- 1 Chemical versus mechanical denudation in meta-clastic and carbonate
- 2 bedrock catchments on Crete, Greece, and mechanisms for steep and high
- 3 carbonate topography

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ABSTRACT

On Crete — as is common elsewhere in the Mediterranean — carbonate massifs form
high mountain ranges whereas topography is lower in areas with meta-clastic rocks. This
observation suggests that differences in denudational processes between carbonate-rich rocks and
quartzofeldspathic units impart a fundamental control on landscape evolution. Here we present
new cosmogenic basin-average denudation rate measurements from both $^{10}\mathrm{Be}$ and $^{36}\mathrm{Cl}$ in meta-
clastic and carbonate bedrock catchments, respectively, to assess relationships between
denudation rates, processes, and topographic form. We compare total denudation rates to
dissolution rates calculated from 49 new and previously published water samples. Basin-average
denudation rates of meta-clastic and carbonate catchments are similar, with mean values of \sim
0.10 mm/a and $\sim 0.13 \text{ mm/a}$, respectively. The contribution of dissolution to total denudation
rate was $\leq 10\%$ in the one measured meta-clastic catchment, and $\sim 40\%$ for carbonate
catchments (~ 0.05 mm/a), suggesting the dominance of physical over chemical weathering at
the catchment scale in both rock types. Water mass-balance calculations for three carbonate
catchments suggests 40-90% of surface runoff is lost to groundwater. To explore the impact of
dissolution and infiltration to groundwater on relief, we develop a numerical model for carbonate
denudation. We find that dissolution modifies the river profile channel steepness, and, together
with infiltration changes, the fluvial response time to external forcing. Furthermore, we show
that infiltration of surface runoff to groundwater in karst regions is an efficient way to steepen
topography and generate the dramatic relief in carbonates observed throughout Crete and the
Mediterranean.

1 INTRODUCTION

In many regions, carbonates form high cliffs and topography in comparison with other sedimentary and sometimes magmatic and metamorphic rocks (Atkinson & Smith, 1976). This observation is ubiquitous throughout the Mediterranean (Godard et al., 2016). For example, on the island of Crete, the landscape is dominated by high and steep carbonate massifs that define the backbone of the island, whereas areas underlain by silica-rich metamorphic units exhibit lower topography and gentler slopes (Fig.1). While to the prominent topographic expression of carbonate bedrock in the Mediterranean is ubiquitous, the reasons for high carbonate topography remain poorly understood, primarily because of the lack of quantification of the processes acting to denude carbonates.

Karstic terrains are often interpreted as landscapes where chemical dissolution dominates over physical erosion (Frumkin, 2013); this paradigm has survived many decades without quantitative testing. The importance of dissolution on the weathering of limestones was recognized as early as the late 18th century (Hutton, 1795) and studies of dissolution rates in limestones have been carried out since the 19th century (Goodchild, 1890; Spring & Prost, 1884). Today, dissolution rate calculations from dissolved loads in rivers and springs are routine (e.g. (Gaillardet, Calmels, Romero-Mujalli, Zakharova, & Hartmann, 2018; Gombert, 2002; Gunn, 1981; Meybeck, 1987); however, the quantification of mechanical rock removal from the same area has proven difficult, except for certain cave systems (Newson, 1971). With the development of cosmogenic nuclide techniques, a tool is available to quantify total denudation, and thus mechanical rock removal when the chemical weathering component is known.

Stone, Allan, Fifield, Evans, and Chivas (1994) first established the use of the ³⁶Cl cosmogenic nuclide to determine bedrock denudation rates in limestones. Previous studies of ³⁶Cl denudation rates have focused on exposed bedrock denudation rates, where dissolution is the

dominant lowering process (Godard et al., 2016; Schaller et al., 2005; Thomas et al., 2018; Xu et al., 2013). However, quantification of carbonate denudation at the catchment scale is needed to integrate the effects of mechanical denudation processes along hillslopes and channels and to understand its impact on relief generation. Ryb et al. (2014) were the first to compare dissolution rates with bedrock and catchment average denudation rates from cosmogenic nuclides in carbonate catchments. They found that in Israel dissolution rates were lower than ³⁶Cl-derived denudation rates and suggested that this discrepancy was due to higher precipitation rates, and thus higher dissolution rates, in the past. These authors argue that the effects of past, higher dissolution rates are incorporated into the cosmogenic nuclide concentrations that integrate millennial timescales but not chemical dissolution rates derived from water that integrate a much shorter time.

The elevated role of dissolution in denudation of carbonates relative to other rock types is related to the development of karst features and landscapes. Karst development alters hydrology by increasing water infiltration and decreasing surface discharge to river systems, which affects denudational processes. We hypothesize that the partitioning of dissolution and mechanical weathering in carbonates scales differently with slope and relief compared to other lithologies and will, therefore, exhibit different topographic responses to external forcings, such as climate and tectonics. Moreover, several studies have suggested that dissolution can affect the shape of cave stream profiles (Covington, Prelovšek, & Gabrovšek, 2013; Springer, Wohl, Foster, & Boyer, 2003; Woodside, Peterson, & Dogwiler, 2015), however we have limited understanding of how altitude-dependent changes in dissolution rate influence river longitudinal profiles in carbonate settings (Covington et al., 2013).

In this study, we address these knowledge gaps by measuring total denudation, dissolution rate and water mass balance at the catchment scale in carbonates and meta-clastic Phyllite and Quartzite units (PQ) on Crete and construct river profile models to assess the dominant factors (e.g. physical and chemical denudation, and subsurface water infiltration) contributing to elevated and steep carbonate terrain compared to quartz-rich rocks in the Mediterranean. To partition total denudation between chemical and mechanical weathering in karst landscapes, we analyze seven ³⁶Cl samples of detrital carbonate sands from Crete and 49 water samples from published and new data (Fig.1). To test how substrate erodibility might contribute to differences in topography and relief between carbonates and other lithologies, we determined denudation rates from 16 10 Be detrital samples in the PQ unit and assessed the partitioning between chemical and mechanical weathering in this unit with an approach similar to the karst areas. We use a numerical model of carbonate denudation to explore the effects of carbonate denudation processes and hydrology on the generation of topography. We explore how the infiltration of runoff to groundwater in limestone karst areas and dissolution change the relief and the shape of river profiles and how climatic and tectonic conditions influence carbonate topography.

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2 BACKGROUND

The island of Crete occupies a forearc high above the Hellenic subduction zone, a long-lived system in which Africa is presently subducting below Eurasia at ~ 36 mm/a (Reilinger et al., 2006). From the Cretaceous to the early Miocene, subduction of continental slivers or attenuated crustal lithosphere and oceanic domains resulted in a thick nappe pile (van Hinsbergen et al., 2005). The Permian to Oligocene Plattenkalk and Jurassic Trypalion carbonate

underwent high pressure – low temperature (HP-LT) metamorphism (Fassoulas, Kilias, & Mountrakis, 1994; Seidel, Kreuzer, & Harre, 1982; van Hinsbergen et al., 2005). These nappes were exhumed by subsequent extension during the Neogene and now unmetamorphosed, Pindos and Tripolitza carbonate units are juxtaposed against HP-LT metamorphic rocks of the Phyillite – Quartzite (PQ) and Plattenkalk units (Rahl, Anderson, Brandon, & Fassoulas, 2005; van Hinsbergen & Meulenkamp, 2006). Miocene to recent multidirectional extension has created numerous sedimentary basins filled with marine and terrestrial deposits (Peterek & Schwarze, 2004; van Hinsbergen & Meulenkamp, 2006; Zachariasse, van Hinsbergen, & Fortuin, 2008). Arc normal and parallel extension of the upper crust is ongoing, as evidenced by active normal faults bounding some of the mountain ranges on Crete (Caputo et al., 2010; Gallen et al., 2014). The PO and carbonate units on Crete are from various positions within the former nappe pile, and there is no clear relationship between stratigraphic position in the former nappe pile and modern elevation in the landscape. Carbonate massifs on Crete reach 2500 m and are characterized by steep flanks with high relief that bound internal areas of lower relief (Fig. 2a). The three main massifs (Lefka Ori, Psiloritis and Dikti) contain several internally drained basins and also have areas that are nearly vegetation free, comprised of natural vegetation of semi-shrub

deserts (Fig. 2a) (Bohn & Gollub, 2000). Areas underlain by the PQ bedrock form mountainous

landscapes but with elevations generally below 1000 m and are more densely vegetated with

olive, kermes oak-mastic and pine forest (Fig. 2b) (Bohn & Gollub, 2000).

units were underthrusted below the Carboniferous to Middle Triassic PO unit, and all units

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3 METHODS

3.1 Cosmogenic nuclide denudation rates

To compare denudation rates in silicic and carbonate bedrock, we collected quartz- and carbonate-rich fluvial sands to measure ¹⁰Be and ³⁶Cl concentrations, respectively, using Accelerator Mass Spectrometry (AMS) at ETH Zurich. Samples for ¹⁰Be were gathered from rivers draining the meta-clastic PQ unit that is mainly exposed in western Crete. We collected ³⁶Cl samples from carbonate catchments spanning a wide range of elevations and relief to see if denudation scales with topographic metrics, which would imply a contribution of mechanical denudation.

3.1.1 Detrital ¹⁰Be denudation rates

We collected 16 samples of fluvial sands with grain size 0.25 - 0.71 mm for ¹⁰Be analysis. All sampled catchments lie within the meta-clastic PQ unit. Most sampled catchments are underlain almost entirely by the PQ unit, except samples WC-616-11 and WC-616-16 where other rock types, such as carbonate and flysch, span ~40% of the catchment. We separated quartz from the bulk sediment by magnetic separation and repeated etching in hydroflourosilicic and diluted hydrofluoric acid. A ⁹Be carrier was added before dissolving the quartz with hydrofluoric acid. Beryllium was extracted using standard column chromatography (Bierman et al., 2002). The ¹⁰Be/⁹Be isotope ratio was measured by 500kV TANDY at ETH Zürich and calibrated using S2007 N ¹⁰Be standard (Christl et al., 2013). Blank corrected ¹⁰Be concentrations were used to calculate erosion rates based on the equations from Brown, Stallard, Larsen, Raisbeck, and Yiou (1995) (detailed AMS data in Tab. S5). Production rates were calculated in a pixel-based approach with the scaling scheme of Stone (2000), assuming a rock density of 2.7 g/cm³ and attenuation lengths for neutrons, slow and fast muons from Braucher, Merchel, Borgomano, and Bourlès (2011) on a 30 m SRTM elevation model, averaged for every catchment and corrected

for topographic shielding. Denudation uncertainties include analytical uncertainty and 2.5%, 50% for the production uncertainties of neutrons and muons, respectively (Lupker et al., 2012).

3.1.2 Detrital ³⁶Cl denudation rates

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Seven samples of fluvial sediments with grain size 1-4 mm were collected for catchment average denudation rates in areas dominated by carbonates using ³⁶Cl. We sampled catchments within the Plattenkalk, Trypalion, and Pindos units. The grain size, which is larger than that used for ¹⁰Be, was chosen to avoid aeolian material transported by strong coastal winds in some catchments and to be able to sample small steep catchments lacking sand size river sediments. The samples were crushed to grain size < 0.5 mm. Meteoric ³⁶Cl was removed by etching with 2 M HNO₃ and repeated rinsing. A ³⁵Cl- enriched spike (Ivy-Ochs, Synal, Roth, & Schaller, 2004) was added, and the samples were dissolved with HNO₃ (Prager, Ivy-Ochs, Ostermann, Synal, & Patzelt, 2009). Measurements of ³⁶Cl were conducted at the Laboratory of Ion Beam Physics 6 MV TANDEM AMS system at ETH-Zürich using a gas-filled magnet to separate the isobar ³⁶S (Vockenhuber, Miltenberger, & Synal, 2019) and calibrated with the internal K382/4N standard (Christl et al., 2013). The AMS data with and without blank correction are shown in the supplementary table S6. The chemical composition of the target and the bulk rock was determined by separate ICP-MS analysis of the bulk sample and the undissolved sample remnant (if not all the material dissolved). The chemical composition of the bulk samples and target used for calculation is reported in the supplement (Tab. S4).

³⁶Cl concentrations and carbonate chemistry were used to calculate denudation rates following the approach of Schimmelpfennig et al. (2009) with the scaling by Stone (2000) and a topographic shielding correction. The scaling factors of nucleonic and muonic production were calculated with CRONUS (Marrero et al., 2016) using the mean basin latitude and altitude

calculated by a pixel-based approach on a 30m SRTM digital elevation model. The Schimmelpfennig et al. (2009) denudation rate calculation does not include an uncertainty propagation. However, the current version of CRONUS is incapable of calculating erosion rates and uncertainties for low concentration ³⁶Cl samples. Therefore, we used the relative error for our samples calculated with a reference total nuclide concentration of 4e⁵ at/g (close to the mean of our measured concentrations) and ensured that the effect of total nuclide variation on the expected relative error is negligible. The samples were run in CRONUS with this reference nuclide concentration while all other sample specific input parameters (nuclide concentration uncertainty, sample chemistry) were adjusted. This approach allowed us to incorporate the analytical uncertainty from the AMS measurement and Cl-concentration as well as the production rate uncertainties. Additional uncertainties incorporated for pressure, sample thickness, bulk density and effective attenuation length are 10 hPa, 0.01 cm, 0.1 g/cm³ and 10 g/cm², respectively.

3.2 Dissolution rates

3.2.1 Water sample collection and analysis

We collected a set of seven water samples from springs, rivers, and Lake Kournas. We used our data in conjunction with 42 water samples from springs, wells, rivers and lakes from Crete, collected by the National Water Monitoring Network, Greece and compiled from Papaioannou (2007) and Kallianis and Chatzitheodorou (2003) (Tab. S1). We collected two samples per site for cations and anions in HDPE bottles and filtered samples on-site through a 0.2 μm filter. Alkalinity was measured in the field with a sulfuric acid titrant and a bromcresol green - methyl red indicator. Cation samples were acidified, and anion samples kept refrigerated until measurement. To gauge the source altitude of water samples, we measured δ¹⁸O on each

sample. See supplement for details about ion concentration and $\delta^{18}O$ measurements. Cation and anion concentrations were corrected for precipitation input by using the lowest [Cl⁻] measurement from the Zaros spring, which is sourced by meteoric water from the high-elevation (> 1000 m), evaporite-free Psiloritis Mountains and assuming all [Cl⁻] is precipitation derived. The other ions were then corrected by scaling with seawater ratios (Stallard & Edmond, 1981).

3.2.2 Dissolution rate calculation

We use a trimonthly, three year time series from seven sites collected by the National Water Monitoring Network, Greece (Tab. S1) to assess chemostatic behavior (Godsey, Kirchner, & Clow, 2009) (*i.e.*, constant chemical concentrations even as discharge varies) of the respective water source (Fig. 3). Variations of $[Ca^{2+}]$ are < 10% and $[Mg^{2+}]$ < 20% throughout the year; therefore, we assume chemostatic behavior of the cations and anions for our analysis. Time-averaged concentrations of Ca^{2+} and Mg^{2+} (in mg/l) are used to calculate average carbonate dissolution rates D (in mm/a) with the following equation:

219 (1)
$$D = \frac{\left([Ca] + \frac{[Ca]}{M_{Ca}} * M_{CO_3} \right) * (P - AET)}{\rho_{Calcite} * 10^6} + \frac{\left([Mg] + \frac{[Mg]}{M_{Mg}} * M_{CO_3} \right) * (P - AET)}{\rho_{Dolomite} * 10^6}$$

Where precipitation (P) and actual evapotranspiration (AET) are in mm/a and the densities (ρ) in g/cm³. M_{Ca} , M_{Mg} , and M_{CO3} are the molar masses of the respective ions. The factor 10^6 converts the dissolution rate into units of mm/a. This calculation assumes dissolution mainly at or close to the surface, an assumption verified by Gunn (1981). Saturation indices are calculated in PHREEQC (Parkhurst & Appelo, 1999) assuming chemical equilibrium.

Information on the potential discharge ($Q_{pot} = P - AET$) available for dissolution is required to determine the amount of total dissolution. To calculate Q_{pot} , we used the average monthly P data provided by WorldClim at 1 km spatial resolution (Fick & Hijmans, 2017),

which was averaged for the catchment area of the water sample. We used annual estimates of AET derived from the Parameterization of Vegetation Indices for Evapotranspiration (PaVI-E) model (Helman, Givati, & Lensky, 2015). The PaVI-E model uses spectral vegetation indices from the MODerate resolution Imaging Spectroradiometer (MODIS) onboard the Terra satellite and information on water vapor flux from the eddy covariance tower international net (FLUXNET) to derive empirically annual AET from 2000 to 2016 at 1 km spatial resolution (a detailed description of the estimation of AET is in the supplement). Analogous to P, AET was averaged for each catchment. Finally, Q_{pot} was calculated at 1 km resolution from P and AET. Groundwater recharge areas were estimated from topography and local geology (Fig. S1). These areas were used to calculate Q_{pot} for springs and aquifer samples, while surface water catchment areas were used to calculate Q_{pot} for surface water samples.

However, recharge areas of karst aquifers commonly differ from their topographic delineation (e.g. Malard, Sinreich, and Jeannin, 2016); therefore we applied a 10% uncertainty on P and 20% uncertainty on AET (together with the 5% analytical uncertainty of the reported concentrations). To further test the reliability of the water fluxes incorporated into the dissolution rates we also calculated dissolution rates based on (1) the mean P and AET on all Cretan carbonate areas between 23.8 and 25.8° longitude and (2) the mean P and AET for every carbonate massif separately from, which our samples are recharged (Tab. S1). To assess the contribution of chemical weathering in meta-clastic catchments, we calculated the chemical rock removal by dissolution for the Topolia catchment (location in Fig. 4a). From this water sample, we calculated the total dissolved flux from this catchment and compared it to the total flux predicted by the ¹⁰Be measurement at the same location.

3.3 Numerical modelling of carbonate denudation

To test the effect of different denudation processes acting in carbonate landscapes, we use a 1D numerical model of topographic evolution. The model domain is uplifted at a uniform rate (U = 0.2 mm/a), and denudation is calculated with two separate process rules - mechanical erosion (E) and dissolution (D) to simulate chemical weathering rate. The rate of surface elevation change can then be described as:

$$(2) \frac{dz}{dt} = U - E - D$$

257 E can be described by the detachment-limited stream power model,

258 (3)
$$E = K * Q_{pot}^{m} * S^{n}$$
,

- where K is a coefficient incorporating erodibility of the substrate, climate, and hydrology, Q_{pot} is
- the potential discharge (P AET), and S is the local channel slope. m and n are empirical
- 261 constants depending on basin hydrology, channel geometry and erosion process (Whipple &
- Tucker, 1999). We use the common values of n = 1 and m = 0.5 for our numerical models
- 263 (Whipple, Hancock, & Anderson, 2000).
- Assuming that runoff rapidly reaches saturation, limited by temperature and pCO₂, the
- 265 mass loss can be expressed as a surface lowering rate due to dissolution (in mm/a) after White
- 266 (1984) as

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$$(4) D = \frac{1}{\rho 10\sqrt[3]{4}} * \left(\frac{K_c K_1 K_{CO_2}}{K_2}\right)^{\frac{1}{3}} pCO_2^{1/3} * (P - AET)$$

- where ρ is the density of limestone (in g/cm³), K_c , K_1 , K_2 and K_{CO2} are the equilibrium constants
- of the calcite-CO₂-H₂O system and P and AET are in mm/a (not to be confused with K, the
- erodibility parameter). The change of surface elevation can then be described as:

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$$(5)\frac{dz}{dt} = U - K(Q_{pot}(z) * I)^m S^n - \frac{1}{\rho 10\sqrt[3]{4}} * \left(\frac{K_c(z)K_1(z)K_{CO_2}(z)}{K_2(z)}\right)^{\frac{1}{3}} pCO_2(z)^{\frac{1}{3}}$$

$$(P(z) - AET(z))$$

- We introduce the infiltration parameter (I) to the stream power portion of the equation to
- analyze the infiltration of runoff to groundwater $(I = \frac{Q}{Q_{pot}})$, as commonly observed in karst
- landscapes. To quantify the magnitude of infiltration, we calculated water budgets for three
- 276 catchments on Crete from the P, AET, and gauging data (see Table S2).
- In the case of a steady-state $(\frac{dz}{dt} = 0)$, equations (2) and (3) can be combined and solved
- for the steady-state local channel slope S and the infiltration parameter can be added:

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$$(6) S = \left(\frac{U - D}{K}\right)^{1/n} * (Q_{pot} * I)^{-\frac{m}{n}}$$

The form of this equation is the same as Flint's (1974) law:

$$(7) S = k_s * Q^{-\frac{m}{n}}$$

- where k_s is the steepness index. As such $k_s = \left(\frac{U-D}{K}\right)^{\frac{1}{n}}$. In similar expressions that exclude the
- 283 dissolution term in equation 2, the first term in equation 6 takes on the form $\left(\frac{U}{K}\right)^{\frac{1}{n}}$ and many
- researchers have used the relationship between k_s values to infer tectonic signals from
- 285 topography. Following these previous studies, we refer to the first term on the right-hand side of
- equation 6 as the dissolution steepness index, k_{sd}.

288 4 RESULTS

- 289 4.1 Cosmogenic denudation rates
- 290 4.1.1 Denudation rates and patterns

Denudation rates derived from ¹⁰Be in meta-clastic catchments range from 0.048 to 0.26 mm/a with a mean of 0.1 mm/a (Tab. 1), which are lower than late Pleistocene coastal uplift rates (0.2-1 mm/a) (Gallen et al., 2014; Ott et al., 2019) (Fig. 4). The denudation rates show no obvious spatial pattern among the meta-clastic PQ unit (Fig. 4). ³⁶Cl-derived catchment-averaged denudation rates from carbonate sands in limestone dominated catchments range from 0.052 – 0.56 mm/a (Tab. 1, Fig. 4). The highest value of 0.56 mm/a is in the Samaria Gorge, one of Europe's deepest gorges with ~ 2 km of relief (Fig. 4a). The next highest denudation rate is 0.18 mm/a from a similar gorge nearby. The low concentrations measured for the Samaria Gorge represent an outlier in our dataset, which is not unexpected. The Samaria gorge has vertical walls of several hundreds of meters that are prone to rockfall and landsliding; therefore, a contribution of rockfall or landslide material may explain the low ³⁶Cl concentration, and we consequently removed this data point from our analysis. Carbonate denudation rates along the high mountain ranges are slightly elevated compared to denudation rates in meta-clastic catchments (Fig. 4a, b). Carbonate denudation rates along the lower relief topography of northeastern Crete are similar to the rates from western Crete (Fig. 4c). Generally, the denudation rates in the carbonate catchments are similar to the rates in meta-clastic catchments but are slightly higher in areas of steep and high topography (Fig. 4a, b).

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4.1.2 Correlation between denudation rates and topographic/climatic metrics

We compare our denudation rates with topographic and climatic metrics for each catchment (Fig. 5). We use a correlation coefficient matrix to assess the strength of parameter relationships within the dataset (Fig. 5g). The correlations between denudation rate and other catchment metrics within the 10 Be dataset are generally weak (2 < 0.4) (Fig. 5 a-f). However, the meta-clastic catchments exhibit similar topography and erosion rates and thus do not span

enough range to assess correlations adequately (Fig. 5). Carbonate catchments, in contrast, cover a larger topographic and climatic gradient than meta-clastic catchments and exhibit stronger correlations of denudation rates with several topographic metrics as well as precipitation (Fig. 5). The correlation coefficient matrix shows the same pattern, with weaker correlations in meta-clastic catchments and stronger correlations ($r^2 > 0.5 \mid r^2 < -0.5$) in carbonate catchments (Fig. 5g). For example, in carbonate catchments, the mean catchment elevation scales inversely with AET (Fig. 5g), likely due to decreasing vegetation cover and consequently of plant transpiration (usually the main component in AET) at higher elevations. The effect of orographic precipitation is noted by the correlation of precipitation with mean elevation and total relief (Fig. 5g). P-values for correlation significance are sample size dependent, an effect manifested in the small 36 Cl dataset (n = 7) that shows some p-values that are higher than in the larger 10 Be dataset (n = 16) (Tab. S7). However, the p-values for correlations between topographic metrics and denudation rate in the 36 Cl catchments are lower compared to 10 Be catchments, and as such the correlation between 36 Cl denudation rate and local relief is still significant at a 90% level (Tab. S7).

In summary, the good correlations between denudation rate and topographic metrics in carbonate catchments likely imply a causal relationship between topography and denudation. The available data in meta-clastic catchments span too narrow a range to assess mutual relationships between denudation rates and topographic metrics adequately.

4.2 Water chemistry and dissolution rates

The chemical signature of Cretan water samples varies among the different water sources (Fig. 6). Water samples from carbonate catchments have a mean [Ca²⁺] of 63 mg/l and a mean [Mg²⁺] of 20 mg/l. River water has higher [Ca²⁺] and [Mg²⁺] compared to subsurface water from wells and springs. Groundwater samples also have less variability in their concentrations. The

values of pCO₂ also vary by water source. Most river samples were oversaturated and we also calculate the pCO₂ at which they would have been in equilibrium (Fig. 6c). The pCO₂ for spring and groundwater are higher as for river water. If we take into account potential degassing of CO₂ in river water and calculate the pCO₂ at which the river water would have been in equilibrium, this relationship is reversed (Fig. 6c), however these theoretical pCO₂ should not be taken at face value becacuse the Sr/Ca ratios of > 2 (ppb/ppm) indicate likely secondary carbonate precipitation (Bickle, Tipper, Galy, Chapman, & Harris, 2015) (Tab. S1). The values for [Na⁺], [SO₄²⁻] and [Cl⁻] vary widely between different catchments and are potentially related to small amounts of evaporites in certain catchments (Tab. S1). We focus our analysis on [Ca²⁺] and [Mg²⁺] because we are mostly interested in the rates of carbonate dissolution.

[Ca²⁺] and [Mg²⁺] vary not only with water source but also with catchment elevation (Fig. 7). [Ca²⁺] decreases with elevation of the recharge area, as does the saturation index with respect to calcite (Fig. 7a). Crete displays a clear increase of precipitation with elevation due to orography and a simultaneous decrease in evapotranspiration (Fig. 5). In terms of [Ca²⁺] fluxes or dissolution rates, there is little change with elevation because the reduced concentrations at high altitude are offset by increased water flux due to the coupled effects of increased precipitation due to orography and reduced evapotranspiration at higher elevations on Crete (Fig. 7c). A matrix of all correlation coefficients (analogous to Figure 5g) for our water sample data is available in the supplement (Fig. S2). Our calculated carbonate dissolution rates vary between 0.016 and 0.150 mm/a, with a mean of 0.048 \pm 0.011 mm/a. The different methods to estimate water fluxes for dissolution described in the methods produced overall similar results with some variations in individual catchments but an overall mean dissolution rate of ~ 0.05 mm/a (Tab. S1). To assess the chemical weathering component in the meta-clastic PQ unit we compared the

total denudational flux from ¹⁰Be to the total dissolved load in the Topolia catchment (Tab.2). We found that < 10% of the denudational flux is from chemical weathering, comparable to other siliciclastic catchments (Dixon & Blanckenburg, 2012).

4.3 Water budgets

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For three Cretan catchments with long-duration gauging data, we calculated annual water budgets spanning years 1996 – 2003 for two catchments and 1996 – 2001 for the other basin and determined the approximate ratio of surface to subsurface runoff for each year in each catchment (Tab. S2). We found that on average, 40 to 90% of the potential runoff left the catchments as subsurface flow. The two catchments with lower loss rates of 40 and 57% are in mixed lithologies, whereas the 90% loss rate is from a carbonate-dominated catchment. We lack water budgets for meta-clastic catchments; however, we assume subsurface water flow to be minor similar to other regions underlain by metamorphic rocks (Driscoll, 1987). Since we do not have detailed information about springs and their discharges downstream and upstream of sampling points that could modify the water budgets, these results are subject to considerable uncertainties. However, the finding of large amounts of subsurface flow in the highly karstified Cretan landscape is not surprising, given the occurrence of multiple dry valleys, poljes, and large aquifers. Based on these findings, we use an infiltration of 70% (I = 0.3) of potential discharge as input for our carbonate denudation models in section 5.3.2. It is important to note that the high uncertainties associated with this estimate of I do not influence any of our main conclusions. Below 500 m we use a linear decrease from 70 to 0% infiltration loss, mimicking the increase of infiltration as the topography rises higher above the karst water table. The karst water table below the Lefka Ori carbonate massif on Crete, for example, has been shown to be more than 1 km below the land surface in some locations (Nikolaidis, Bouraoui, & Bidoglio, 2013).

5 ANALYSIS AND MODELLING

This study presents an analysis of water and cosmogenic samples from Crete to quantify the mechanical and chemical components to total denudation and relief production in meta-clastic and carbonate catchments. We will first discuss the processes that influence altitude-dependent dissolution rates and how we can use dissolution rates together with the total denudation rates in the carbonate and meta-clastic catchments to quantify catchment-scale chemical and mechanical weathering. We then use numerical simulations to assess the primary factors controlling the various topographic expressions of carbonate and meta-clastic bedrock catchments assuming that carbonate denudation occurs by both mechanical erosion and chemical weathering, while silicic-clastic units erode mostly through mechanical erosion. Simulations were then used to predict the response of carbonate catchments to different forcings.

5.1 Processes controlling water composition of carbonate catchments on Crete

We observed higher [Ca²⁺] and [Mg²⁺] surface water concentrations compared to water traveling via the subsurface (Fig. 6). These differences could be related to changes in the CO₂ availability for dissolution. Soil CO₂ concentration is typically an order of magnitude higher than atmospheric values (Brook & Box, 1984), and thus offers an important source of carbonic acid for carbonate dissolution. Soil CO₂ concentrations scale with primary productivity, particularly in semi-arid environments (Cotton & Sheldon, 2012), which typically decreases with increasing elevation (Daubenmire, 1943; Humboldt, 1806). In Crete, natural vegetation varies from forests dominated by olive trees at sea level to pine and cypress-dominated woodlands at intermediate altitudes, and grasslands or barren land at high elevations (Bohn & Gollub, 2000). The origin of the water sampled from groundwater aquifers and springs are the high elevation mountain

ranges, as evidenced by more negative $\delta^{18}O$ values similar to the high elevation water samples (Tab. S1). At the top of these mountains, vegetation is quite sparse with mostly barren lands with patches of thin soil (Fig. 2a). Therefore, CO₂ availability for dissolution is mostly restricted to atmospheric CO₂ (and not soil CO₂). Water in streams may represent a mixture between such high elevation low [Ca²⁺] water and water sourced from lower elevations where more CO₂ for dissolution is available. This interpretation is further supported by the high pCO₂ – typical for soil systems- at which the supersaturated water samples would be in equilibrium (Fig. 6c). However, pCO₂ values for the rivers are difficult to interpret due to supersaturation suggesting CO₂ degassing and Sr/Ca (ppb/ppm) ratios > 2 indicative of secondary carbonate precipitation (Bickle et al., 2015).

Calcite solubility increases with decreasing temperature (Plummer & Busenberg, 1982) and therefore, the potential for dissolution should increase with elevation if all other parameters are held constant. However, the aforementioned processes override the effect of a general calcite solubility increase with lower temperatures. Decreases of [Ca²⁺] with altitude due to a decrease in soil CO₂ have also been observed in river and spring water in Swiss Jura mountains (Calmels, Gaillardet, & François, 2014) and limestone dissolution tablet measurements in the Alps (Plan, 2005). However, the runoff available for dissolution increases due to orographic rainfall and lower evapotranspiration rates at high altitudes. On Crete, this effect almost balances the decrease of [Ca²⁺] and leads to a less pronounced decrease in the carbonate dissolution rate with altitude (Fig. 7c).

5.2 Chemical versus mechanical denudation in carbonates and clastics

Our calculation of total dissolved loads in the Topolia meta-clastic catchment (Tab. 2) showed that < 10% of the total denudational flux in this catchment is through the dissolved load. This result is supported by Gaillardet, Dupré, Louvat, and Allègre (1999) and Larsen et al. (2014) who estimated that globally chemical weathering in silicates is responsible for < 5% of the denudation. Therefore, we assume that chemical weathering in the PQ bedrock is minor and that surface lowering is largely achieved by mechanical weathering.

The average calculated dissolution rate for carbonate catchments is ~ 0.05 mm/a (Tab. S1). This rate is nearly a factor of 3 lower than the average carbonate denudation rate of 0.13 mm/a, but implies that dissolution shoulders a greater proportion of denudation in carbonate catchments ($\sim 40\%$) compared to the PQ catchment. Figure 8 shows the difference between total denudation rates and dissolution rates on Crete. Denudation rates in carbonate catchments increase with topographic metrics such as local and total relief, normalized channel steepness (k_{sn}) and elevation (Fig. 5g), whereas carbonate dissolution rates do not show this correlation. The correlation of total denudation with topographic metrics indicates that the discrepancy between total denudation rates and dissolution rates is best explained by mechanical erosion in the carbonate catchments.

A previous study by Ryb et al. (2014) explained a similar discrepancy through higher precipitation (MAP) rates in the past. A higher paleo-MAP would result in a higher water flux available for dissolution and thereby increase paleo-dissolution rates. Average catchment erosion rates in our case integrate over a time window of 4 – 6 ka. Tritium concentrations of well and spring data range from 0.7 to 5 tritium units and indicate a mix of submodern (recharged prior to 1952) and modern water for most aquifers (Polychronaki, Pavlidou, & Zouridakis, 2009). Our water samples are snapshots of modern to submodern conditions depending on the water source.

Ryb et al. (2014) observed the same discrepancy between total denudation and dissolution rates and argued that this reflects a higher paleo-MAP, based on the weak correlation of slope with their denudation rates. However, paleo-MAP reconstructions for 6 ka show no relevant difference from modern precipitation rates on Crete (Hijmans, Cameron, Parra, Jones, & Jarvis, 2005). Moreover, our carbonate denudation rates do scale with topographic metrics, whereas a simple increase in paleo-MAP is not expected to cause relief-dependent changes in denudation rates. For these reasons, we suggest carbonate dissolution rates have remained relatively steady over the time-scale integrated by our cosmogenic nuclide derived erosion rates and that the discrepancies between total denudation and dissolution in carbonates primarily reflect the roles of physical weathering and erosion.

The correlation of denudation rates with precipitation and the lack of correlation between dissolution rates and precipitation (Fig. S4) illustrates that increases of carbonate denudation with precipitation are not necessarily the result of increased dissolution due to a higher water flux. Alternatively, discrepancies between erosion and dissolution can be caused by increased mechanical erosion in steep and high landscapes.

5.3 Mechanisms for high carbonate topography and low PQ topography

5.3.1 Erodibility in carbonates and meta-clastic rocks and landscape response

Coastal uplift data from western and southern Crete show that the PQ-unit and the carbonate mountain ranges are often fault bound (Ott et al., 2019; Wegmann, 2008) but that despite similar uplift rates the topographic expression of the carbonate mountains is much higher. Another explanation for higher and steeper carbonate topography is a difference in erodibility (K) between the PQ-unit and the different carbonate units. Under this reasoning, carbonate catchments would have a lower K-value that would require them to steepen to maintain

denudation rates similar to those in the meta-clastic catchments (equation 3). However, K incorporates climatic and hydrologic parameters as well as substrate erodibility within the simple stream power framework. On Crete, we do not expect a substantial climatic difference between our catchments, whereas our water budgets imply marked differences in hydrology between meta-clastic and carbonate catchments. The substrate erodibility is relatively challenging to quantify directly, and the carbonates exhibit landforms diagnostic of karst hydrology. We, therefore, choose to assess the impact of hydrologic behavior by incorporating hydrologic parameters into the discharge term Q of equation 5 and explore the effects of infiltration of surface runoff in karst areas on topography. We can use this characterization of infiltration to test if the difference in hydrology between units is sufficient to explain differences in relief or if an additional difference in substrate erodibility is required. Also, the combined action of dissolution and mechanical erosion in the carbonate catchments should scale differently with elevation than denudation in the PQ unit that is dominated by mechanical processes, demonstrating the need to explore plausible mechanisms of relief generation.

5.3.2 Landscape response to infiltration and dissolution

To explore the landscape response to dissolution and the infiltration of runoff to subsurface flow, we model the evolution of topography over time explicitly accounting for mechanical and dissolution-related denudation as well as the effect of subsurface infiltration. The erodibility parameter K was chosen to be 10^{-5} to mimic the fluvial relief of Cretan mountain river systems. To calculate the portion of denudation due to carbonate dissolution, we scale AET, precipitation, temperature, and soil pCO_2 with altitude. Scaling of AET and precipitation was accomplished by fitting functions to P/AET versus altitude plots for satellite data on Crete within carbonate terrain (Fig. S2). Soil CO_2 was scaled with AET based on the relationship from Brook,

Folkoff, and Box (1983). The range of log₁₀(pCO₂) predicted by the model (-3.05 to -2.40) is similar to the values observed in our water samples (Fig. 6c). Soil CO₂, effective precipitation, and temperature are then used to calculate a carbonate lowering rate with equation (4). We run the model until a steady-state topography is achieved. The empirical equations used, a listing of parameters and a detailed scheme of the model setup are in the supplement (Tab. S3, Fig. S3).

Figure 9 shows the steady-state topography of 1D river profiles for different denudation processes in a homogeneously uplifting mountain range (for the transient response see Fig. S5). Precipitation on Crete increases with elevation whereas AET decreases. These trends lead to a strong net increase in runoff with altitude that is available for both dissolution and mechanical river incision (Fig. 9b). The decrease of AET with elevation is linked to a decrease of primary productivity and therefore scales with the production of soil CO₂. Dissolution scales with the cube root of CO₂ but linearly with the availability of water (see EQ. 4). Therefore, dissolution in our model increases with elevation despite a strong decrease in soil CO₂. This increase of carbonate dissolution rate predicted by the model is not observed in our dissolution rates from Crete (Fig. 7c); however we lack water samples from streams at high elevations.

The dissolution rates calculated within our model from equation four are landscape averages. Studies by Palmer (1991) and Covington et al. (2013) showed that dissolution rates in cave streams can be higher than these landscape average rates. However, all water samples we collected from local streams were saturated with respect to calcite and therefore would not permit additional dissolution along the stream. In this case, only where undersaturated water from springs or the surroundings enters a stream, can additional dissolution occur. Our model dissolution is a representation of locally adding undersaturated water from the surroundings, but does not include additional dissolution, nor discharge related to potential springs at the base of

carbonate massifs. Hence, we note that there is uncertainty on the dissolution rates along the stream profile which inhibits a direct relation of the modelled magnitude of the dissolution effect to natural stream profiles. However, the uncertainty of in-stream dissolution rates does not affect any of the conceptual conclusions drawn below.

Model runs that include dissolution exhibit lower relief and gentler slopes compared to models with only detachment limited stream power incision (Fig. 9a). The contribution of dissolution to the surface lowering acts to decrease the river gradient because less mechanical erosion is required to balance the uplift. This is in agreement with findings from Springer et al. (2003) who observed this behavior in stream profiles in West Virginia, USA.

The variation of dissolution rate with altitude will also alter the shape of a river profile by changing the dissolution channel steepness index k_{sd} along the profile. While standard detachment limited stream profiles will exhibit a constant channel steepness k_s , altitude dependent dissolution processes will lead to a non-uniform channel steepness k_{sd} . If dissolution is not considered but is important in the longitudinal profile evolution, such variations in steepness might be interpreted as changes in channel concavity. A dissolution rate that increases with altitude, as the one we modelled will lead to a progressive upstream decrease in k_{sd} because an increasing portion of denudation will be through slope-independent dissolution (Fig. 9d). We also used the empirical equation derived from the linear fit to our carbonate dissolution rates with altitude (Fig. 7c). In this model run dissolution decreases with altitude due to infiltration (with a constant value above 1300 m to avoid dissolution rates that reach zero). When the empirical dissolution rate is applied, k_{sd} is lower than in a stream power model (SPM) and the difference reduces with altitude (Fig. 9d).

While dissolution acts to reduce the steepness of river profiles, the opposite occurs when infiltration of surface runoff is included in the model (Fig. 9a). The associated reduction in stream discharge generates a steady-state topography that is higher and steeper by a factor of $I^{\text{-m/n}}$ (~ 1.8 in our model) compared to the standard stream power model (based on the integration of equation 7). Equation (1) implies that a decrease in discharge will need to be compensated by an increase of slope to achieve the same erosion rate. Therefore rivers in carbonate catchments that typically experience high amounts of infiltration to the karst system will steepen their gradient to maintain the same denudation rate, consistent with the hypothesis of Gallen and Wegmann (2017) posed for rivers in south-central Crete. The PQ unit is predicted to denude through mechanical processes without substantial infiltration of runoff and therefore are less steep than carbonate catchments to match the same rock uplift. This mechanism provides one possible explanation for the limited range in the values of the topographic metrics observed in the PQ-catchments (Fig. 5).

These results also imply that topography in carbonates will respond differently to changes in uplift rate compared to areas with little subsurface infiltration and without dissolution. The landscape response time is defined by the timescale required for a perturbation (e.g. a change in tectonic or climatic forcing) to propagate from the river outlet to the channel head (Howard, 1994). Infiltration and dissolution will change the original response time equation from Whipple and Tucker (1999) to:

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$$(9) \tau = \int_0^x \frac{dx}{K * Q_{pot}^m * I_{up} *^m S^{n-1} + D}.$$

Infiltration during the uplift of a limestone mountain range (decreasing I) above the karst water table will increase the time of landscape adjustment. Here, we have modified I to I_{up} to highlight that amount of infiltration loss upstream of the knickpoint sets the response time. This

rate of upstream infiltration rate might change as the knickpoint migrates. In a block experiencing a sudden uplift increase without a prior karst network, infiltration rates are likely to increase upstream and will therefore mainly affect the response time of the upper channel reaches. Fabel, Henricksen, Finlayson, and Webb (1996) showed this behavior in an Australian carbonate stream, where the development of karst lead to the abandonment of the stream channel during normal flow conditions and therefore almost stalled the migration of stream knickpoints. In our example model, the response time of a 10 km long mechanically denuding basin (e.g. PQ -unit) is 7.5 Ma. This increases to 13.7 Ma for a carbonate catchment and 11.5 Ma when an increase in orographic precipitation is included. However, these calculations were performed with a constant I_{up}, whereas changes of infiltration rates are likely to happen in carbonate terrains; depending on the initial stage and evolution of the karst network. High infiltration rates to the karst system will facilitate the growth of internally drained basins in the central parts of the uplifted area due to the long time needed for integration. When dissolution is considered without infiltration, it has the opposite effect of decreasing the response time to external forcing. In our model, this decrease is < 1 Ma. The long response time of streams experiencing infiltration explains the limited fluvial integration of the internal parts of carbonate massifs on Crete and the occurrence of internally drained basins since these areas started uplifting after 10 Ma (van Hinsbergen & Meulenkamp, 2006).

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6 DISCUSSION

High carbonate massifs on Crete are probably related to a mix of all the effects mentioned above. Some of the ranges are bound by active faults (Fig. 1) and might have exhibited higher uplift rates in the past. For instance, the southern Lefka Ori Mountains are

bound by the same active faults as the mountains in western Crete within the PQ unit; however, we do not know if other, now inactive, faults caused higher uplift rates in the Lefka Ori in the Pliocene or Early Pleistocene. Yet, Pleistocene uplift rates along the southwest coast of Crete, show that the mountain ranges PQ mountain ranges in western Crete and the Lefka Ori carbonate massif experience uniform coastal uplift rates (Ott et al., 2019) while exhibiting >1.5 km more relief in the carbonates. As these carbonate mountain ranges started uplifting, substantial infiltration of up to 90% of surface runoff increased the response time to tectonic forcing and fostered the establishment of internally drained basins in the center of the massifs. Internally drained basins such as the Lassithi Plateau are now common within the uplifted Cretan carbonate massifs.

Our results imply that in a Mediterranean climate, mechanical rock removal is responsible for more than half of the surface lowering and that uplift above the groundwater table induces infiltration that needs to be compensated by steepening topography. This explanation provides a mechanism for why carbonate areas in the Mediterranean typically exhibit high local relief and areas with different tectonic and climatic forcing such as the Appalachian Mountains in the US or southern Ireland and England comprise low relief carbonate topography (Gallen, 2018; Mills, 2003; Simms, 2004). Carbonates in regions with high localized uplift and unfavorable conditions for dissolution will respond to the uplift forcing mainly by mechanical rock removal that will require more steepening as the topography is elevated above the groundwater table. In contrast, areas of lower uplift and conditions more favorable for dissolution, rock uplift will be balanced mostly by dissolution. Steep slopes are not required for steady-state in such conditions (U minus D close to zero and therefore low k_{sd} value).

Variable dissolution and hydrology along stream profiles as they occur in carbonate settings are parameters that could be implemented into landscape evolution models. By constraining the hydrology and dissolution behavior of a study area, landscape evolution models and river profile analysis could be used to extract information such as tectonic and climatic forcings from otherwise hard to analyze karstic terrains. On Crete, the limited number of karst springs suggests that a substantial fraction of the infiltrated water leaves through groundwater submerged springs into the sea. However, in some areas on Crete and many worldwide groundwater daylights in large karst springs. These sudden changes of discharge along a river profile are another mechanism that will greatly influence the shape of stream profiles and show the importance of characterizing the hydrologic behavior of a karst system before analyzing the topography.

7 CONCLUSIONS

We present new water chemistry and detrital 36 Cl-denudation data from Crete to partition the total denudation in carbonate catchments between chemical dissolution and mechanical erosion. Water chemistry data show that on Crete, $[Ca^{2+}]$ and $[Mg^{2+}]$ decrease with elevation and are lower for aquifer water derived from high elevation massifs due to less vegetation and higher infiltration rates. An increase in runoff with altitude slightly offsets this decrease in dissolution rates resulting in a more moderate decrease with elevation. These carbonate dissolution rates account for $\sim 40\%$ of the total surface lowering of 0.13 mm/a, as measured with 36 Cl in carbonate catchments on Crete. The remainder is accomplished through mechanical rock removal as implied by the correlation of k_{sn} , relief and elevation with total denudation rate. In meta-clastic catchments denudation rates were similar (~ 0.1 mm/a) with less than 10% of chemical rock removal.

Our numerical model of carbonate denudation showed that there are distinct differences between siliciclastic or metamorphic bed channels and carbonate bed channels that are affected by both water infiltration into karstic features and dissolution of bedrock. Dissolution affects river profiles through chemical lowering of channel beds. Changes in dissolution rate with elevation, thus alter the shape of river profiles by changing the channel steepness. In contrast, the infiltration of runoff to groundwater in karstic areas will increase the maximum elevation and relief of a landscape. The infiltration in karst areas will increase the response time to external forcing and facilitate the establishment of internally drained basins in the center of an uplifted block, whereas dissolution will decrease the response time of a landscape. This interpretation implies that high and steep carbonate topography in the Mediterranean is likely related to high local uplift rates, where dissolution is not able to balance uplift, and mechanical rock removal requires steep slopes as runoff is mostly lost to the subsurface.

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8 FIGURE CAPTIONS

Figure 1: (a) Topography of Crete and sampling sites for cosmogenic nuclides and water. The rectangle depicts the locations of the close-ups in figure 4. Inset: Location of Crete within the Mediterranean. (b) Simplified geologic map of Crete after Creutzburg (1977) with faults and outlines of sampled basins for catchment average denudation rates. (c) Cumulative hypsometric curve of the main lithologies on Crete. The colors for lithologic units in (b) and (c) are identical. Note that carbonate lithologies are disproportionately over-represented at high elevations.

Figure 2: Typical appearance of the high elevation areas within the carbonate massifs (a) and meta-clastic PQ catchments (b) on Crete.

Figure 3: Schematic diagram of the steps involved in the carbonate dissolution rate calculation.

Figure 4: Catchment average denudation rates from cosmogenic nuclides. Catchments with dominantly PQ bedrock where ¹⁰Be was measured are outlined in black, ³⁶Cl carbonate catchments in red. The locations for (a), (b) and (c) are indicated in figure 1a. The catchment labels correspond to the numbers in Tab. 1.

Figure 5: Correlation between denudation rates and catchment metrics. (a – f). Local relief was calculated with a 500 m radius on a 30m Shuttle Radar Topography Mission (SRTM) digital elevation model (DEM).Regression lines and their correlation coefficients are shown in black for ¹⁰Be catchments and red for ³⁶Cl catchments. (g) Matrix of correlation coefficients within the dataset (HI – Hypsometric Index). The upper triangle with the black outline corresponds to data from the ³⁶Cl catchments. The lower triangle shows data from the ¹⁰Be catchments. The catchment metrics are listed on the sides and the correlation coefficients are colored by their respective value. All correlation coefficients were calculated with uncertainties reported in the methods section. A matrix of p-values for this table is provided in the supplement (Tab. S7).

Figure 6: Boxplots of $[Ca^{2+}]$ (a), $[Mg^{2+}]$ (b) concentrations and $log_{10}(pCO_2)$ for different types of water sources with boxes representing 25% and 75% percentiles and the mean (red bar) (all n = 15; river n = 6, drill n = 4, spring n = 5). For locations with time-series, only one-time averaged value was incorporated. The dark grey boxes include the theoretical pCO_2 at which

samples oversaturated with respect to calcite would have been in equilibrium prior to degassing. We calculated the theoretical pCO₂ values for supersaturated samples by adding acidity from CO₂ until saturation is achieved. Subsequently, the new pH estimate of the solution is used in PhreeQC with all other inputs held equal to calculate the pCO₂ prior to degassing.

Figure 7: Calcium concentration (a), calcite saturation index (b) and calculated carbonate dissolution rate (c) against the mean elevation of the estimated recharge area with 1 σ error bars.

Figure 8: Total denudation rates from ³⁶Cl measurements against local relief with regression line (red line) and correlation coefficient. Blue circles depict carbonate dissolution rates calculated from water data. We interpret the difference between the two rates as the effect of mechanical weathering within the carbonate catchments.

Figure 9: Carbonate denudation model. (a) Steady state elevations of river profiles for 4 different denudation process combinations. SPM – stream power model incision. (b) Rates of different input parameters for the SPM & infiltration & dissolution model run. P, AET and soil CO₂ are empirical functions of altitude and therefore change along the model domain with the uplifting topography. (c) Rates of total, mechanical denudation and dissolution of the same model as in (b). (d) Dissolution channel steepness index k_{sd} for the SPM model, a model with the calculated dissolution rate and a model using the regression line equation from figure 7c as dissolution-scaling. Note, stream profiles without dissolution will have a constant k_s value along stream, while stream profiles with altitude dependent dissolution will change their channel steepness along profile.

Table 1: Cosmogenic nuclide samples, AMS concentrations and derived denudation rates.

				catchment	mean				
#	Lab ID	Latitude	Longitude	area	elevation			[¹⁰ Be]	Erosion rate
		°N	°E	km²	m			10 ⁴ atoms/g	[mm/a]
1	WC-616-2	35.4298	23.924	14.1	333.8			3.72 ± 0.32	0.088 ± 0.013
2	VVC-010-2	33.4238	23.324	14.1	333.8			3.72 ± 0.32	0.088 ± 0.013
	WC-616-3	35.4518	23.7903	21.6	612.7			2.63 ± 0.23	0.154 ± 0.020
3		05.4545	22 7264	20.4	560.0			4.05 . 0.40	
4	WC-616-4	35.4515	23.7864	28.1	560.8			4.35 ± 0.18	0.090 ± 0.010
	WC-616-5	35.4054	23.6829	30.9	576.3			8.04 ± 0.33	0.049 ± 0.005
5									
6	WC-616-6	35.4056	23.7094	15.4	543.3			3.33 ± 0.26	0.115 ± 0.015
	WC-616-7	35.3297	23.5582	34.7	577.9			3.78 ± 0.24	0.104 ± 0.013
7		00.0207	10.0001	3	07710			0.70 2 0.2 1	0.20 . 2 0.020
	WC-616-9	35.4166	23.5698	9.8	490.2			4.38 ± 0.28	0.084 ± 0.010
8	WC-616-10	35.273	23.6664	18.6	570.3			4.71 ± 0.23	0.083 ± 0.010
9	VVC-010-10	33.273	23.0004	18.0	370.3			4.71 ± 0.23	0.083 ± 0.010
	WC-616-11	35.2599	23.6022	14.4	497.3			6.88 ± 0.42	0.059 ± 0.010
10									
11	WC-616-12	35.2735	23.6606	34.2	638.4			5.70 ± 0.26	0.072 ± 0.008
	WC-616-13	35.2708	23.6853	70.9	576.2			6.32 ± 0.28	0.063 ± 0.008
12									
42	WC-616-14	35.2861	23.8176	20.5	637.5			1.59 ± 0.14	0.262 ± 0.035
13	WC-916-15	35.4633	23.8323	51.4	590.2			4.07 ± 0.19	0.099 ± 0.013
14	VVC-310-13	33.4033	23.0323	31.4	330.2			4.07 ± 0.13	0.033 ± 0.013
	WC-617-16	35.1635	24.4742	123.3	490.3			2.90 ± 0.43	0.128 ± 0.023
	Sample	Latitude	Longitude	catchment area	mean elevation	[CaO]	[CI]	[³⁶ CI]	Erosion rate
	1 112		G :					1	
		°N	°E	km²	m	%	ppm	10 ⁵ atoms/g	[mm/a]
15	01.647.5	05.515	05 05 05		45.5		00.55	0.50 . 5.55	0.400 / 5.515
16	Cl-617-2	35.342	25.6565	8.9	481.3	47.4	90.96	3.59 ± 0.26	0.109 ± 0.014
	Cl-617-3	35.3333	25.6213	16.1	453.0	48.8	8.17	4.89 ± 0.28	0.052 ± 0.007
17									
	Cl-617-6	35.2787	24.8296	57.2	1408.7	50.3	3.98	3.60 ± 0.17	0.141 ± 0.016

18									
	Cl-617-8	35.377	24.1351	24.1	1213.2	37.1	205.66	5.83 ± 0.80	0.16 ± 0.041
19									
	Cl-617-10	35.3764	24.201	87.4	727.2	48.5	79.87	3.31 ± 0.21	0.128 ± 0.016
20									
	Cl-617-15	35.1975	24.0572	26.3	1061.6	38.9	168.06	4.74 ± 0.44	0.175 ± 0.031
21									
	WC-616-8	35.2707	23.9638	38.4	1213.0	50.0	51.48	0.99 ± 0.23	0.556 ± 0.068

Tab. 2: Chemical weathering contribution to the total denudation in the Topolia meta-clastic catchment from the total dissolved load (TDS). Calculation of annual total and chemical denudational flux for 1 m² in the catchment.

TDS	Р	AET	Q = P- AET	Q in m³	Solutional flux	
mg/l	mm/a	mm/a	mm/a	m³/a	kg/m³/a	
58.16	751	553	198	0.198	0.0115	
			Be rate	Rock density	Total flux	% Chemical
			mm/a	kg/m³	kg/m³/a	
			0.049	2700	0.1323	8.71

Figure 1

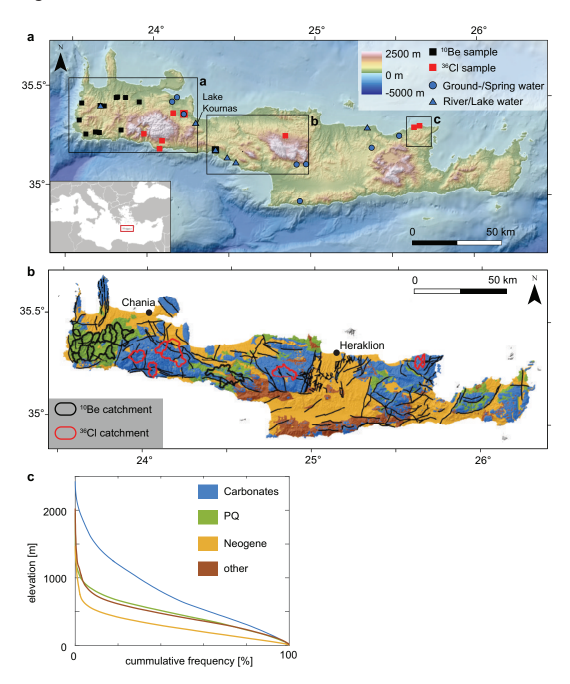


Figure 2





Figure 3

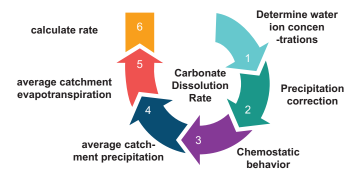


Figure 4

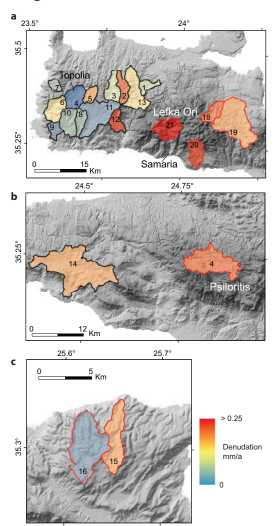
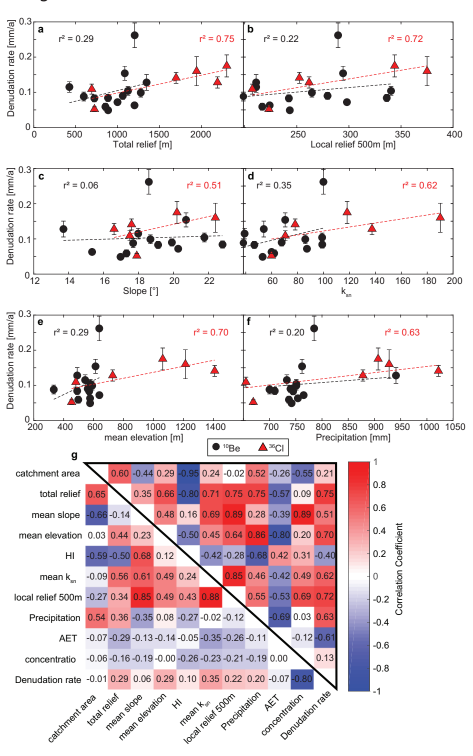


Figure 5





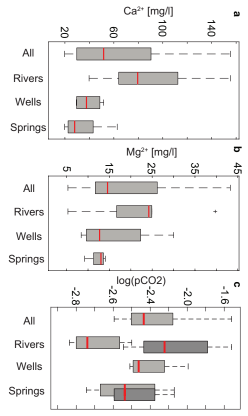
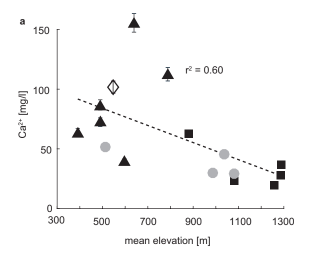
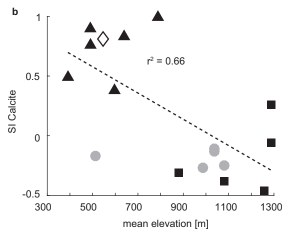


Figure 7





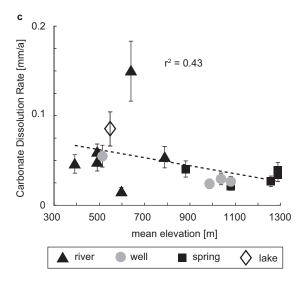


Figure 8

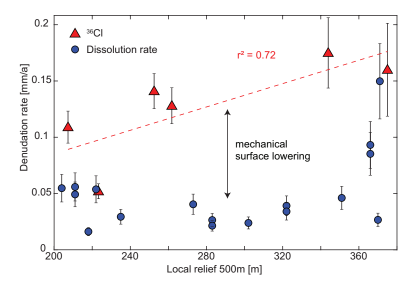


Figure 9

