.69
Ryusilu (Ryu)	1.05	0.00	1.05	0.04	0.00	0.04	1.09	0.08
Sumatra (Sum)	1.01	0.00	1.01	0.04	0.00	0.04	2.09	0.28
Sunda Strait (SuS)	0.40	0.00	0.40	0.70	0.00	0.70	0.76	0.07
Java (Java)	1.00	0.00	1.00	0.28	0.00	0.28	1.71	1.04
Java (Jav) Ball/Lombolz (BL)	1.05	0.00	1.05	0.02	0.00	0.02	1.71	0.72
Tonga (To)	2.20	0.00	0.57	0.08	0.00	0.08	0.79	0.00
Aloutions (Alo)	2.20	0.01	1.04	0.21	0.00	0.21	1.49	0.09
Alaska (Ala)	0.70	0.00	0.70	0.44	0.00	0.44	1.48	1.02
AldSKa (Ald)	0.79	0.00	0.79	0.09	0.00	0.09	0.00	0.28
Maxico (Max)	1.50	1.50	0.00	0.05	0.05	0.00	0.00	2.02
Aptillos (Apt)	0.20	0.00	0.00	0.15	0.13	0.00	0.00	0.71
North Chile (NC)	1.00	0.00	1.00	0.20	0.00	0.20	216	0.71
Control Chile (CC)	1.35	0.00	1.33	0.17	0.00	0.17	2.10	0.45
South Sandwich (SS)	2.23	0.00	2.23	0.19	0.00	0.15	1.00	0.00
Carbonate_rich lithologies	0.75	0.00	0.75	0.25	0.00	0.25	1.00	0.12
	0.82	0.00	0.66	2.00	0.00	0.34	1.01	176
Marianas (Mar)	0.02	0.00	0.00	1.03	0.00	0.54	0.57	0.79
North Philippines (NPh)	172	0.00	172	15.60	0.00	216	3.88	0.82
South Philippines (SPh)	3.21	0.00	2.57	104	0.00	0.10	2 75	0.32
New Britain (NB)	1.83	0.00	1.05	1.04	0.00	1.73	2.75	3.28
Solomon Sol)	1.05	0.00	0.52	-4.55 0.77	0.00	0.62	114	2.28
Vanuatu (Van)	3.43	0.00	3.42	5.49	0.00	2.05	5.47	7.40
New Zealand (NZ)	0.45	0.01	0.65	10.64	0.00	2.05	2.47	1.45
Custemala_ELSal (CES)	0.05	0.00	0.05	8 55	0.00	0.73	162	1.07
Nicaragua (Nic)	1.00	0.00	1.00	8.01	0.01	1.50	2.68	3.42
Costa Pica (CP)	0.47	0.00	0.47	4.21	0.00	0.74	1 20	1.26
Colombia-Ecuador (CE)	0.47	0.00	0.47	11.20	0.00	2 00	3.07	1.20
Doru (Dor)	0.30	0.02	0.35	11.05	0.00	2.33	1.52	/ 0.0/
Calabria (Cal)	0.07	0.00	0.30	18 37	0.00	0.58	0.08	5.61
Total	39.49	2.00	31.90	10.57	0.00	0.00 21.47	53.46	50 57
iotai	55.45	2.11	31,33	105.00	0.20	21.77/	JJ.=0	50.57

Note: The estimates of the  $CO_2$  flux emitted from subducting slabs reported here (as a result of the metamorphic decarbonation model) were calculated using the composition and thickness of each sediment lithology (Plank and Langmuir, 1998; Plank, 2014; Clift, 2017), the entire length of each arc sector and the subduction speed (van Keken et al., 2011) (Supplementary Table 1).  $CO_2$  fluxes emitted from subducting AOC basalts through metamorphic decarbonation are calculated considering the upper basaltic portion (300 m of thickness) of the altered oceanic crust (Kelemen and Manning, 2015; Gorce et al., 2019; Merdith et al., 2019; Li et al., 2019). Metamorphic decarbonation along each subducting slab was investigated using a typical hydrated and carbonated AOC basalt (Staudigel et al., 1989) (Supplementary Table 1). The  $CO_2$  fluxes from arc volcances (persistent and diffuse degassing; Supplementary Table 2) were measured from previous studies conducted by <sup>*a*</sup>Burton et al. (2013), <sup>*b*</sup>Aiuppa et al. (2019) and <sup>*c*</sup>Werner et al. (2019).

depths (Table 1) through multiplying these per-metre outputs by the total length of the trench (Syracuse et al., 2010).

### 3. Results and discussion

# 3.1. The efficiency of decarbonation during subduction: the role of bulk composition and thermal regime

Our simulations of metamorphic decarbonation from different lithologies (sediments and AOC basalts) under different thermal regimes indicate that the proportion of  $CO_2$  lost from subducting sediments decreases with increasing initial carbonate content (Fig. 3). Sediments such as siliceous ooze and carbonated clay with <10 vol.% of carbonate minerals (<5 wt.% CO<sub>2</sub> in the bulk composition) experience complete decarbonation beneath forearcs in subduction zones (with rarer high temperature pathways such as at Mexico and Cascadia) and beneath volcanic arcs for more common lower temperature subduction pathways (Fig. 4). In calcareous nannofossil oozes, chalks, and limestones, which contain more than 25 vol.% of carbonate minerals (12–34 wt.% CO<sub>2</sub>), partial decarbonation occurs such that only 5 to 40% of their initial CO<sub>2</sub> is lost (Figs. 3 and 4). Therefore, 60 to 95% of the CO<sub>2</sub> sub-

ducted in these lithologies would be subducted to greater than 170 km depth, beyond where it could be incorporated into arc magma sources (Figs. 3 and 4a), a conclusion consistent with previous studies (Gorman et al., 2006; Kerrick and Connolly, 2001; Connolly, 2005; Thomsen and Schmidt, 2008; Thomson et al., 2016; Wong et al., 2019). These results highlight the importance of an arc-by-arc evaluation of slab CO<sub>2</sub> loss, as the fate of slab carbon depends on the arc-specific vol.% of carbonate minerals subducted and the subduction pressure-temperature pathway (Supplementary Fig. 1).

Our simulations indicate that metamorphic decarbonation from AOC basalts is 100% efficient in most of the subduction zones under intermediate and warm thermal regimes (Fig. 5; Table 1). Decarbonation is incomplete at sub-arc depths in the coldest sub-ducting slab (Tonga, Solomon, Honshu, Hokkaido, Kuriles, Calabria and Kamchatka; Fig. 5; Table 1) and in three subduction zones with intermediate thermal regimes (Izu, South Philippines and New Britain). Thus, our results are consistent with the findings of Li et al. (2019) who proposed that altered oceanic crust may transport subducted carbon to the sites of diamond formation. We note that cold thermal regimes are more efficient in releasing CO<sub>2</sub> form carbonate-poor sediments than from AOC basalts (Figs. 4 and 5; Table 1). Intermediate thermal regimes can promote a complete



**Fig. 3.** Extent of CO<sub>2</sub> loss during metamorphic decarbonation. Bold black lines represent the percentage of CO<sub>2</sub> lost from carbonate-bearing sediments. Carbonated clay with a bulk composition containing <5 wt.% CO<sub>2</sub> (<10 vol.% of carbonate minerals) is representative of carbonate-poor sediments. Carbonate minerals within siliceous ooze and carbonated clay can be completely dissolved during subduction. Medium and rich carbonate lithologies, such as calcareous nannofossil ooze, chalk and limestones with a bulk composition containing 12–34 wt.% CO<sub>2</sub>, are only partially dissolved during subduction. The blue arrow indicates an increase of the extent of CO<sub>2</sub> loss as a function of the thermal regime, which goes from a colder (Calabria) to a warmer (New Zealand) thermal regime.

decarbonation from AOC basalts and carbonate-poor sediments, and incomplete decarbonation from chalk and limestone. Decarbonation of sediments and AOC basalt occur at sub-arc depths under intermediate thermal regimes. Complete decarbonation of AOC basalts is observed beneath fore-arcs in warm subduction zones (Mexico and Cascadia) (Fig. 5; Table 1); this is similar to what we observed for carbonate-poor sediments in warm thermal regimes. We note that the approach of our model does not consider the transfer of CO<sub>2</sub> derived from AOC basalts to sediments. If we consider the scenario in which CO<sub>2</sub> released from basalts is transferred to carbonate-beating sediments, decarbonation may be slightly less efficient within carbonate-rich sediments (Fig. 3). Therefore, our model may overestimate the CO<sub>2</sub> flux released from the carbonate-rich sediments.

Under cold and intermediate thermal regimes, which account for 94% of the arcs considered here, our models indicate that metamorphic decarbonation of carbonate-bearing sediments and AOC basalt occurs mostly between ~80 and 170 km depth (~2.3 to 5.5 GPa) (Figs. 4 and 5; Table 1). This depth range corresponds to the zone beneath volcanic arcs, therefore, this CO<sub>2</sub> may be incorporated into the mantle wedge sources of arc magmas. Our models indicate that CO<sub>2</sub> is released in sharp pulses at fore- and sub-arc depths (Figs. 4b and 5b). Multiple decarbonation pulses may affect the timescale and efficiency of the CO<sub>2</sub> transfer from the slab to the surface.

Figs. 4b and 5b show that decarbonation occurs at lower pressures under warm thermal regimes and to gradually higher pressures under intermediate and cold thermal regimes. The calcareous nannofossil ooze models (with 18 wt.% CO<sub>2</sub>; Fig. 3) demonstrate that warmer thermal regimes (e.g., New Britain) can generate greater CO<sub>2</sub> loss for a single bulk composition (as percentage of the bulk composition) than colder geothermal paths (e.g., Solomon and New Zealand) (Figs. 3 and 4a, b). This is also observed for sub-

ducting chalks and limestones (with  $\sim$ 25 wt.% CO<sub>2</sub>; Fig. 3), where warmer thermal regimes such as Colombia-Ecuador can produce greater CO<sub>2</sub> loss than North Philippines, Nicaragua and Costa Rica (with the exception of South Philippines (Figs. 3 and 4a, b).

It is only under the warmest thermal regimes on Earth that metamorphic decarbonation of carbonate-poor sediments occurs at fore-arc depths (40–60 km depths), probably before the depth where  $CO_2$  can be incorporated into arc magmas (Figs. 4b and 5b). Our results are consistent with the aforementioned geochemical findings for subducted carbonate-bearing rocks from the Aegean reported by Stewart and Ague (2020). Our modelling of 33 subduction zones concurs that metamorphic decarbonation is promoted at temperatures above 500 °C but such conditions are usually reached between 2.3 and 2.7 GPa under cold and intermediate thermal regimes, i.e. beneath the sites of arc magma genesis.

The approach of Syracuse et al. (2010) did not incorporate shear heating along the plate interface. We explored the consequence of adding reasonable levels of shear heating at the plate interface following van Keken et al. (2018) on metamorphic decarbonation We determined that shear heating only slightly affects the depths where decarbonation occurs. The quantitative effect on decarbonation is negligible (Supplementary Fig. 2). The lack of impact of shear heating is in part due to the inability of shear heating to significantly heat the slab interior even if the slab top is hotter (see, e.g., Figure 3 in van Keken et al., 2019).

# 3.2. Contribution of decarbonation on the deep carbon cycle: $\rm CO_2$ transport from slab to surface

For each arc we can compare our modelled CO<sub>2</sub> flux from carbonate-bearing sediments and AOC basalts into the base of the mantle wedge with the volcanic CO<sub>2</sub> degassing output. Table 1 lists the CO<sub>2</sub> flux into the wedge over the 2.3–5.5 GPa pressure range for the entire length of each arc sector calculated from our model. If we assume that these margins are in a near steady-state then the CO<sub>2</sub> output flux produced by the volcanic arc (through persistent and diffuse degassing) should match this input flux, unless other processes of CO<sub>2</sub> loss or gain are in play. As noted in Aiuppa et al. (2019), current limits on gas monitoring probably do not capture entire arc CO<sub>2</sub> outputs. Furthermore, the carbonate abundance we measure in sediments today may be different from those currently breaking down under mantle wedges. Notwithstanding these limits we may gain insight into the global geological CO<sub>2</sub> cycle by comparing the input and output fluxes. We are helped by the fact that a large proportion of arc-CO<sub>2</sub> loss is from degassing volcanic systems whose fluxes have been measured or estimated. Therefore, we can test an initial steady-state hypothesis that the presentday  $CO_2$  outputs (from the Aiuppa et al. (2019) and Werner et al. (2019) datasets) are balanced by the inputs from slab lithologies currently subducted under each margin. We do this by comparing the CO<sub>2</sub> lost from that sector by magmatic degassing to that supplied to each arc sector by metamorphic decarbonation of carbonate-bearing sediments and AOC basalts (Fig. 6a).

Fig. 6a shows that the correlation between  $CO_2$  slab input and magmatic  $CO_2$  output from several arcs cluster around and below the 1:1 line, while this correlation clusters above the 1:1 line only for a few subduction zones. This result supports the steady-state hypothesis and demonstrates that slab decarbonation processes followed by transport of  $CO_2$  through the mantle wedge melting zone to magmatic systems have the potential to play an important role in controlling the magnitude of arc  $CO_2$  degassing in most of the subduction zone.

In some subduction zones the input of  $CO_2$  into the mantle wedge from carbonate-bearing sediments and AOC basalts exceeds the arc output (i.e. those arcs that plot below the 1:1 line in Fig. 6a), while in others the volcanic  $CO_2$  output exceeds that



**Fig. 4.** Metamorphic decarbonation of carbonate-bearing sediments in 33 subduction zones. We used representative sediment compositions (Supplementary Table 1) for the subduction zones investigated. Decarbonation trend (a) and  $CO_2$  fluxes (b) as function of depth in each subduction zone. The hottest subduction zones (Mexico and Cascadia) are fully decarbonated in the fore-arc. The rest of the subducting slabs with colder thermal regimes than Mexico and Cascadia lose significant  $CO_2$  between ~2.3 and 5.5 GPa (between ~80 and 170 km depth). See Table 1 for abbreviations. Estimates of the  $CO_2$  flux (Table 1), reported here (b), were calculated using the thickness of each sediment lithology (Plank and Langmuir, 1998; Plank, 2014; Clift, 2017) and the speed of subduction (van Keken et al., 2011). The subduction zone colours are coded by slab temperature considering the temperature at 100 km depth within the sediments (van Keken et al., 2011). (For interpretation of the colours in the figure(s), the reader is referred to the web version of this article.)



**Fig. 5.** Metamorphic decarbonation of carbonated AOC basalts in 33 subduction zones. The simulations of the 33 subducting slabs were performed using a representative AOC basalt (Supplementary Table 1). Decarbonation trend (a) and  $CO_2$  fluxes (b) as function of depth in each subduction zone. See Table 1 for abbreviations. Estimates of the  $CO_2$  flux (Table 1), reported here (b), were calculated using a thickness of 300 m and the speed of subduction (van Keken et al., 2011). (For interpretation of the colours in the figure(s), the reader is referred to the web version of this article.)



Fig. 6. Comparison of CO<sub>2</sub> released from volcanic arcs (Aiuppa et al., 2019; Werner et al., 2019) with the  $CO_2$  input from sub-arc slab. (a) The  $CO_2$  input from sub-arc slab obtained through metamorphic decarbonation. (b) The CO2 input from slab between 0 and 230 km depth obtained through carbonate dissolution (Farsang et al., 2021). (c) The CO<sub>2</sub> input from slab between 0 and 230 km depth obtained through the combination of metamorphic decarbonation and carbonate dissolution (Farsang et al., 2021). Only the subducting slabs in which metamorphic decarbonation is not 100% efficient in liberating CO2 at fore- and sub-arc depths are considered in the panel (c), as carbonate dissolution may be more efficient at sub-arc depths (Galvez et al., 2016; Farsang et al., 2021); so metamorphic decarbonation and dissolution might occur together at sub-arc depths. Circles: Subduction of low-carbonate sediments. Squares: Subduction of high-carbonate sediments. Inclined lines illustrate the magnitude of the volcanic output compared to the sub-wedge input of CO2. Colour indicates geographic region: Green - subduction of East Pacific lithosphere; Yellow - subduction of West Pacific lithosphere; Dark blue - subduction of Northeast Indo-Australian lithosphere; Grey - subduction of Philippine Sea Plate lithosphere; Pale blue - subduction of North Indian-Australia lithosphere (Sunda Arc); Fuchsia subduction of Atlantic or Mediterranean lithosphere. See Table 1 for abbreviations.

which we have modelled being added to the wedge from sediment and AOC basalts (i.e. those arcs that plot above the 1:1 line in Fig. 6a). We observe a general tendency for carbon-rich (squares) or carbon-poor (circles) sediments and AOC basalts to lead to one or the other of these situations. We observe that volcanic CO<sub>2</sub> outputs exceed in some subduction zone with carbonate-rich sediments (e.g., Calabria, Solomon, Izu). This result indicates that metamorphic decarbonation of chalk and limestone, in addition to the CO<sub>2</sub> lost from AOC basalts, may be not efficient in liberating enough CO<sub>2</sub> from the slab under cold thermal regimes to balance the arc volcanic CO<sub>2</sub> emissions. In general, we observe that there are no excess volcanic CO<sub>2</sub> emissions in the majority of subduction zones with carbonate-poor sediments (Fig. 6a); this indicates that metamorphic decarbonation of AOC basalts and clay could liberate sufficient CO<sub>2</sub> from the slab to balance the arc volcanic CO<sub>2</sub> outputs. The exceptions are Cascadia and Mexico that are displaced above the 1:1 line, as volcanic  $CO_2$  outputs exceed the  $CO_2$  slab input at sub-arc depths.

The results discussed in our study on carbon mobilization and transport from the slab to the surface should be contextualized considering that in our decarbonation model we assume oxidizing conditions (Debret and Sverjensky, 2017). However, the redox state of subducting slabs may differ widely (Galvez et al., 2013; Debret and Sverjensky, 2017). For instance, reduced fluids may react with carbonate minerals that can favour their saturation in carbon leading to precipitation of graphite (Galvez et al., 2013). This may affect the mobilization and transport of subducting carbon, hindering carbon recycling to the surface. Rock-carbonation by fluid–rock interactions may also have an important impact on the direct transfer of carbon to the mantle wedge, providing a mechanism to stabilize and retain a part of the total carbon within a subducting slab (Piccoli et al., 2016).

The rate of water flow within the slab also affects carbon mobilization. It has been demonstrated thermodynamically that pervasive flow promotes greater mobilization of carbon than channelized flow from the slab to the mantle wedge (Gorman et al., 2006). Plümper et al. (2017) also show that both pervasive and channelized water flows occur within the slab, but that pervasive flow is considered subordinate to channelization. In order to consider a scenario in which the water flow is faster we increased the transport rate by a factor of 100 (from 0.25 m yr<sup>-1</sup> to 25 m yr<sup>-1</sup>; Wilson et al., 2014). By increasing the water flow velocity within the AOC basalts and sediments, we provide less H<sub>2</sub>O to the system that will be accounted for by the metamorphic solid-solution reactions. This approach allows us to consider only a percentage of the water supplied by the slab serpentinization (van Keken et al., 2011) for the equilibrium. Despite the transport rate increases by a factor of 100, thus reducing the H<sub>2</sub>O available for the metamorphic solid-solution reactions, decarbonation is still efficient considering fast transport rates (Supplementary Table 3). This indicates that decarbonation remains an important process in releasing CO<sub>2</sub> from the slab even if the amount of H<sub>2</sub>O available for metamorphic reactions is reduced by faster transport rates.

# 3.2.1. Comparison between metamorphic decarbonation and carbonate dissolution

Our thermodynamic model considers only  $CO_2$  mobilisation into the mantle wedge by metamorphic decarbonation. Carbonate dissolution (Frezzotti et al., 2011; Ague and Nicolescu, 2014; Kelemen and Manning, 2015; Galvez et al., 2016; Farsang et al., 2021) and melting (Poli, 2015; Mann and Schmidt, 2015) may also mobilise  $CO_2$ . It has been shown that carbonate melting will not occur in cool and medium temperature subduction pathways (Mann and Schmidt, 2015) and, therefore, we do not consider carbonate melting in our modelling. Carbonate dissolution is an important mechanism for carbon removal up to sub-arc depths in several subduction zones (Kelemen and Manning, 2015; Galvez et al., 2016; Farsang et al., 2021). We compare our results obtained from metamorphic decarbonation with the results obtained from Farsang et al. (2021) on carbonate dissolution below. This comparison helps us to better understand the potential contribution of both the processes in liberating  $CO_2$  from the slab to the mantle wedge in different subduction zones on Earth (Fig. 6a, b). This also provides insights into other potential processes that might release  $CO_2$  between the mantle wedge and the surface. These processes include diapirism of sediments intruding into the mantle wedge, in which  $CO_2$  from the slab may be added by assimilation of these sediments (e.g., Chen et al., 2021) and assimilation of carbon from the lithosphere (Deegan et al., 2010; Troll et al., 2012; Carter and Dasgupta, 2015; Aiuppa et al., 2017).

Fig. 6b compares the CO<sub>2</sub> that Farsang et al. (2021) determined may be supplied to each arc sector by carbonate dissolution of carbon-bearing sediments (inorganic and organic carbon) and AOC basalts with the CO<sub>2</sub> lost from that sector by magmatic degassing (Aiuppa et al., 2019; Werner et al., 2019). Farsang et al. (2021) considered carbonate dissolution at fore- and sub-arc depths (to 230 km depth), although the main contribution is obtained at subarc depths (Fig. 6b). Fig. 6b shows that in subduction zones with carbonate-poor sediments, with the exception of Mexico, volcanic CO<sub>2</sub> outputs do not generally exceed the inputs. However, in several subduction zones with carbonate-rich sediments (chalk and limestones) carbonate dissolution does not produce enough CO<sub>2</sub> from the slab to balance the arc volcanic CO<sub>2</sub> emissions (Fig. 6b). These results are broadly similar to our results obtained considering metamorphic decarbonation (Fig. 6a). Most of the water is released from the slab at sub-arc depths (van Keken et al., 2011), therefore decarbonation and dissolution might occur at the same depths in liberating CO<sub>2</sub>. For this reason, we combine fluxes produced by metamorphic decarbonation and mineral dissolution and we compare to fluxes at arcs where either process on their own is insufficient to explain the arc fluxes (e.g., Solomon, New Britain, Vanuatu, Nicaragua, Colombia-Ecuador, Izu, Marianas) (Fig. 6c). Metamorphic decarbonation can be efficient at fore- and sub-arc depths depending on the P-T path (Figs. 4 and 5), while carbonate dissolution may be more efficient at deeper depths such as subarc depths (Galvez et al., 2016; Farsang et al., 2021). Therefore, if metamorphic decarbonation is not efficient at fore-arc depths, it is possible that decarbonation and dissolution might occur at sub-arc depths. Particularly, if metamorphic decarbonation is not 100% efficient in liberating CO<sub>2</sub> from the subducting slab at sub-arc depth, dissolution may release the rest of the CO<sub>2</sub> within sediments and AOC basalts (Fig. 6c). We highlight that whether these two processes can operate at the same time and whether the operation of one affects the efficiency of the other is under debate. It is likely that the interaction between decarbonation and dissolution is very complicated because decarbonation will change the activity of CO<sub>2</sub> and H<sub>2</sub>O in the fluid, which will affect dissolution and vice versa. This implies that the increased CO<sub>2</sub> activity and decreased H<sub>2</sub>O activity will decrease the efficiency of both processes, therefore, the combined CO2 fluxes released from slabs reported in Fig. 6c are maximum fluxes. This suggests that the combined effects of metamorphic decarbonation and carbonate dissolution together may produce sufficient decarbonation to explain arc volcanic CO<sub>2</sub> outputs in subduction zones with carbonate-rich sediments (Fig. 6c).

Previous studies suggested that diapirism of buoyant sediment into the mantle wedge may provide a reservoir from which carbonate can be assimilated during ascent of arc magma, thus contributing to the carbon flux in volcanic arcs (Kelemen and Manning, 2015; Chen et al., 2021). Furthermore, assimilation of carbon from the lithosphere (magma-carbonate interaction) may be invoked to explain the difference between  $CO_2$  input and output (Carter and Dasgupta, 2015; Aiuppa et al., 2017). A more detailed discussion is provided below to understand the controls on  $CO_2$  fluxes and the contribution of metamorphic decarbonation. We first consider subduction zones on a geographic basis.

### 3.2.2. East Pacific

Most of the East Pacific subduction zones with carbonate-rich sediments (Columbia-Ecuador, Nicaragua, Guatemala-El Salvador, Costa Rica and Peru) plot close to the 1:1 line, implying that the CO<sub>2</sub> flux produced through decarbonation of sediment and AOC basalts can balance the volcanic outputs (Fig. 6a). This indicates that decarbonation may be an important process for the deep carbon cycle in these subduction zones. This is in agreement with previous studies, based on gas geochemistry, which suggest that decarbonation of subducting carbonate-rich sediments and AOC basalts may provide an important contribution in balancing the output from the arcs (Sano and Marty, 1995; Hilton et al., 2002; Shaw et al., 2003). We note that our model does not predict forearc decarbonation along the Costa Rica slab, which is in contrast with recent gas geochemistry studies (Shaw et al., 2003; Barry et al., 2019). Similar results can be observed in the scenario in which carbonate dissolution dominates the CO<sub>2</sub> liberation from the slab (Fig. 6b), with the exception of Nicaragua that plot more markedly above the 1:1 line.

CO<sub>2</sub> fluxes emitted from the slabs of North and Central Chile greatly exceed the volcanic emissions (Fig. 6a), which suggests that decarbonation can account for the entire volcanic output while transport of carbon to the surface may not be efficient and direct. This result corroborates evidence that the mantle wedge contains fluids transferred from the subducted lithosphere (Hickey-Vargas et al., 2002). We observe the same result if we consider the scenario in which the CO<sub>2</sub> fluxes of the Chilean segments are produced by carbonate dissolution (Fig. 6b). Therefore, CO<sub>2</sub> may be sequestered as carbonated rocks (Sieber et al., 2018) somewhere into the mantle wedge or arc lithosphere over long time scales (Hickey-Vargas et al., 2002).

Our models indicate that the Mexican and Cascadian slabs are sufficiently warm to purge most of their CO<sub>2</sub> before reaching 2.3 GPa. We have plotted these with a nominal slab input for comparative purposes, which makes clear that their volcanic CO<sub>2</sub> fluxes are far in excess of our model predictions (Fig. 6a). Mexico's 3.92 Mt/yr volcanic flux is also large compared to the majority of other arcs and is all the more notable for being mostly generated by a single volcano, Popocatépetl, which emits more CO<sub>2</sub> than any other terrestrial volcano (Aiuppa et al., 2019; Werner et al., 2019). Therefore, our model indicates that metamorphic decarbonation of the carbonate-bearing sediments and AOC basalts subducted beneath Mexico is not a viable source for the high CO<sub>2</sub> flux from Popocatépetl. The scarce efficiency of carbonate dissolution (Fig. 6b) along the Mexican slab (Farsang et al., 2021) indicate that dissolution is also not the process that promotes the high CO<sub>2</sub> flux from Popocatépetl. Previous studies suggested that the prodigious CO2 flux from Popocatépetl should result from magma assimilation of limestone in the upper crust (Schaaf et al., 2005; Carter and Dasgupta, 2015; Aiuppa et al., 2017). Likewise, our results also indicate that metamorphic decarbonation cannot be the dominant process supplying CO2 to the warm and complementary Cascadian arc (Figs. 4 and 5). Dissolution of inorganic and organic carbonate (Farsang et al., 2021) may be the dominant process along this warm slab (Fig. 6b). However, if all CO<sub>2</sub> in the sediments and AOC basalts is lost via metamorphic decarbonation at the fore-arc depths (Figs. 4 and 5), which means that by the time the slab reaches sub-arc depths it is essentially CO<sub>2</sub> free and there should be no inorganic carbonate available for dissolution at sub-arc depths. Therefore, organic carbonate could be the source for the Cascadian volcanic arc CO<sub>2</sub> emissions. These findings are supported by a field-based studies in an exhumed slab

from Greece which found that between 40 and 65 vol.% of the CO<sub>2</sub> in subducted sediments may be released into a fore-arc at  $\sim$ 500 °C (warm slab) via carbonate mineral dissolution (Ague and Nicolescu, 2014) and metamorphic decarbonation reactions (Stewart and Ague, 2020). The scenario offered by Mexico and the Cascadian, on the one hand, and the other eastern Pacific subduction zones, on the other hand, provides a framework to understand patterns of CO<sub>2</sub> input and output in many other convergent margins in the Atlantic, Mediterranean and Indian oceans.

### 3.2.3. West Pacific

Our modelled CO<sub>2</sub> input from subducted sediment and AOC basalts exceeds or equals the CO<sub>2</sub> emitted by western Pacific arcs in most of the cases except for Izu and Marianas (Fig. 6a). This is especially true of Kamchatka, Kurile, Hokkaido, Honshu, Tonga, Aleutians, Alaska and New Zealand. This result is supported by studies based on gas geochemistry, which indicate that the initial subducted carbon, dominantly sourced from AOC basalts (particularly Kurile and Kamchatka as reported by Fischer et al., 1998 and Hilton et al., 2002) and sediment, is mostly lost from the slab during metamorphic decarbonation at sub-arc depth (Sano and Marty, 1995; Hilton et al., 2002; Lopez et al., 2017; Epstein et al., 2021). The low volcanic CO<sub>2</sub> output may be a sampling artefact, since some of the volcanic outputs data rely on arcs that are (geographically or bathymetrically) remote and difficult to sample or where the current volatile output may not capture historic highs. Sequestration of CO<sub>2</sub> may also occur into the mantle wedge or arc lithosphere, which can prevent the transport of COH fluids through volcanic arcs. Nevertheless, the overall impression is that most of these arcs emit a relatively low proportion of the CO<sub>2</sub> that our models suggest is transferred from subducted sediments and AOC basalts to their mantle wedges. The exceptions are Izu and Marianas, as their volcanic CO<sub>2</sub> outputs exceed the slab-derived CO<sub>2</sub>, therefore, subducted sediments and AOC basalts may be not the main source of the CO2 emitted from Izu and Marianas volcanic arcs (e.g., Macpherson et al., 2010).

### 3.2.4. Northeast Indo-Australia and Philippine Sea Plate

The volcanic CO<sub>2</sub> output of the northeast Indo-Australia Plate, including the Solomon microplate, slightly exceeds the input of sedimentary and AOC basaltic CO<sub>2</sub> indicated by our decarbonation models while Vanuatu and New Britain plot close to the 1:1 line (Fig. 6a). The arc lithosphere in these subduction zones (Vanuatu, New Britain, and Solomon) is geologically young and does not host large deposits of sedimentary carbonate. Therefore, upper plate crustal contamination cannot be invoked as a substantial source of the excess CO<sub>2</sub> emitted from these arcs. Likewise, carbonate dissolution also fails to generate sufficient CO<sub>2</sub> from the slabs to account for the volcanic outputs and so can be considered an important source of volcanic carbon (Fig. 6b), but not the unique one. Therefore, we consider the slab derived CO<sub>2</sub> fluxes obtained from the combination of metamorphic decarbonation and carbonate dissolution (Fig. 6c), which implies that the two processes together can occur efficiently at sub-arc depths. The combination of decarbonation and dissolution can equal the CO<sub>2</sub> emitted by northeast Indo-Australia volcanic arcs (Fig. 6c). This suggests that the volcanic CO<sub>2</sub> output of the northeast Indo-Australia volcanic arcs may be fed by decarbonation and carbonate dissolution along subducting slabs at sub-arc depths.

CO<sub>2</sub> fluxes emitted from the slabs of the Philippines Sea plate exceed volcanic emissions (Fig. 6a). Regarding the Kyushu, Ryukyu and South Philippine arcs, decarbonation of carbonated AOC basalts provides the main contribution to the CO<sub>2</sub> released from their slabs into the mantle wedge. At these margins older oceanic lithosphere is subducted than at the Vanuatu, New Britain, and Solomon margins. Since secondary carbonate appears to build up in oceanic crust in the few million years after accretion (Li et al., 2019), the older altered oceanic crust in these subduction zones could contribute more  $CO_2$  than slabs feeding the northeast Indo-Australian arcs. The imbalance between the input and output of  $CO_2$  in the arcs of the Philippines Sea plate suggests that  $CO_2$  may be accumulated and stored into the mantle wedge or arc lithosphere before to reach the surface. This observation is also supported from the scenario in which carbonate dissolution was the main process in liberating  $CO_2$  from the slab (Fig. 6b).

#### 3.2.5. Sunda Arc

Metamorphic decarbonation of sediments and AOC basalts is an important process that may explain the Javan volcanic emissions (Fig. 6a). Carbonate dissolution may also provide an important contribution to balance the Javan volcanic outputs (Fig. 6b). Assimilation of crustal carbonate has been proposed as a further source of the emissions (Deegan et al., 2010; Troll et al., 2012; Aiuppa et al., 2017). In the Sumatra and Bali-Lombok sectors our estimate of how much CO<sub>2</sub> is added to the mantle wedge is slightly larger than their volcanic outputs (Table 1; Fig. 6a). Therefore, decarbonation of subducting sediments and AOC basalts may be the principal source of CO<sub>2</sub> emitted from these two sectors, which is also consistent with the findings of carbon isotope and  $CO_2/^3$ He studies (Poorter et al., 1991; Varekamp et al., 1992; Halldorsson et al., 2013). Results reported from Farsang at el. (2021) indicate that carbonate dissolution may also have an important role in recycling carbon from the Sumatra subducting slab (Fig. 6b). This suggests that decarbonation and/or dissolution may be important processes in liberating CO<sub>2</sub> from the Sumatra slab. The volcanic CO<sub>2</sub> output from the Sunda Straight can be comfortably accounted for by sediment and AOC basalt input (Fig. 6a). Indeed, at face value the low volcanic output in Sumatra, Bali/Lombok, and Sunda Straight may indicate substantial carbon sequestration in the mantle wedge.

### 3.2.6. Atlantic

In the Atlantic, less CO<sub>2</sub> is released from the South Sandwich Arc than is added to its mantle wedge, similar to the situation in the majority of other subduction zones with carbonate-poor sediments of the East and West Pacific, northeast Indo-Australia, Philippines Sea Plate, and Sunda arc. In contrast, the volcanic CO<sub>2</sub> output from the Lesser Antilles Arc is almost two times higher than the CO<sub>2</sub> input we determined from decarbonation of subducted sediments and AOC basalts (Fig. 6a). Despite decarbonation being 100% efficient and dominantly occurring at sub-arc depth in the Lesser Antilles Arc (Table 1; Figs. 4 and 5), as also observed by Li et al. (2020), it cannot be the unique process in liberating CO<sub>2</sub> from the slab. Only inorganic carbon is considered in our decarbonation model, but we note that organic carbon is abundant in the sediments of Lesser Antilles (Clift, 2017) and carbonate dissolution mobilizes organic (reduced) carbon from this slab (Farsang et al., 2021). Therefore, the contribution of carbonate dissolution might balance the volcanic CO<sub>2</sub> output from the Lesser Antilles Arc (Fig. 6b). Assimilation of crustal sediments is also a widely documented process in the geochemical evolution of Antilles magmatism (Davidson, 1987; Bezard et al., 2015) and recent tectonic models have strengthened the case for this sediment residing in the upper plate (Allen et al., 2019). Prior geochemical modelling of Lesser Antilles magmatism has focused on assimilation of clastic sediment but the excess CO2 arc emissions relative to our model suggests there may also be a potential remobilisation of upper plate carbonate-bearing sediments (Davidson, 1987).

### 3.2.7. Calabria

The Calabrian margin subducts Mesozoic oceanic crust of the Ionian Sea (Piana Agostinetti et al., 2009; Polonia et al., 2011), while its accretionary wedge comprises Plio-Quaternary units and

Messinian evaporites overlying Tertiary and Mesozoic sediments (Cernobori et al., 1996; Polonia et al., 2011). The Mesozoic carbonates comprise approximately 2000 m of limestone (Polonia et al., 2011) that lies on the underlying oceanic crust (Piana Agostinetti et al., 2009). Detachments in the accretionary wedge occur at the base of the Messinian evaporites and at the top of the Mesozoic carbonates, therefore, the Mesozoic limestones have the highest probability of subduction with the oceanic crust (Piana Agostinetti et al., 2009; Polonia et al., 2011). To investigate how subduction of carbonate-bearing sediment and AOC basalts contributes to the CO<sub>2</sub> fluxes emitted from Aeolian volcanoes and Etna, we assumed that the slab beneath Calabria is able to transport 1000 m of limestones to sub-arc depths and 300 m of carbonated AOC basalts (Supplementary Table 1). Our model indicates that in total, considering decarbonation of limestones and AOC basalts, 0.98 Mt/yr CO<sub>2</sub> can be released in the mantle at sub-arc depths. The estimate obtained with our decarbonation model is insufficient to account for the volcanic flux from the Calabria volcanoes (Etna, Stromboli, and Vulcano combined; Table 1; Fig. 6a). Our results indicate that decarbonation of limestones and AOC basalts may be one of the important sources of CO<sub>2</sub> that contribute to the high volcanic output of Calabria, which is consistent with the  $CO_2/^3$ He ratios and estimated sources of carbon in gas and fluids from Vulcano (Sano and Marty, 1995). Our results are also consistent with Tonarini et al. (2001), who proposed that the altered Ionian oceanic crust and its sedimentary cover are the sources of the fluids contaminating the mantle source of Mt. Etna magmas. However, this would require some portion of the volcanic CO<sub>2</sub> be derived from other sources. Etna gases are characterized by unusually high CO<sub>2</sub>/S, which have been attributed to magma-limestone interactions in the upper crust (Aiuppa et al., 2017, 2019). We suggest that a proportion of the CO<sub>2</sub> emitted from Etna is fed by decarbonation of subducting limestones and AOC basalts but that this is augmented by CO<sub>2</sub> liberated by magma-crust interaction. Furthermore, the role of carbonate dissolution in the Calabrian slab has not been investigated, but it may be an important CO<sub>2</sub> source that could be studied in future works.

## 4. Conclusion

Our study investigates the potential contribution of metamorphic decarbonation on the deep carbon cycle considering 33 subduction zones. A key result of our study is that the pervasive hydration of subducted sediments promotes intense sub-arc metamorphic decarbonation of carbonate-bearing sediments and AOC basalts, under cold and intermediate thermal regimes, directly beneath the observed locations of arc volcanism. Metamorphic decarbonation of AOC basalts is not 100% efficient under cold thermal regimes. The warmest slabs on Earth (Cascadia and Mexico) promote metamorphic decarbonation of carbonate minerals at fore-arc depths. Our modelling shows also that all of the CO<sub>2</sub> in carbonpoor lithologies is likely to be mobilized and lost from the slab. In contrast, the CO<sub>2</sub> from carbonate-rich lithologies, such as calcareous nannofossil ooze, chalk, and limestone, is only partially mobilized via decarbonation. Dissolution of inorganic and organic carbon (Kelemen and Manning, 2015; Farsang et al., 2021) is the other process that has been proposed to play an important role in liberating CO<sub>2</sub> from the slab. Both metamorphic decarbonation or carbonate dissolution by itself can liberate sufficient  $CO_2$  from carbonate-rich lithologies (calcareous nannofossil ooze, chalk and limestones) to balance volcanic CO<sub>2</sub> outputs for some arcs. The efficient occurrence of metamorphic decarbonation and carbonate dissolution together at sub-arc depths is needed to equal the CO<sub>2</sub> emitted by, for instance, northeast Indo-Australia, Nicaragua and Izu-Bonin volcanic arcs. We propose that the combination of decarbonation and dissolution is the best explanation for high CO<sub>2</sub> emissions from carbonate-rich subduction zones in Calabria, Nicaragua, Colombia-Ecuador, Vanuatu, New Britain, Solomon, Izu and Marianas. This is also applicable to some subduction zones with carbonate-poor sediments such as Lesser Antilles, Cascadia and Mexico.

Assimilation of carbon from the lithosphere may also contribute to the strong  $CO_2$  emission of Popocatépetl in Mexico. We observe that slab-derived  $CO_2$  exceeds the volcanic  $CO_2$  emissions in subduction zones with carbonate-poor sediments. This suggests that the mantle wedge or arc lithosphere may accumulate and store  $CO_2$ . The mantle wedge and lithosphere of any active arc may be dynamic reservoirs in which magmatism may add and remove multiple generations of  $CO_2$  at multiple levels. The pathways and processes by which this occurs are still difficult to investigate. Further modelling and experimental investigations are required to elucidate the fluid transfer processes within the mantle wedge.

### **CRediT authorship contribution statement**

**Fabio Arzilli:** Conceptualization, Data curation, Formal analysis, Investigation, Methodology, Software, Supervision, Validation, Visualization, Writing – original draft. **Mike Burton:** Conceptualization, Funding acquisition, Investigation, Methodology, Project administration, Resources, Supervision, Visualization, Writing – original draft. **Giuseppe La Spina:** Data curation, Formal analysis, Methodology, Software, Validation, Writing – review & editing. **Colin G. Macpherson:** Conceptualization, Data curation, Investigation, Methodology, Resources, Supervision, Visualization, Writing – original draft. **Peter E. van Keken:** Methodology, Resources, Software, Writing – review & editing. **Jamie McCann:** Formal analysis, Validation, Writing – review & editing.

# **Declaration of competing interest**

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

### Data availability

Data will be made available on request.

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### Appendix A. Supplementary material

Supplementary material related to this article can be found online at https://doi.org/10.1016/j.epsl.2022.117945.

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