# Detrital carbonate minerals in Earth's element cycles

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#### Abstract

We investigate if the commonly neglected riverine detrital carbonate fluxes might balance several chemical mass balances of the global ocean. Particulate inorganic carbon (PIC) concentrations in riverine suspended sediments, i.e., carbon contained by these detrital carbonate minerals, was quantified at the basin and global scale. Our approach is based on globally representative datasets of riverine suspended sediment composition, catchment properties and a two-step regression procedure. The present day global riverine PIC flux is estimated at  $3.1 \pm 0.3$  Tmol C/y (13% of total inorganic carbon export and 4 % of total carbon export), with a flux-weighted mean concentration of  $0.26 \pm 0.03$  wt%. The flux prior to damming was  $4.1 \pm 0.5$  Tmol C/y. PIC fluxes are concentrated in limestone-rich, rather dry and mountainous catchments of large rivers in Arabia, South East Asia and Europe with 2.2 Tmol C/y (67.6 %) discharged between 15 °N and 45 °N. Greenlandic and Antarctic meltwater discharge and ice-rafting additionally contribute  $0.8 \pm 0.3$  Tmol C/y. This amount of detrital carbonate minerals annually discharged into the ocean implies a significant contribution of calcium (~ 4.75 Tmol Ca/y) and alkalinity fluxes (~ 10 Tmol(eq)/y) to marine mass balances and moderate inputs of strontium (~ 5 Gmol Sr/y), based on undisturbed riverine and cryospheric inputs and a dolomite/calcite ratio of 0.1. Magnesium fluxes (~ 0.25 Tmol Mg/y), mostly hosted by less-soluble dolomite, are rather negligible. These unaccounted fluxes help elucidating respective marine mass balances and potentially alter conclusions based on these budgets.

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# Detrital carbonate minerals in Earth's element cycles

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Abbreviations: IC: Inorganic carbon, PIC: Particulate inorganic carbon, DIC: Dissolved inorganic carbon, POC: Particulate Organic Carbon, DOC: Dissolved Organic Carbon, SOC: Soil organic carbon, TC: Total river carbon, SVM: Support Vector Machine, MGGP: Multi-Gene Genetic Programming, SR: Symbolic Regression, RMSE: Root mean squared error, MC: Monte Carlo, VI: Variable Influence, fwm: flux-weighted mean, med: median, mod: modelled, wo: without, S: related to change in sediment discharge, D: related to damming, SC: Source Carbonate, hdi: human development index, gdp: gross domestic product, nli: night light index, pop: population count, N: Number of simulations.

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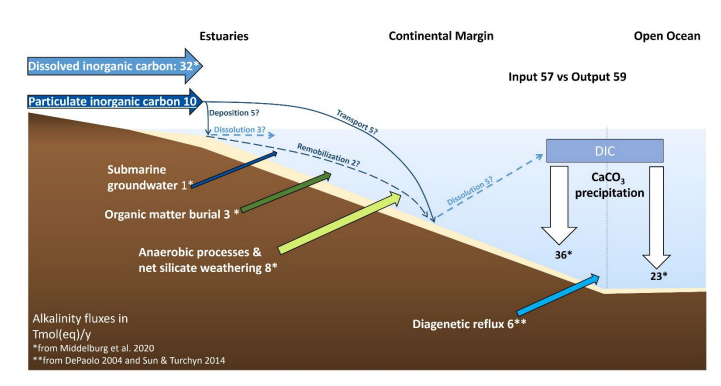
# **Key points:**

- The present day riverine detrital carbonate flux from land to sea is 3.1 ± 0.3 Tmol C/y
   (= 0.037 Pg C/y).
- Associated calcium, alkalinity and strontium contribute significantly to their global
   biogeochemical cycles.
- Damming reduced the riverine PIC flux by 25 % (from naturally 4.1 ± 0.5 Tmol C/y =
   0.049 Pg C/y).

**Abstract.** We investigate if the commonly neglected riverine detrital carbonate fluxes might balance several chemical mass balances of the global ocean. Particulate inorganic carbon (PIC) concentrations in riverine suspended sediments, i.e., carbon contained by these detrital carbonate minerals, was quantified at the basin and global scale. Our approach is based on globally representative datasets of riverine suspended sediment composition, catchment properties and a two-step regression procedure. The present day global riverine PIC flux is estimated at  $3.1 \pm 0.3$  Tmol C/y (13% of total inorganic carbon export and 4 % of total carbon export), with a flux-weighted mean concentration of  $0.26 \pm 0.03$  wt%. The flux prior to damming was  $4.1 \pm 0.5$  Tmol C/y. PIC fluxes are concentrated in limestonerich, rather dry and mountainous catchments of large rivers in Arabia, South East Asia and Europe with 2.2 Tmol C/y (67.6 %) discharged between 15 °N and 45 °N. Greenlandic and Antarctic meltwater discharge and ice-rafting additionally contribute  $0.8 \pm 0.3$  Tmol C/y. This amount of detrital carbonate minerals annually discharged into the ocean implies a significant contribution of calcium (~ 4.75 Tmol Ca/y) and alkalinity fluxes (~ 10 Tmol(eq)/y) to marine mass balances and moderate inputs of strontium ( $\sim 5$  Gmol Sr/y), based on undisturbed riverine and cryospheric inputs and a dolomite/calcite ratio of 0.1. Magnesium fluxes ( $\sim 0.25$  Tmol Mg/y), mostly hosted by less-soluble dolomite, are rather negligible. These unaccounted fluxes help elucidating respective marine mass balances and potentially alter conclusions based on these budgets.

Plain Language Summary. Earth surface conditions, including climate and sea level, are largely controlled by the cycling of carbon and biogeochemically coupled elements. However, most elemental budgets cannot be consentaneously balanced for the present state. Here, we investigate the possible role of riverine carbonate minerals in biogeochemical cycles. We derive individual river basin export fluxes, the global export flux to the ocean and its reduction by human influence, utilizing state-of-the-art regression techniques and published global-scale datasets. Results point to a significance of riverine detrital carbonates for the global mass balances of carbon, calcium, alkalinity and strontium, which might help solving this long-standing problem.

# **Graphical Abstract**



### 1 Introduction

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Erosion and weathering of Earth's surface not only shape landscapes, but also influence the global carbon cycle, thereby maintaining the habitability of our planet (Berner et al., 1983; Ebelmen, 1845; Ferrier and West, 2017; Penman et al., 2020; Urey, 1952; West et al., 2005). Oceanic mass balances of carbon (C) and biogeochemically coupled elements provide a powerful tool to investigate these processes and their role in the Earth system globally and over longer time-scales (classically > 100 ka) (Berner and Berner, 2012; Dickens, 2001; Krabbenhöft et al., 2010; Tipper et al., 2010). They also allow quantification of hardly measurable processes, such as global rates of marine carbonate burial or hydrothermal activity (Shalev et al., 2019; Tipper et al., 2006; van der Ploeg et al., 2019). However, some of the most prominent and most frequently considered budgets presented in that context remain unbalanced and/or highly debated, such as those of Ca, Mg, Sr and alkalinity (Berner and Berner, 2012, 1987; Gislason et al., 2006; Jones et al., 2012; Krabbenhöft et al., 2010; Lebrato et al., 2020; Milliman, 1993; Tipper et al., 2010, 2006). This is usually explained by disequilibrium, i.e., the present state strongly differs from average Pleistocene conditions, by proposing a variety of smaller-scale marine processes and/or by invoking yet unaccounted input fluxes (Krabbenhöft et al., 2010; Middelburg et al., 2020; Milliman, 1993; Shalev et al., 2019; Tipper et al., 2010). For most marine mass balances, riverine dissolved loads are traditionally considered the only major input term, reflecting the catchment-integrated result of chemical rock weathering as transported by the Earth-spanning fluvial networks (Berner and Berner, 2012, 1987). Some authors recognized submarine groundwater discharge as another important flux to the ocean with a probable magnitude of 0.7 to 6 % of the global river discharge (Mayfield et al., 2021; Milliman, 1993; Zhou et al., 2019). In addition to these dissolved inputs, it is generally accepted that organic and biogenic riverine particles exert major control on the biogeochemical cycling of carbon (C), nitrogen (N) and phosphorous (P) (Berner, 1999, 1982;

Boyer and Howarth, 2008; Froelich et al., 1982; Hilton and West, 2020), and of silicon (Si) (Conley, 2002; Sutton et al., 2018). Moreover, the importance of ions and complexes sorbed to the surfaces of riverine sediments was highlighted (Berner et al., 1983; Tipper et al., 2021). A similar importance was proposed for particulate inorganic forms (mineral detritus) of silicon (Si) (Mackenzie and Garrels, 1966, 1965), calcium (Ca) (Gislason et al., 2006), strontium (Sr) (Hong et al., 2020; Jones et al., 2012), iron (Fe) (Luo et al., 2020; Poulton and Raiswell, 2002) and other elements (e.g., Abbott et al., 2019; Jeandel et al., 2011), based on experimental and field-measured element release rates. Recently, based on a limited dataset, Middelburg et al. (2020) suggested that riverine particulate inorganic carbon (PIC) fluxes to the ocean may be about 1/3 of riverine dissolved inorganic carbon (DIC) fluxes and that the ocean alkalinity budget is close to balance when this is considered an additional alkalinity input. While basaltic minerals, ashes and glasses of volcanic origin are currently considered to be the major host minerals of particulate Ca and Sr fluxes (Gislason et al., 2006; Jones et al., 2012; Torres et al., 2020), significant riverine PIC fluxes would imply substantial additional Ca, Sr and Mg delivery in particulate forms.

Carbonate dissolution and recrystallization are well known to occur in estuaries (Aller, 1982; Gattuso et al., 1998; Santos et al., 2019) and the ocean (Krumins et al., 2013; Milliman, 1974; Sulpis et al., 2017), providing evidence for the (partial) release of Ca, Mg, Sr, inorganic C (IC) and alkalinity from detrital sources to the oceanic inventories. Dissolution of PIC in the ocean could, thus, represent a major missing term in oceanic mass balances, potentially altering the conclusions deduced from those budgets (e.g., Berner and Berner, 2012; Krabbenhöft et al., 2010; Paytan et al., 2021; Tipper et al., 2006, 2010). Notably, recrystallization within the sediment column, i.e., dissolution and direct re-precipitation, may result in an exchange of elements and isotopes between PIC and seawater (DePaolo, 2004; Fantle et al., 2010; Paytan et al., 2021).

High solubility and rapid dissolution kinetics of carbonate minerals cause the dominant mass of IC to be transported in dissolved form (Lasaga, 1984). Therefore, the significance of detrital carbonate minerals in river sediments is often neglected. However, detrital carbonates are commonly observed constituents of suspended sediments in rivers (Mackenzie and Garrels, 1966; Müller et al., 2021a) and even authigenic carbonate production in calcite-saturated rivers is common (Grosbois et al., 2001; Kempe and Emeis, 1985; Négrel and Grosbois, 1999). Such authigenic carbonate formation on land represents a (temporary) sink of weathering-derived cations, alkalinity and carbon, with implications for the location of gas exchange and global mass balancing (Rovan et al., 2021; Zhao et al., 2016). Sr-isotopic constraints suggest that 30 -50 % of the carbonate minerals within the Gulf of Lyon sediments are detrital, even more during glacial periods (Pasquier et al., 2019). Additionally, the isotopic composition of carbonates from turbidites in the Bengal fan, one of the largest sediment dispersal systems on earth (Mouyen et al., 2018), suggests a mixture of biogenic (> 85 wt%), detrital (up to 10 wt%) and diagenetic (1.2 – 4 wt%) origin (France-Lanord et al., 2018). This indicates that the PIC delivery may indeed be a relevant flux to the marine realm, but its size and dissolving fraction remain unclear because a global assessment is lacking.

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We aim to better constrain these important numbers based on several approaches. First, we establish a first-order calculation based on published average PIC and CaO concentrations and sediment fluxes. Next, we quantify PIC concentrations and fluxes globally at the basin-scale, using published datasets of riverine suspended sediment and catchment characteristics, and a two-step regression procedure, involving regressive classification and symbolic regression (for details see section 2.2; Regression and Upscaling & Supplementary Information SI 2). Controlling factors of the global PIC flux, human influence and the fate of the delivered detrital carbonates in the ocean are then discussed, including implications for oceanic mass balances and carbon cycling.

# 2 Methods and procedures

To calculate the global PIC flux, we need a gapless set of PIC concentrations and sediment fluxes of all rivers in the world, which is not realistically achievable from measurements. However, the latter can be generated using advanced models that provide suspended sediment fluxes of global rivers in space and time, based on water balance and catchment properties (WBMSed 2.0, Cohen et al., 2014). The WBMSed 2.0 provides anthropogenically disturbed suspended sediment flux data, as well as natural background values. These data, along with the locations of the river mouths, were taken from a compilation of the *GlobalDelta* project (Nienhuis et al., 2020). No such model is available for PIC concentrations yet. Hence, we here develop a statistical, spatially-explicit model that predicts PIC concentrations from catchment properties. Modelled PIC concentrations were combined with both, the natural and the anthropogenically disturbed suspended sediment fluxes (WBMSed 2.0) to arrive at the corresponding PIC fluxes.

Annual median PIC concentrations were calculated for all locations in the GloRiSe v1.1 database (Müller et al., 2021b) from direct measurements, mineralogical and petrographic observations or empirically from major element composition (Supplementary Information SI 1). The uncertainty of these concentrations was defined as the mean relative deviation of single measurements from the flux-weighted mean of available time-series (Müller et al., 2021b). A large set of hydro-environmental and physiographic variables was derived from the HydroBasins database (Linke et al., 2019) by spatially assigning each GloRiSe-location to the corresponding sub-basin (at Pfafstetter level 7). Annual averages for the upstream catchment of nine variables were selected based on correlation analysis and/or a causal link to PIC concentrations (Supplementary Information SI 1). These variables cover topography, vegetation, hydrology, climate and human impact (Table 1). As the carbonate in the catchment is the source of riverine PIC, a proper indication of this 'source carbonate' (SC) was extracted

from global maps of lithology (GLiM, Hartmann and Moosdorf, 2012), unconsolidated sediments (GUM, Börker et al., 2018) and soils (WISE, Batjes, 2012). For soils, the carbonate content was given directly, while for each rock and sediment class a global representative estimate of the carbonate content was taken from literature (Supplementary Information SI 1). Area-weighted upstream averages were calculated individually for the carbonate content of GliM, GUM and WISE in each basin. Next, these were summed and normalized to 100 % to represent the SC, i.e., carbonate available to be transported as PIC. All the predictor variables are summarized in Table 1. Catchments with SC < 10 % were assumed to be PIC-free, as dissolution usually dominates over detrital carbonate transport in (undersaturated) rivers (see 1 Introduction).

Table 1 Predictor variable selection to model PIC concentrations. Variables are taken from HydroBasins (Linke et al., 2019), except for the potential source carbonate, which was calculated from global soil, sediment and lithological maps (Batjes, 2012; Börker et al., 2018; Hartmann and Moosdorf, 2012). All variables represent the upstream-average of a specific HydroBasins sub-basin at Pfafstetter level 7. Abbreviations: hdi: human development index, gdp: gross domestic product, nli: night light index, pop: population count.

Topography & Vegetation	Underground & Humans	Climate & Hydrology
Elevation	Potential source carbonate (rock, sediment, soil)	Precipitation
Upstream catchment area	Soil organic carbon content	Temperature
Forestation	Human factor (log(hdi+gdp+nli+pop))	Extent of water bodies (rivers, lakes, reservoirs)
Bare areas (rock, desert, tundra, open shrub land)		

#### 2.1 Regression & Upscaling

To estimate PIC concentrations in the remaining ~65 % of the global suspended sediment discharge, we employed a two-step regression procedure consisting of (I) a qualitative indication of the presence of PIC in a catchment (yes/no) and (II) a quantitative regressive

estimation of the PIC concentration. This two-step procedure was necessary because PIC concentrations are not only log-normally distributed, but also frequently close or equal to zero, thus hampering the regression procedure, which is a well-known problem in ecology (Fletcher et al., 2005).

For the qualitative model, we applied a Support Vector Machine (SVM), a standard technique from the MATLAB 2019b Machine Learning toolbox. This model was trained and forced by only five variables, because SVMs have been found to achieve better results with less variables (Kitsikoudis et al., 2013). We chose to use SC, precipitation, elevation, forestation and human factor, covering the most diverse aspects of sources and preservation potential of PIC. PIC concentration was assumed not present if it was below 0.1 wt%, approximating the uncertainty of most measurements included in *GloRiSe*. SVM was chosen, because it performed slightly better than alternative methods, such as logistic regression and ensemble techniques.

For the quantitative model, symbolic regression (SR) by means of multi-gene genetic programming (MGGP) was used, providing a fully data-driven tool to find both the model structure and its parameters. SR was chosen, because it performed better than simple linear regression or alternative machine learning techniques (available e.g., in the MATLAB Regression Learner Toolbox) in terms of both, accuracy and precision. The implemented SR-algorithm pseudo-randomly creates linear combinations of (potentially non-linear) terms, which are tested and evolved to best fit the observed PIC concentrations as assessed by the root mean squared error (*GPTIPS* 2.0, Searson et al., 2010). Thus, SR is able to cover non-linear relationships between the variables and its performance seems comparable to artificial neural networks, while it still results in comparably simple equations that can be related to the governing processes (Gandomi et al., 2015; Jin et al., 2019; Kitsikoudis et al., 2013). Variable selection and SR intrinsically determine the importance of individual variables for, and their

direction of relationship to PIC concentrations, which we quantitatively assess using the linear correlation coefficient and coefficient of determination (R and  $R^2$ , respectively, at p < 0.01) between the median result of 830 accepted Monte Carlo simulations and each variable. This method reduces biases due to multi-collinearity and non-linearity and is commonly applied to the evaluation of canonical correlations analyses (Kuylen and Verhallen, 1981).

The global riverine PIC flux is the sum of the products of sediment fluxes and PIC concentrations in each basin draining directly to the coastal ocean. For a proper re-estimation and uncertainty analysis, the regression and prediction procedure was repeated 2,000 times with a (pseudo-)random perturbation of sediment fluxes and PIC concentrations within the range of their respective uncertainties, including the full model derivation via SVM and SR. The final result is the mean of 830 accepted simulations that produced less than 0.3 % outliers in respect to the 10 % and 90 % percentile (10 of 3364 coastal basins) and its uncertainty is the standard deviation of these models (Koehler et al., 2009). For comparison, we also provide literature-based first-order estimates of riverine PIC fluxes (Supplementary Information S3). Because much less detailed data is available for atmospheric and cryospheric PIC contributions, these fluxes were estimated using published PIC concentrations and sediment fluxes (Supplementary Information S3).

# 217 3 Results

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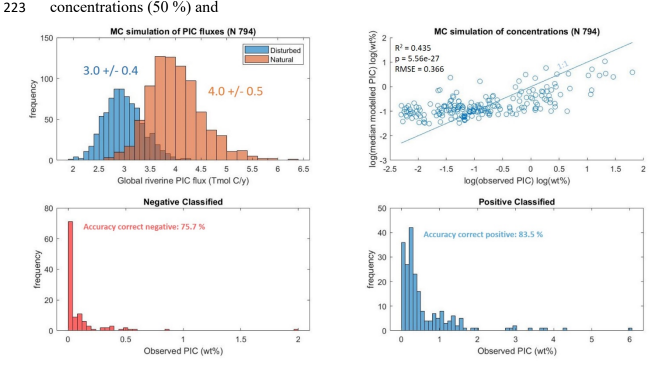
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We calculate that currently  $3.1 \pm 0.3$  Tmol PIC are annually discharged to the coastal ocean. The pre-human flux was  $4.1 \pm 0.5$  Tmol PIC/y (Fig. 1), accounting for damming and soil erosion (by the underlying WBMSed 2.0 model (Cohen et al., 2014, 2013)). The 25 % reduction is dominated by particle retention in reservoirs. The uncertainty of 10 % appears low, considering the much larger uncertainty of sediment fluxes (50 %), observed PIC concentrations (50 %) and



**Figure 1**: Results (upper) and performance (lower) of the Monte Carlo (MC)-refined regression procedure. Histograms (upper left panel) show the distribution of natural and anthropogenically disturbed global PIC fluxes in Tmol C/y (times 0.012011 yields Pg/y). The upper right panel assesses the performance of the quantitative prediction via SR (1:1 line = perfect prediction). The lower panels evidence the performance of an exemplary qualitative model (left: negative classifications (= No PIC present, correct predictions are < 0.1 wt%); right: positive classifications (= PIC present, correct predictions are > 0.1 wt%). N is the number of accepted MC simulations. RMSE is the root mean squared error.

Table 2 Comparison of the herein presented results and literature-based estimates of global average PIC concentration (cPIC, flux-weighted mean, median and mixture of median and mean, respectively), suspended sediment discharge (fTSS, global sum) and PIC flux (fPIC, global sum). References: 1: Meybeck (1982), 2: Viers et al. (2009), 3: Savenko (2007), 4: Bayon et al. (2015), 5: Beusen et al., (2005), 6: Milliman and Farnsworth (2011), 7: Syvitski and Kettner (2011), 8: Cohen et al. (2014), 9: Middelburg et al. (2020) based on Canfield, (1997) and Beusen et al. (2005), 10: Meybeck (1993), 11: Journet et al. (2014), 12: Jickells et

al (2005), 13: Overeem et al. (2017), 14: Raiswell et al. (2008), 15: Wadham et al. (2013). Abbreviations: med: median, fwm: flux-weighted mean, obs: observations, wo: without,. 'Literature' indicates values and ranges that were calculated from published values ('first-order' estimates, Supplementary Information SI 2, grey columns). 'This study' refers to values we derived in this contribution (2 Methods & Procedures, Supplementary Information SI 1). Bold numbers indicate the values suggested for further use. Conversion to Pg/y by a factor 0.012011.

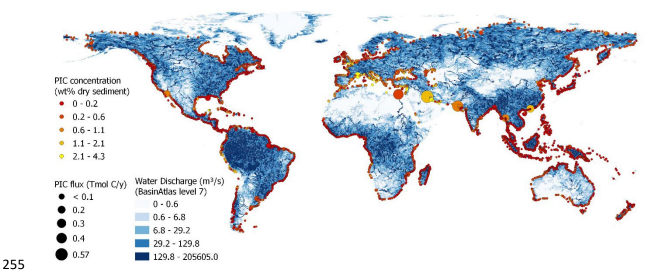
Variable	cPIC	cPIC	cPIC	fTSS	fPIC	fPIC	fPIC	fPIC	fPIC
(unit)	(wt%)	(wt%)	(wt%)	river	river,	river,	river,	atmosphere	cryosphere
				(Gt/y)	pre-human	present	actual	(Tmol C/y)	(Tmol C/y)
					(Tmol C/y)	day	(Tmol C/y)		
						(Tmol C/y)			
Value	0.26	0.42	0.7	16	4.1	3.1	10.4	0.25	0.78
Range	0.24 -	0.1 – 0.7	0.4 - 1	12 - 20	3.6 – 4.6	2.8 – 3.4	4.0 - 16.7	0.10 - 0.40	0.48 – 1.12
Reference	This	This	Literature	Literature	This study	This study	Literature	Literature	Literature
	study	study	(1-4)	(5-8)	(model)	(model)	(1 – 10)	(11,12)	(13-15)
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quantitatively modelled PIC concentrations (factor 4) (Supplementary Information S2.3). The reason is the low uncertainty of PIC presence: correct negative classifications (= no PIC present) (75.7 % accuracy) have a lower range of 0 – 0.1 wt% and basins with less than 10 % source carbonate are assumed to have a PIC concentration and error of 0 wt%, reducing variability of results and errors. Positive classifications (= PIC present) are similarly accurate (83.5 %). Moreover, modelled PIC concentrations are within a smaller range than observed values, i.e., the model is biased. Very small values, i.e., PIC < 0.1 wt%, will not drastically affect results, especially because the global flux is dominated by a few large rivers (see below). Miscalculations in PIC-rich rivers, could be more critical to the assessment, e.g., the Rhone river is a comparably small river in terms of sediment discharge, but a major contributor to the

global PIC flux because of high PIC concentrations. However, for most of the important rivers measurements are available, and thus this uncertainty is accounted for (Supplementary Information S2.3). Therefore, the global flux and flux-weighted average concentrations are rather robust. Notably, these uncertainties do not account for inaccuracies in the input datasets.

For instance, the global lithological map (GLiM) has an accuracy of only  $\sim 60\%$  compared to point observations (Hartmann and Moosdorf, 2012). The flux-weighted mean PIC concentration of  $0.26 \pm 0.03$  wt% is lower than the median of PIC-bearing rivers only ( $0.41 \pm 0.01$  wt%, excluding PIC-free rivers) implying  $\sim 40\%$  of the riverine sediment flux to be PIC-free. Both are statistically indistinguishable from the median of observed basinal averages ( $0.35 \pm 0.3$  wt%), covering  $\sim 35\%$  of the global sediment flux (Cohen et al., 2014). Additionally, some authors used mean values, which are more susceptible to outliers caused by small rivers and are typically higher than medians (because of log-normal distributions). High PIC concentrations are rarely found in rivers with high discharge, except for a few large rivers draining markedly dry (e.g., the Nile and Euphrates-Tigris systems) and/or mountainous (e.g., the Indus system) catchments (Fig. 2).

From a total of 3365 catchments considered, the biggest 862 basins contribute ~ 99 % of the total riverine PIC flux, while the biggest 10 catchments, situated in South-East Asia, Arabia, Europe and North America already sum up to ~ 53 %. The Euphrates-Tigris system (13.3 %), the Indus (10.3 %) and the Nile (8.9 %) alone contribute 32.6 % of the total global PIC flux, followed by Yangtze (4.5 %), Salween (4.4%), Colorado (USA, 3.4 %), Rhone (2.8 %), Huanghe (2.1 %), Mississippi (2.0 %) and the Ganga-Brahmaputra system (1.6 %).



**Figure 2** Map of the model results. Point data along the coast are the result of this study (Mean of 794 accepted Monte Carlo simulations). Size scales with the magnitude of the PIC flux (Tmol C/y), based on pre-human sediment discharge) and color is related to PIC concentration (wt %). For comparison, blue colors indicate natural annual mean water discharge (m³/s) (Linke et al., 2019). Conversion of fluxes to Pg/y by a factor 0.012011.

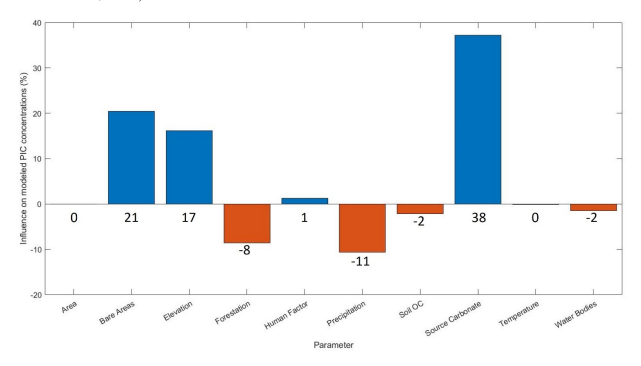
About two-thirds (2.7 Tmol PIC/y) is delivered to the coastal ocean between 15 °N and 45 °N, contrasting riverine DIC, OC, total solute and bulk sediment fluxes (Hartmann et al., 2014; Ludwig et al., 1996; Milliman and Farnsworth, 2011). The anthropogenic reduction of the global PIC flux is dominated by the decreasing contribution of the Nile due to intense damming (~8 of 25 %).

The present PIC flux related to atmospheric dust deposition is  $0.25 \pm 0.15$  Tmol C/y, which is ~ 8 % of the riverine PIC flux and ~ 0.3 % of the total riverine carbon flux (~ 71 Tmol C/y, Supplementary Information S3). Thus, the atmospheric contribution is negligible in global mass balances. PIC related to meltwater discharge and ice-rafted debris from Greenland and Antarctica together contribute another  $0.8 \pm 0.3$  Tmol PIC/y, which is ~ 26 % of the present day riverine PIC flux and ~ 1 % of the total river carbon flux (Supplementary Information S3). In total, ~ 4 Tmol PIC arrive in the ocean annually (~ 5 Tmol when considering natural river discharge, see Table 1).

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#### 4.1 Natural controls of PIC and their variation through time

The relevance of each variable to the model was assessed through the coefficients of correlation and of determination between the individual variable and the median model outcome, being independent of non-linearity and multi-collinearity (Fig. 3). A strong positive influence of SC on PIC concentrations is eminent from these procedures (+38%, Fig. 3). SC includes carbonate from soils ( $\sim3\%$  carbonate on average) and unconsolidated sediments ( $\sim2-4\%$ ) but is dominated by lithological (bedrock) contributions ( $\sim20\%$  on average). This is because terrestrial carbonate weathering is dissolution-dominated, which arises from fast dissolution and high solubility (Lasaga, 1984; Morse and Arvidson, 2002). This contrasts with the precipitation-dominated behavior of silicates, producing clay minerals and oxides characteristic to soil assemblages (Brantley et al., 2008; Lasaga, 1984; Ma et al., 2011; Morse and Arvidson, 2002).

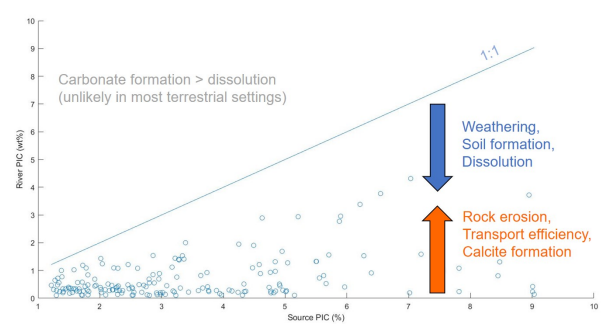


**Figure 3** Relative importance of the different variables to our model results as assessed by the coefficient of determination  $(R^2)$  between the variable in question and the median result of 794

high-quality Monte Carlo simulations. The correlation coefficient gives the direction of influence (orange: negative, blue: positive). Individual values are indicated above the bars. OC: Organic Carbon. Variables as in Table 1.

Generally, riverine suspended sediment is a mixture of source rocks, their solid weathering products (soil and sediment), organic matter and material of anthropogenic origin, with additional in-stream processing. Thus, the differences between SC and PIC may arise from preferential dissolution of carbonates compared to silicates in the weathering zone (= soil) before erosion, and also from in-stream dissolution (Dornblaser and Striegl, 2009), precipitation (e.g., Kempe and Emeis, 1985; Négrel and Grosbois, 1999) and particle sorting during transport (e.g., Bouchez et al., 2011; Garzanti et al., 2011). According to our results, humans did not (yet) influence PIC concentration significantly on a global scale (influence of human factor is only 1%, Fig. 3), while they severely reduced PIC fluxes through their impact on suspended sediment discharge (see section 4.2 Human activities and riverine carbon). A thick soil cover can only develop if chemical weathering rates exceed material removal by erosion (Ferrier and West, 2017; West, 2012), and it is promoted by biological activity. Especially forestation stabilizes the soil, disintegrates pristine rocks and introduces organic acids and ligands, increasing mineral solubilities (Brantley et al., 2017; Calmels et al., 2014). This view is supported by the negative impact of variables in favor of soil formation and dissolution, such as precipitation (- 11 %) and forestation (- 8 %). In contrast, the organic carbon content of soils and temperature, which may influence dissolution kinetics, do not seem to play a major role for PIC concentrations, nor do catchment size or the extent of water bodies, (relatable to the residence time of the particles within the fluvial system). The more prominent influence of (rock) erosion on PIC concentrations is evident from the large influence of related variables, namely, elevation (+ 17 %) and the extent of bare areas (+ 21 %). Rapidly eroding, mountainous terrains are characterized by fast, efficient transport and diminutive sediment

storage (Hilton and West, 2020; Milliman and Syvitski, 1992), limiting both the extent of soil formation (Dixon and von Blanckenburg, 2012; Jenny, 1941) and in-stream dissolution.



**Figure 4** Relationship of river PIC and source PIC. Source PIC (12 % of SC) includes sediment and soil contributions but is dominated by rocks. In-stream dissolution and contributions of weathered material decrease river PIC, while rock erosion has a pronounced positive effect by contributing source rock. Transport efficiency and in-stream precipitation can further enhance PIC concentrations at the river mouth. The unit of Source PIC (%) is wt% PIC in the given percentage of carbonate within the upstream outcrop area.

This interpretation that PIC concentrations increase with erosion is apparently inconsistent with increasing carbonate dissolution following pyrite oxidation and sulfuric acid production upon accelerated erosion as observed in shale-dominated terrains (Bufe et al., 2021; Calmels et al., 2007; Torres et al., 2014). However, we do not only consider shale-rich, but all carbonate-bearing (> 10 %) terrains in this analysis, which could obscure such relationships. Such an apparent inconsistency was noted for other global scale compilations as well (Bufe et al., 2021). Moreover, PIC is predominantly produced by the physical disintegration of pristine rocks and soils, while this same process promotes dissolution and oxidation kinetics. Thus, trends in PIC concentrations and carbonate dissolution do not strictly oppose each other but may even covary in rapidly eroding terrains.

In summary, the rather slowly changing  $(10^3 - 10^5 \text{ y})$  tectonic, physiographic and

lithological settings seem to exert a dominant control on PIC concentrations as demonstrated by the eminent role of SC and elevation in our model. Superimposed on this base-line situation, much faster variations in climatic and vegetation patterns seem to affect the relative contributions of weathered, PIC-poor soil and pristine, PIC-rich source rock.

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Although heavily discussed (Caves Rugenstein et al., 2019; Foster and Vance, 2006; Willenbring and Von Blanckenburg, 2010), many observations suggest that soil formation and/or chemical weathering decreased during cold, dry periods (Berner et al., 1983; Dixon et al., 2016; Jenny, 1941; Schachtman et al., 2019), potentially increasing the ratio of pristine source rock to weathered soil in suspended sediments, thus, the riverine PIC flux. Moreover, glacial activity during cooler periods may accelerate erosion, thus, PIC production and potentially preservation (because of decreased residence times). This is supported by an increase of the detrital carbonate fraction in glacial sediments of the Gulf of Lyon, compared to sediments deposited during interglacial periods (Pasquier et al., 2019). An indication of increased cryogenic PIC deposition in response to ice-sheet dynamics is provided by the socalled 'Heinrich-events', which are unusual accumulations of coarse carbonate-rich detritus in marine sediment (Bond and Lotti, 1995; White et al., 2016). In contrast, thawing permafrost exposes old, but fresh organic matter that is rapidly respired (e.g., Walz et al., 2017), potentially increasing PIC dissolution (Aller, 1982; Archer et al., 1989; Calmels et al., 2014; Oelkers et al., 2011; Zolkos et al., 2018). Consistently, although not solely related to carbonate weathering, an increased riverine export of dissolved inorganic carbon in response to the recent warming was reported from large Arctic rivers (Drake et al., 2018; Zolkos et al., 2020).

However, recent observations and theories challenge this simple view (Caves Rugenstein et al., 2019; Foster and Vance, 2006; Willenbring and Von Blanckenburg, 2010), implying more complex and transitional spatio-temporal dynamics of erosion (Chen et al., 2018; Foreman et al., 2012; van de Schootbrugge et al., 2020) and carbonate weathering

(Gaillardet et al., 2019; Zeng et al., 2019) and, consequently, of riverine inorganic carbon export. Additionally, environmental conditions and, consequently, carbonate dissolution in the (coastal) ocean are expected to change over multiple time-scales, ranging from seasons and decades (Cai et al., 2011; Wallace et al., 2014) to geological time-scales (Broecker, 1982; Ganeshram et al., 2000; Sluijs et al., 2013), with implications for the magnitude and timing of contribution of PIC to oceanic inventories.

### 4.2 Human activities and riverine carbon

Rivers annually deliver about 31.5 Tmol DIC, 19.1 Tmol DOC and 17.4 Tmol POC to the ocean (Table 3). Including 3.1 Tmol PIC/y increases the total riverine carbon export (TC) to 71.1 Tmol C/y, equating a contribution of 4 %, which is within the uncertainty of the estimates excluding PIC (Table 3). An accurate and precise knowledge of the riverine carbon export is necessary to understand the distribution and fate of anthropogenic carbon perturbations (Friedlingstein et al., 2020; Resplandy et al., 2018). Over the past century these riverine carbon fluxes have changed and continue doing so, likely in response to climate change and local human activities, such as industrialization, changes in land-use, hydrology and agricultural practices (Drake et al., 2018; Lambert et al., 2017; Li et al., 2019; Liu et al., 2020; Noacco et al., 2017; Raymond and Hamilton, 2018; van Hoek et al., 2021; Zeng et al., 2019).

The net effect of human activity on riverine sediment discharge is a ~10 % reduction, dominated by damming (Cohen et al., 2014; Syvitski et al., 2005), resulting in an even higher reduction of riverine PIC (~24 %, Table 3) and OC fluxes (~13 %, Maavara et al. (2017)). The differences between those fractions are related to the non-even spatial patterns of riverine carbon and sediment discharge (Ludwig et al., 1996; Milliman and Farnsworth, 2011, Figure 2). Damming also increases the residence time of particles in the riverine realm (Rueda et al., 2006), where PIC and POC are commonly remobilized by dissolution/degradation. However,

organic matter degradation and burial in reservoirs are very heterogeneous and dependent on reservoir ages (Maavara et al., 2017). Low importance for the model (- 2 %) for the extent water bodies, including reservoirs, indicate a rather negligible effect of reservoirs on PIC concentrations on the global scale. Our human factor is not an important predictor in the model (1 %), despite the expected influence of lime-fertilizers (Haynes and Naidu, 1998; Shoghi Kalkhoran et al., 2019; Zeng et al., 2019), of cement (Horvath, 2004), and of human-induced soil erosion, the latter affecting the active weathering zone (Govers et al., 2014). PIC could also be reduced by increasing dissolution through industrial or agricultural acids (Perrin et al., 2008; Webb and Sasowsky, 1994; Wicks and Groves, 1993). Eventually, the lack of resolution between those positive and negative influences in our human factor obscures a clearer relationship. Thus, more detailed studies on the different human influences on riverine carbonate are required. Notably, the human influence is correlated to observed PIC concentrations, but this spurious relationship stems from collinearity of SC and human population, both being high in southeast Asia, Europe and North America, confirming our method is correcting for multi-collinearity.

A 24 % reduction of PIC fluxes equates to only 1.4 % of the total riverine carbon flux (TC). The damming-related decrease in organic carbon fluxes (13 % of OC, Maavara et al. (2017)) results in another 6 % reduction of TC. In contrast, carbonate dissolution-related DIC fluxes likely increase(d) by ~ 13.5 % in the period 1950 – 2100 as a consequence of climate-change and land-use change (Zeng et al., 2019). Such an increase of DIC fluxes would result in a 5.6 % increase of TC, partially compensating the reduction of OC and PIC in terms of total carbon export (total disturbance: - 2 %). This is consistent with the estimation of a somewhat stable riverine TC export as a result of in-stream removal of anthropogenic carbon by POC deposition and respiration (Cole et al., 2007; Regnier et al., 2013; van Hoek et al., 2021).

The bulk anthropogenic effect on total global riverine DIC fluxes remains elusive

(Raymond and Hamilton, 2018) and human activities other than dam-building impact terrestrial and freshwater carbon cycling (van Hoek et al., 2021). Notably, humans also change conditions at the site of riverine PIC deposition: The current coastal ocean acidification in response to anthropogenic emissions and eutrophication (Borges and Gypens, 2010; Carstensen and Duarte, 2019), could enhance PIC dissolution, acting as a heterogeneous buffer (Middelburg et al., 2020).

**Table 3** Summary of the riverine carbon export (in Tmol C/y). DIC: (Amiotte Suchet et al. (2003); Gaillardet et al. (1999); Hartmann et al. (2014); Li et al. (2017); Ludwig et al. (1996, 1998); Meybeck (1982), DOC: (Aitkenhead and McDowell (2000); Dai et al. (2012); Harrison et al. (2005); Li et al. (2019); Ludwig et al. (1996, 1998), POC: Beusen et al. (2005); Galy et al. (2015); Li et al. (2017); Ludwig et al. (1996, 1998); Meybeck (1982), Superscripts: S: By changes in sediment flux only, D: By damming only (Maavara et al., 2017; This study); \*: By climate change and land-use change for carbonate weathering only (Zeng et al., 2019). Human disturbance of TC is the bulk effect as indicated for DIC, PIC, DOC and POC. 'Literature' indicates averages and ranges taken from the above mentioned studies (grey columns). TC represents the sum of our PIC estimate and DIC, DOC and POC estimates from literature. Conversion to Pg/y by a factor 0.012011.

	DIC	PIC	DOC	POC	TC
Modern global river export (Tmol C/y)	31.5	3.1	19.1	17.4	71.1
Percentage of TC	44.3	4.4	26.9	24.5	100
					57.8 to 89.7
Range (Tmol C/y)	26.6 to 36.3	2.8 to 3.4	14.2 to 30.0	14.2 to 20.0	
Human disturbance (%)	+13.5*	-24 <sup>s</sup>	-13 <sup>D</sup>	-13 <sup>D</sup>	- 2
Range (%)	+9.8 to +17.1	- 5.6 to - 39.0	-12.8 to -13.2	-12.8 to -13.2	-4 to + 0.7
Source	Literature	This study	Literature	Literature	This study & Literature

### 4.3 Implications for oceanic mass balances

The fate of the detrital carbonate flux in the marine realm, i.e., PIC burial or dissolution, determines the implication of the global PIC flux for oceanic mass balances (Middelburg et al., 2020). PIC preservation may affect global estimates of marine carbonate burial, while PIC

dissolution would translate to an additional input of Ca, Mg, Sr, C and alkalinity to the marine solute inventories. Because the scientific community lacks a reliable global quantification of these aspects, we discuss the following questions:

1. Where is river PIC deposited?

- 2. Does PIC deposition influence global estimates of carbonate burial?
- 3. Does PIC dissolve and alter oceanic mass balances of Ca, Mg, C, Sr and alkalinity?

#### 4.3.1 Where is river PIC deposited?

On time scales of years to centuries, a major fraction of the riverine suspended matter remains in the estuary (often  $\sim 40$  - 60 %), while the rest is deposited along the shelves and continental slopes with little escape towards the deep sea (Dyer, 1995; Meade, 1972; Wright and Nittrouer, 1995). However, sediment dynamics in river-dominated ocean margins are highly variable in space and time, including deposition near the river mouth and subsequent lateral advection to more calm environments as well as transport towards the slope (Geyer et al., 2004; McKee et al., 2004). Saderne et al. (2019) emphasize that some coastal ecosystems, such as mangrove forests and seagrass meadows, efficiently trap and dissolve such detrital carbonate from external sources.

Global sea level fall under cooler climates (average Pleistocene state) exposes the PIC-rich shelf to erosion, shifting depocenters to the slope, where PIC may be further transported to and/or dissolved in the open ocean (Filippelli et al., 2007; Kump and Alley, 1994; Tsandev et al., 2010). Thus, on larger time-scales (>10<sup>3</sup> years), most of the riverine PIC that does not dissolve on short time-scale (1 to 10<sup>2</sup> years) will be transported to the slope. A significant fraction may, however, have dissolved before re-mobilization or remain at the initial site of deposition (preservation of the former estuary/shelf).

#### 4.3.2 Does PIC deposition influence global estimates of carbonate burial?

Riverine PIC burial on the shelves may be implicitly included in carbonate mass accumulation (CMA) rate estimates derived from carbonate content, density and sediment accumulation rates, although microscopic criteria were established to distinguish biogenic and detrital carbonates (Milliman, 1974). However, hot spots of carbonate burial do generally not coincide very well with hot spots of riverine suspended sediment deposition (i.e., carbonate-poor shelves) (O'Mara and Dunne, 2019). Moreover, the dissolving PIC fraction and the fraction that remains at the initial site of deposition (i.e., is not re-eroded from the former estuary over longer time-scales) does not contribute to estimates of carbonate burial on the slope. Therefore, we believe that the effect of riverine PIC deposition on CMA-derived estimates of biogenic carbonate burial is rather limited. In contrast, carbonate burial estimates derived from mass balances (e.g., van der Ploeg et al., 2019) or solution chemistry (e.g., Chung et al., 2003) are directly affected by PIC dissolution.

# 4.3.3 Does PIC dissolve and alter oceanic mass balances of Ca, Mg, C, Sr and alkalinity?

Marine surface waters are supersaturated with respect to most carbonate minerals (Milliman, 1974; Peterson, 1966). Therefore, provision of carbonate mineral surfaces by PIC discharge, energetically favoring nucleation of these same minerals, may trigger inorganic carbonate precipitation in the water column (Wurgaft et al., 2016). TIC/TOC ratios of sediments from the Huanghe estuary, China (Gu et al., 2009; Yu et al., 2018), and trends in alkalinity/DIC ratios in the marginal Red Sea support this view (Wurgaft et al., 2016). Compared to marine carbonate compensation, PIC will rapidly settle in the shallow coastal ocean and the degree of carbonate saturation varies with depth and across different local environments at the seafloor (Aller, 1982; Boudreau and Canfield, 1993). As chemical conditions, especially pH, vary within the sediment column, carbonate may even be dissolved in the upper parts of the sediment column, but formed in the lower, more alkaline parts (Aller, 1994).

Carbonate dissolution at the sediment-water interface and in diagenetic settings is well

known (Aller, 1982; Archer et al., 1989; Sulpis et al., 2017). In these settings, aerobic degradation of organic matter may drive carbonate dissolution via the production of CO<sub>2</sub> and other acidic compounds (Aller, 1982; Oelkers et al., 2011). Anaerobic degradation produces reduced metabolites such as ammonium, sulfide and iron(II), most of which form strong acids upon upward migration and subsequent re-oxidation in the bioturbated zone, which drastically reduces carbonate saturation (Aller, 1994; Boudreau and Canfield, 1993) and may alter the carbon cycle-coupling of subsequent dissolution (Beaulieu et al., 2011; Huang et al., 2017; Liu et al., 2018; Torres et al., 2017). Carbonate dissolution may also be influenced by biological activity such as seagrass root oxygen loss, sponge boring and bioturbation (Burdige et al., 2008; Mackenzie and Andersson, 2011; Saderne et al., 2019). Substantial riverine PIC dissolution was observed in the maximum turbidity zone of the eutrophic Loire estuary, France (Abril et al., 2003). Moreover, (detrital) carbonate dissolution driven by eutrophication-related bottom water acidification was observed in the Gulf of St. Lawrence, Canada (Nesbitt and Mucci, 2021) and in the Chesapeake Bay, USA (Shen et al., 2019). The proposed total flux of  $\sim 5$  Tmol PIC/y corresponds to  $\sim 10$ Tmol(eq)/a of alkalinity ( $\sim 30$  % of dissolved equivalent), which is  $\sim 30$  % lower than the estimate of Middelburg et al. (2020). Despite integration of groundwater discharge, marine organic matter burial, anaerobic processes and marine silicate weathering, the modern ocean alkalinity budget is marked by an imbalance of ~25 % of the output by carbonate burial (59 Tmol(eq)/y in the coastal and open ocean). Half of this imbalance can be closed by inclusion of riverine PIC fluxes, assuming terrestrial PIC will either dissolve or biases estimates of carbonate burial (Graphical Abstract). Part of the residual imbalance could be attributed to other diagenetic processes in the coastal zone, such as marine alumo-silicates weathering (Gislason et al., 2006; Hong et al., 2020; Jones et al., 2012; Torres et al., 2020) and carbonate diagenesis (DePaolo, 2004; Fantle et al., 2010; Paytan et al., 2021). However, part of the

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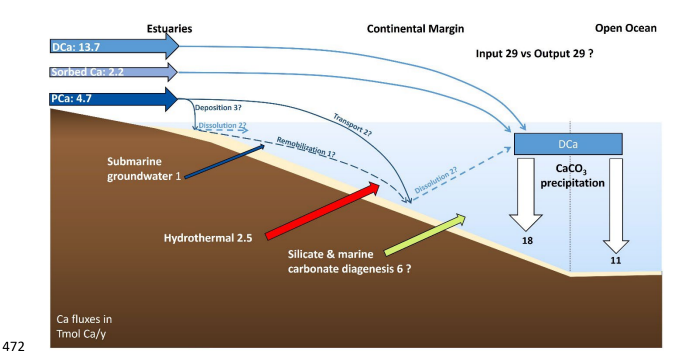
imbalance could be real, considering that the residence time of carbonate ions in the ocean (~ 100 ky) is larger then the time since the last glaciation (Middelburg et al., 2020; Milliman, 1993).

The inputs of ~5 Tmol PIC/y (rivers + cryosphere) further imply ~4.75 Tmol Ca/y, ~0.25 Tmol Mg/y and ~5 Gmol Sr/y, assuming ideal stoichiometry, 10 % dolomite (typical value in *GloRiSe* v1.1) and 1000 ppm Sr in calcite and dolomite. This equates to ~34.7 % (Ca), ~4.6 % (Mg) and ~8.9 % (Sr) of the respective dissolved equivalents, representing the current major input terms of the respective marine mass balances (Berner and Berner, 2012; Krabbenhöft et al., 2010; Mayfield et al., 2021; Tipper et al., 2010, 2006). As dolomites typically exhibit much lower dissolution rates than calcites (Pokrovsky et al., 2005), Mg addition by PIC dissolution is probably even smaller and thus negligible. So far, none of these highly discussed budgets could be consensually balanced, neither at the present state nor in reconstructions of the past – a conundrum persisting already for decades (Berner and Berner, 2012, 1987; Hong et al., 2020; Jones et al., 2012; Krabbenhöft et al., 2010; Mayfield et al., 2021; Middelburg et al., 2020; Milliman, 1993; Shalev et al., 2019; Tipper et al., 2006, 2010).

Riverine PIC input also impacts the Ca-cycle (Figure 5). Apart from riverine dissolved Ca fluxes, submarine groundwater discharge (1 Tmol Ca/yr, Mayfield et al., 2021) and hydrothermal processes (2 - 3 Tmol Ca/yr, DePaolo, 2004) were invoked to balance the high output fluxes by carbonate burial, but still leave an imbalane of 36 %, that can be reduced by further 16 % through consideration of PIC fluxes. The remaining 20 % could be attributed to submarine weathering of volcanogenic silicate debris (Gislason et al., 2006; Hong et al., 2020; Jones et al., 2012; Torres et al., 2020) and/or carbonate diagenesis (3 - 5 Tmol Ca/yr, DePaolo, 2004; Fantle et al., 2010). However, carbonate precipitation in early diagenetic settings currently represents an additional sink of ~ 1 Tmol Ca/yr and ~ 2 Tmol(eq)/y of alkalinity and is related to anaerobic oxidation of organic matter and silicate weathering (Schrag, 2013; Sun

and Turchyn, 2014; Torres et al., 2020). This flux is implicitly included into the mass balance of alkalinity (Graphical Abstract), by reducing the alkalinity source of the diagenetic reflux through carbonate dissolution (from DePaolo, 2004).

The fraction of detrital carbonates in coastal margin sediments was estimated to < 10 and 50 % in the Bengal fan (France-Lanord et al., 2018) and in the Gulf of Lyon (Pasquier et al., 2019), respectively. If the long-term biogenic carbonate burial on the slope is ~2 Tmol C/y (Milliman, 1993) and all riverine PIC (natural: 4.1 Tmol C/y) either dissolves or is transported



**Figure 5** Illustration of the modern ocean calcium budget and how it may be complemented by the inclusion of the riverine PCa flux. The fate of PCa (particulate Ca) in the ocean is, however, uncertain (details in the main text). Fluxes are given in Tmol Ca/y. CaCO3 burial fluxes are from Middelburg et al. (2020), Milliman (1993) and O'Mara and Dunne (2019). River sorbed Ca is from Müller et al., (2021), DCa (dissolved Calcium) and groundwater inputs are from Mayfield et al. (2021), the hydrothermal flux and marine carbonate diagenesis (3 – 4 Tmol Ca/yr)is from DePaolo (2004). The silicate diagenesis flux of Ca is assumed to fill the residual imbalance of  $\sim$  2 Tmol Ca/yr.

to the slope on long time-scales, then 2.1 to 3.9 Tmol PIC/y (51-95%) would need to dissolve in order to match these detrital fractions. This back-of-the-envelop calculation is not a valid quantification of the globally dissolving PIC fraction, but illustrates PIC dissolution may

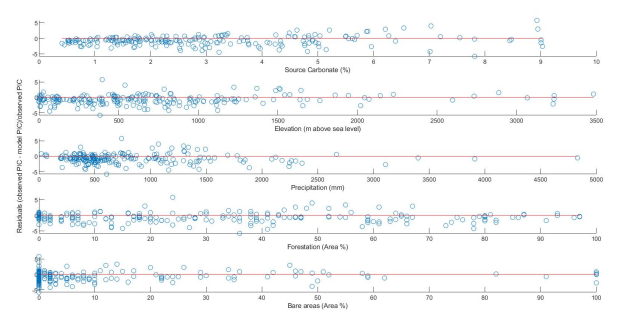
indeed be significant. However, as argued above, the coastal ocean is a heterogeneous region with locally very spatio-temporally variable conditions supporting carbonate dissolution, preservation as well as precipitation. Importantly, recrystallization still leads to an exchange with the marine element and isotope inventories (e.g., Fantle et al., 2010; Kastner, 1999; Paytan et al., 2021). A global estimation of the dissolving PIC fraction should account of this spatio-temporal variability and complex interactions of organic and inorganic particles within coastal sediments. Therefore, consideration of the detrital mineral flux of rivers to the ocean and its isotopic composition may help to solve longwithstanding debates about imbalances in global biogeochemical cycles.

# 5 Next steps

Improving our understanding of the role of PIC in global biogeochemical cycles involves a qualitative understanding and quantitative estimation of its fate in the marine realm, as discussed in the previous section. This might be accomplished by tracing detrital components through isotopic approaches (e.g., France-Lanord et al., 2018; Pasquier et al., 2019) and by diagenetic modeling of riverine particles involving all the important biogeochemical processes driving carbonate precipitation an dissolution (Meister et al., 2022; Torres et al., 2020).

Apart from this, the accuracy of the modeled riverine PIC concentration needs to be improved. As seen from Figure 3, the 'Source Carbonate' is the most important predictor used in the model, but it is also the least well constrained, critically increasing the relative misfit at high concentrations and downward biasing at low concentrations (Figure 6) This might be improved by a more thorough assignment of carbonate content to the different lithological and unconsolidated sediment units through integration of chemical or mineralogical analysis into the corresponding maps. However, limited precision of low concentration measurements might pose an analytical limitation. In highly forested regions, PIC concentrations seem to be

systematically underestimated, which might be related to the type of forest. There is no obvious correlation of residuals to other important predictors (elevation, precipitation and bare areas), suggesting these variables are reasonably well captured. However, as discussed in section 4.2 (Human activities and riverine carbon), the human factor used in this study is rather vague (Table 1) and might mask certain opposing effects, such as liming or locally increased dissolution by acid introduction. Therefore, a detailed analysis of these factors would be warranted.



**Figure 6** Analysis of distribution of relative residuals (fractional deviation from observations) among the most important predictors in the model.

Another important feature may be the accuracy and precision of measured PIC concentrations, as most studies on sediment composition do not involve PIC analysis, or at least do not report the carbonate content that was removed during sample preparation of organic carbon analysis (Müller et al., 2021a). A greater spatial and temporal (time series) coverage of such observations, especially in remote regions, such as Greenland and Antarctica, assist a more accurate upscaling. Similarly, vertically integrated sediment flux measurements, including estimates of bed load transport, could improve the quantification of not only PIC, but all river sediment related biogeochemical fluxes (e.g., Galy and France-Lanord, 2001). Finally,

contributions of Arctic continental ice-rafting and from coastal erosion to PIC fluxes could be significant and lack any reliable estimate.

### **6 Conclusion**

The riverine flux of PIC, i.e., discharge of detrital carbonate minerals, represents a significant, yet mostly unaccounted chemical mass transfer in the Earth system (3.1  $\pm$  0.3 Tmol C/y), currently contributing ~4.4 % to the total riverine carbon export. The pre-human flux was 4.1  $\pm$  0.5 Tmol PIC/y; the 24 % reduction is caused by particle retention in reservoirs, especially of the Nile river. Considering perturbations of riverine particulate and dissolved, inorganic and organic carbon species in concert, the riverine export flux seems to have remained rather stable, while carbon speciation changed.

Although the fate of PIC in the ocean remains quantitatively unknown, oceanic element and isotope inventories of Ca and alkalinity are most probably affected by detrital carbonate dissolution in the coastal ocean, with implications for conclusions deduced from their highly debated, but frequently used mass balances. PIC contributions to the oceanic budgets of Sr and total C are less important and Mg fluxes are insignificant.

Naturally, the concentration of PIC is controlled by catchment topography and surface lithology, i.e., slowly changing tectonic factors ( $10^3 - 10^5$  y scales), but also by climate and vegetation, which are subjected to much faster spatio-temporal variations. Similarly, marine conditions change through time, so that related PIC dissolution may also vary. An additional, significant amount of detrital carbonate ( $0.8 \pm 0.3$  Tmol C/y) is exported from Greenland and Antarctica and responds to ice-sheet dynamics, while eolian contributions can be neglected (at the present). These results imply a response of the global PIC flux to human activity and to natural changes in environmental and climatic conditions, but also to the tectonic evolution of our planet.

# 7 Data and script accessibility 540 All data and scripts used for this study, along with a detailed manual and the supplementary 541 information, can can be accessed via: https://doi.org/10.5281/zenodo.6125880 (DOI: 542 543 10.5281/zenodo.6125880). Data from which figures were generated are can also be found as supplementary information alongside the online version of this article. 544 **8 Competing Interest** 545 546 The authors declare that they have no conflict in interest. 9 Funding 547 This work was carried out under the umbrella of the Netherlands Earth System Science Centre 548 (NESSC). This project has received funding from the European Union's Horizon 2020 research 549 and innovation programme under the Marie Skłodowska-Curie, grant agreement No 847504. 550 551 Funding was also provided by BMBF-project PALMOD (Ref 01LP1506C) through the German Federal Ministry of Education and Research (BMBF) as Research for Sustainability 552 inititative (FONA). AS thanks the European Research Council for Consolidator Grant 771497. 553 10 Acknowledgements 554 We thank Olivier Sulpis, Jens Hartmann, Gibran Romero-Mujalli, Stefan Kempe and Jaap 555 Nienhuis for discussion and advice. Robert Hilton and and anonymous reviewer are thanked 556 for their valuable inputs, significantly improving this work. 557 11 References 558 559 Abbott, A.N., Löhr, S., Trethewy, M., 2019. Are clay minerals the primary control on the oceanic rare earth element budget? Front. Mar. Sci. 6, 1-19. https://doi.org/10.3389/fmars.2019.00504 560

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