



AperTO - Archivio Istituzionale Open Access dell'Università di Torino

Glacier melting increases the solute concentrations of himalayan glacial lakes

This is the author's manuscript				
Original Citation:				
Availability:				
This version is available http://hdl.handle.net/2318/1638803 since 2019-12-19T17:44:31Z				
Published version:				
DOI:10.1021/acs.est.6b02735				
Terms of use:				
Open Access				
Anyone can freely access the full text of works made available as "Open Access". Works made available under a Creative Commons license can be used according to the terms and conditions of said license. Use of all other works requires consent of the right holder (author or publisher) if not exempted from copyright protection by the applicable law.				

(Article begins on next page)

1 Glacier melting increases the solute concentrations of Himalayan glacial lakes

2 Franco Salerno,^{*1,3} Michela Rogora,² Raffaella Balestrini,¹ Andrea Lami,^{2,3} Gabriele A. Tartari,² Sudeep Thakuri, ¹

3 Danilo Godone,⁴ Michele Freppaz,⁵ and Gianni Tartari^{1,3}

- 4 ¹ CNR Water Research Institute (IRSA), Via del Mulino 19, Brugherio (MB) 20861, Italy
- 5 ² CNR Institute of Ecosystem Study (ISE), Largo Tonolli 50, Verbania Pallanza (VB) 28922, Italy
- 6 ³Ev-K2-CNR Committee, Via San Bernardino, 145, Bergamo 24126, Italy
- 7 ⁴CNR IRPI Geohazard Monitoring Group, Strada delle Cacce, 73, 10135 Torino, Italy
- 8 ⁵Università di Torino, DISAFA and NatRisk, Largo Paolo Braccini 2, 10095, Grugliasco (TO), Italy
- 9 *Correspondence to Franco Salerno (salerno@irsa.cnr.it) Water Research Institute National Research Council (IRSA-
- 10 CNR), Via del Mulino, 19, 20861 Brugherio, MB, Italy TEL +39 039 21694221 FAX +39 039 2004692

11 ABSTRACT Over the past two decades, we observed a substantial rise in ionic content that was mainly determined by 12 the sulfate concentration at 20 remote high elevation lakes located in central southern Himalaya. At LCN9, which was 13 monitored on an annual basis for the last twenty years, the sulfate concentrations increased over 4-fold. Among the 14 main causes, we exclude a change in the composition of wet atmospheric deposition, as well as a possible influence of 15 decrease in seasonal snow cover duration, which could have exposed larger basin surfaces to alteration processes. 16 Glacier retreat likely was the main factor responsible for the observed increase of sulfate concentrations. We attribute 17 this chemical changes mainly to the sulfide oxidation processes that occur in subglacial environments. Moreover, we 18 observe that the weakened monsoon of the past two decades has only partially contributed to the lakes enrichment 19 through runoff waters that are more concentrated in solutes or lowering the water-table, resulting in more rock exposed 20 to air and enhanced mineral oxidation.

21 INTRODUCTION

High mountain ecosystems are especially vulnerable to climate change, and lakes in particular may act as sentinels.^{1,2} An evaluable opportunity for a fine-scale investigation is particularly evident on the south slope of Mt. Everest (Nepal), which is one of the most heavily glacierized parts of Himalaya³. At the same time, this region is the most densely populated by glacial lakes in the overall Hindu-Kush-Himalaya range⁴. In this region, Lami et al.⁵ observed a significant relationship between the increase in the annual temperature recorded in the area and the enhanced conductivity detected in two glacial lakes located above 5000 m a.s.l. in the last decades, but without inferring any undergoing physical process or source of ionic release. At the same time, an increase in conductivity and sulfate concentrations has been observed in a few lakes of the European Alps, but in this case, the changes have been specifically ascribed to the solute release from an active rock glacier as a response to climate warming⁶. An increase in sulfate from melting ice during drought years is also reported by Williams et al.⁷ in the outflow of a rock glacier in the Colorado Front Range and by Mast et al.⁸ in southwestern Colorado. Even in this last case the authors consider temperature as the main climatic driver of change and sulfate as largely derived from pyrite (watershed weathering sources), although the undergoing physical processes are only supposed.

35 In this study, after exploiting the recently reconstructed⁹ daily temperature and precipitation time series and the 36 available accurate tracing of the glacier shrinkage in the region¹⁰, we discuss changes in the water chemistry of 20 37 remote lakes located on the south slopes of central Himalaya that were sampled approximately after twenty years, starting in the early 1990s. Furthermore, during this period, two selected lakes (named Lake Cadastre Number (LCN)¹¹ 38 39 9 and 10) were monitored yearly. We consider several drivers of changes of the in-lake chemistry, including 40 temperature, precipitation, atmospheric deposition, rocks and soil weathering processes (periglacial environment), 41 seasonal snow cover duration, and, in particular, the role of glaciers (subglacial and surrounding periglacial components). With regard to permafrost, little is known about its distribution in Himalaya.¹² Therefore the subsurface 42 43 ice and its possible degradation is not included in this analysis, although its influence on in-lake chemistry evolution cannot be excluded as reported by other authors^{13,14} in the Colorado Front Range. 44

45 MATERIAL AND METHODS

46 Study area. The monitored glacial lakes are located in Sagarmatha (Mt. Everest) National Park (SNP), Nepal, which 47 lies in the central southern Himalayas (Fig. 1a, 1c). The SNP (1148 km²), crossed by the Khumbu valley, is the world's highest protected area, extending from an elevation of 2845 to 8848 m a.s.l..^{15,16} According to Searle et al.¹⁷, the 48 49 Khumbu valley substratum is dominantly made up of crystalline rocks (e.g., sillimanite gneisses calc-silicates, 50 amphibolites and K-feldspar augen gneisses) and leucogranites that intrude the metasediments as dykes, sills and 51 bodies. The overlying Everest series (weakly metamorphosed shales and pelites with limestone bands), the Yellow band 52 unit (limestones, marbles and calc-silicates) and the Ordovician limestones are only present in the eastern part of the 53 basin. Extensive moraine debris of different ages, alluvial deposits and glacio-fluvial deposits host sub-surface aquifers 54 in close connection with the hydrographic network.

The land cover classification shows that almost one-third of the territory is characterized by temperate debriscovered glaciers, with 75% of glacier surface area lying between 5000 m and 6500 m a.s.l.¹⁰ and with less than 10% of the park area being forested^{15,16}. The soils in the high valleys are primarily Entisols.¹⁸

Salerno et al.¹⁹ reported the presence of 170 glacial lakes and provided an accurate description of their topographic 58 59 characteristics. Most of them are small (median surface of 0.01 km²) and within an elevation zone ranging from 4800 to 60 5300 m a.s.l.. Table 1 shows the main morphometric features of the 20 survey lakes. They have been selected to cover wide elevation (4466-5427 m a.s.l.) (Fig. 1b), basin slope (9°-35°), and basin aspect (120°-231°) ranges. The mean 61 62 glacial coverage is 18%, with three basins without glaciers (LCN40, LCN66, and LCN70). Two sub-basins (LCN10 and 63 LCN67) does not have glaciers within the sub-basin, which directly drain into the lakes, while in this case their 64 upstream lake basin (LCN9 and LCN68, respectively) presents a glacier coverage (all data are reported in Table SI1). 65 Details on these lakes and their basins can be found in Tartari et al.²⁰. All glaciers lying within the survey lake basins 66 are debris-free and without clearly detectable accumulation and ablation zones.

67 The climate is characterized by the monsoons, with a prevailing S-N direction.²¹ During the last twenty years at the Pyramid meteorological station (5050 m a.s.l., Fig. 1c)⁹, which is located just 500 m from LCN9 (Fig. 1c), 90% of 68 69 the precipitation is concentrated during June-September, and the probability of snowfall during these months is very 70 low (4%), whereas the amount of snowfall at an annual level reaches 20%. Snow as a percentage of total precipitation 71 increases with the elevation gradient justifying the wide glacier coverage in SNP. Recently, the daily temperature and 72 precipitation time series of the last twenty years (1994-2013) were reconstructed by Salerno et al.⁹ for these altitudes 73 (Fig. 1b), which correspond to the median elevation of the 20 survey lakes. The total annual cumulated precipitation at 74 this elevation is 446 mm, with a mean annual temperature of -2.45 °C. The mean annual air temperature has increased 75 by 0.9 °C since the early 1990s. The significant increases were experienced just for the months of April, November, and December.9 As the air temperature increased, the average elevation of the freezing line (0 °C isotherm) has moved 76 77 upward. The major changes occurred in April, where it moved upward by 225 m, passing from around the mean 78 elevation of glaciers terminuses to the mean elevation of the surfaces of the same glaciers located in the basins of the 20 79 survey lakes (Fig. 1c). The increase in air temperature observed at these high elevations during April fits with the 80 warming reported by Pal and Al-Tabbaa²², who observed that only April shows significant changes in all Indian regions 81 and the West Himalaya (1901-2003 period). As for precipitation, a substantial reduction of rainfall (47%) and in the 82 probability of snowfall (-10%) has been observed in this area over the last twenty years⁹. According to Yao et al.²³ 83 there is strong evidence of the general monsoon weakening even in the overall Himalayas.

Water chemistry analysis. The chemical composition of 20 remote lakes located in SNP was evaluated in the period between 1990 and 1997 (hereafter early 1990s) and successively from 2008 to 2012 (hereafter early 2010s). Lake sampling was carried out in autumn, at the end of the monsoon season. Samples were collected from the lake shore, possibly close to the lake outlet. Moreover, for the lakes LCN 9 and LCN10, sampling and chemical analyses have been performed annually since 1990 (1991 for LCN9), usually in late September or October, with the exception of the 89 missing years 1995-96 and 1998. For these lakes, three samples were generally collected along the water column from 90 an inflatable boat in the deepest part of the lake. Here we took their average. For detailed information on the 91 hydrochemical and biological characteristics of these lakes, see the work of Lami et al.⁶.

92 All samples were collected using prewashed plastic bottles and stored at 4 °C for successive chemical analysis; these 93 were performed during the 1990s by the CNR Water Research Institute and subsequently by the CNR Institute of 94 Ecosystem Study (Italy). Several intercomparisons have been conducted to ensure data quality and comparability 95 between the data produced by the two laboratories. Samples were analyzed for pH, conductivity, total alkalinity (using 96 the Gran method for acidimetric titration), ammonium, total nitrogen, total phosphorus and reactive silica (by 97 spectrophotometry), sulfate, nitrate, chloride, calcium, magnesium, sodium and potassium (by ion chromatography). 98 Data were checked for internal consistency by means of the ionic balance and the comparison between measured and 99 calculated conductivity. Details on the analytical methods and on the quality controls adopted in the laboratory can be 100 found in Lami et al.⁵.

In this paper, we focus on conductivity values, as an index of total solute concentrations of lakes, H^+ , and major ions: calcium + magnesium (Ca²⁺+Mg²⁺), sulfate (SO₄²⁻) and bicarbonate (HCO₃⁻). These are indeed the prevailing ions in all sampled lakes, representing more than 80% of the total ionic content (Fig. SI1). We assume alkalinity as corresponding to bicarbonate ions, because at circum-neutral pH, as those of the study lakes, dissolved HCO₃⁻ is the dominant species contributing to alkalinity.

106 Morphometric and hydrological analysis. Seasonal snow cover duration of each survey lake basin has been attributed 107 as a mean for the 2002-2012 period of the daily MODIS (Moderate Resolution Imaging Spectroradiometer) imagery 108 (pixel 250 m) (http://modis.gsfc.nasa.gov/). The glacier surface areas within the survey lake basins were derived for the 109 early 1990s from Landsat 5 TM (17 November 1992, pixel 30 m) and for the early 2010s from Landsat 11 ETM (30 November 2011, pixel 15 m) remote imagery. Data and methods are detailed in Thakuri et al.¹⁰. We interpreted the land 110 111 cover of the remaining part of each basin through a visual interpretation of the Advanced Land Observation Satellite 112 (ALOS, October 2008, pixel 10 m) remote imagery and distinguished terrain coverage by bare rocks, bare soils, and 113 shrub vegetation. No rock glaciers were detected in these basins by visual interpretation.

Because we were interested in understanding whether the changes that occurred over the last twenty years in glacier surface areas and snow cover have influenced the solute variations observed during the same period, we had to overcome, as shown below, such limits linked with the resolution and availability of the satellite data.

Regarding glaciers, the satellite resolution of remote imagery available for covering the analyzed period provides a large uncertainty in estimating changes in surface areas, especially for small glaciers, such as those located within the basins of the survey lakes; thus, the estimated changes are dubious¹⁹. This limit has been overcome by enlarging the scale of analysis to the overall SNP. At this scale, the 29 glaciers (with a surface area >1 km²) that were already traced by Thakuri et al.¹⁰ are more than 30 times larger (5.7 ± 0.40 km² as median) than those located within the survey lakes (0.31 ± 0.10 km² as median); thus, changes are more representative and less uncertain¹⁹.

123 Concerning the snow cover, the MODIS imagery is available only since the early 2010s. For the previous decade, 124 we deduced the trend in the coverage based on the estimated snowfall decrease reported in this area since the early 125 1990s⁹. The increase of temperature observed outside the monsoon period, when the precipitation is almost completely 126 composed by snow, brought a significant decrease (-14%, p < 0.05) of the snowfall probability for the last twenty years at 5050 m a.s.l.. The trend reported above has been calculated by these authors according to Fujita and Sakai²⁴ and 127 Ueno et al.²⁵ assuming that the probability of snowfall and rainfall depends on mean daily air temperature, using as 128 thresholds 0 and 4 °C, respectively. The variations that occurred in the 2002-2012 period were computed as the 129 130 difference between the first five and the last five years. Even in this case, the analysis was enlarged to the overall SNP 131 to increase the representativeness within an elevation ranging from 4500 to 5500 m a.s.l. (6152 MODIS pixels), i.e., the 132 altitudinal band of the survey lakes (Fig. 1b).

At this wide scale, we determined that changes in glacier surface areas and seasonal snow cover duration are related to certain morphometric boundary conditions. Elevation, aspect, and slope have been investigated for snow cover duration and glaciers, but for glaciers, we also considered the maximum, minimum and mean elevation, and the down and upstream slopes of the glaciers following the experience of Thakuri et al.¹⁰ and Salerno et al.¹⁹. Detailed computational methods of these morphometric variables are reported in the works of these authors. In the same way, at the scale of each survey lake basin, we determined that the differences that occurred in SO_4^{2-} and HCO_3^{-} are related to the same morphometric boundary conditions investigated for glaciers and snow cover duration at a wider scale.

For simulating the daily melting of the glacier located in the LCN9 basin we used a T-index model²⁶. This model uses the mean daily air temperature, the glacier elevation bands, and a melt factor (0.0087 m d⁻¹ °C⁻¹) provided by Kayastha et al.²⁷ from a field study (Glacier AX010) located close to the SNP.

143 Statistical analysis. We conducted a Principal Component Analysis (PCA) among the lake chemical components and 144 the basin morphometric features to obtain information on the relationships among the data and to look for reasons that 145 could justify the observed changes in major ion concentrations²⁸. The test was performed by using the "princomp" and 146 the "biplot" functions in the R Project environment²⁹. The degree of correlation among data was verified through the 147 correlation coefficient (r) after testing that the quantile-quantile plot of model residuals follows a normal distribution. 148 Otherwise data were log-transformed to meet the statistical requirements for normal distribution; then the residuals of 149 the regressions were tested for homoscedasticity (not shown here) (e.g.,²⁹). The chemical concentrations of the 20 150 survey lakes, measured in the early 1990s and the early 2010s, were compared, and the differences between the two

151 periods were tested by using the non-parametric ANOVA for paired comparisons (Friedman test) at $p < 0.05^{28,30}$.

152 Differences in glacier surface areas and snow cover duration were analyzed by applying the same statistical test. The

153 significance of annual chemical trends was evaluated with the annual Mann Kendall test at p < 0.05.³¹

154 RESULTS AND DISCUSSION

Survey lakes. In the early 1990s, most of the survey lakes exhibited low values of conductivity (25 μ S cm⁻¹ as 155 median) (Table 2, Fig. 2a). After approximately 20 years, the conductivity increased significantly, by 47% (p<0.001) on 156 average. In some cases, the values even doubled. Lakes became significantly (p<0.001) enriched in solutes, primarily in 157 SO_4^{2-} (+57%), then in $Ca^{2+}+Mg^{2+}$ (+41%) and less in HCO_3^{--} (+11%) (Table 2, Fig. 2b,c,d). The increase in SO_4^{2-} was 158 even 4-fold in LCN9 (from 84 to 327 µeq L-1) and LCN10 (from 73 to 285 µeq L-1) (Fig. 2b). For these lakes, a 159 160 significant increasing annual trend (p<0.05) was observed both for conductivity, SO_4^{2-} , $Ca^{2+}+Mg^{2+}$, and HCO_3^{-1} concentrations (Fig. 3b,c,d), with two main peaks in 1998 and 2010 (discussed in the last paragraph). All survey lakes 161 162 (Table 2 and Fig. 2e), as well as LCN9 and LCN10 (Fig. 3e), exhibited a significant increase of hydrogen ion (H⁺) 163 concentrations (126%, p<0.01).

164 The temporal change of the relative contribution of the various ions to the total ionic content of LCN 9 can be 165 observed in Figure SI1: sulfate passed from 22 to 36%, calcium remained stable (41%), while alkalinity contribution decreased from 24 to 11%. To assess the relative change in the ionic composition of the survey lakes, we calculated the 166 167 c-ratio $HCO_3^{-1}(HCO_3^{-1} + SO_4^{2-})$, which is a geochemical indicator that is used to evaluate the relative dominance of the 168 carbonation reaction and sulfide oxidation in meltwaters.³² The c-ratio decreased significantly in the survey lakes (-18%, p<0.01) (Table 2, Fig. 2f). Particularly pronounced is the c-ratio decline observed in LCN9, from 0.56 to 0.29 (-169 170 48%), and in LCN10, from 0.59 to 0.31 (-47%) (Fig. 3f). The only exception with respect to the general pattern is 171 LCN31 (Fig. 2f). The Duwo glacier, which is located within the LCN31 basin, is surging and shows a divergent 172 response to climate change with respect to all of the other glaciers located in the region.¹⁰ Therefore, we think that the 173 two anomalies could be associated. All data presented in this section are reported in Table SI1.

In the following, we discuss potential factors that may have contributed to the increase of solute concentrations in the lakes, with special attention towards SO_4^{2-} and HCO_3^{-} ions: atmospheric deposition, weathering, precipitation, glaciation, and snow cover duration.

Atmospheric Deposition. The chemistry of wet deposition was investigated at the Pyramid station during 1991-1992³³ and 2007-2008³⁴. The precipitation contents for all ions were the lowest ones among those measured at high elevation sites around the world³². The comparison between the studied periods revealed no substantial variations in SO_4^{2-} and other compounds (NO_3^{-} and NH_4^{+}) that resulted from anthropic activities, thus suggesting that the tropospheric

- background concentrations are constant in the Himalayas³⁴. These findings indicate that the atmospheric deposition
- 182 cannot play significant role in the solute increasing trend recorded in Himalayan lakes.

183 Land cover and Snow cover duration. We tested the possible dependence of SO_4^{2-} and HCO_3^{-} concentrations for the 184 survey lakes sampled in 1990s with respect to the basin land cover and seasonal snow cover duration. We preferred to 185 use 1990s data in order to investigate the dependences between these variables before a period (1990s-2010s) influenced decidedly more by higher climate-driven changes⁹. The correlation matrix among all of these variables is 186 presented in Fig. SI3, and an overall overview of their mutual relationships is provided with the PCA in Fig. 4a. The 187 main finding is that the SO_4^{2-} concentrations are directly related to glacier coverage (r=0.72, p<0.001) (Fig. 4b), i.e., 188 189 higher lake solute concentrations are found for basins with more glacial coverage. No relationship can be observed with 190 the other selected variables and in particular with the snow cover duration. Furthermore, we note that the HCO₃-191 concentrations do not have significant relationship with the land cover and the snow cover duration.

192 When observing the morphometric features of the LCN9 basin and the LCN10 sub-basin (Table 1), it becomes clear 193 that the discriminating feature between these lakes is the presence of a small glacier of 0.11 km² lying within the LCN9 194 basin, whereas the LCN10, which is located downstream, receives the glacial melting indirectly from LCN9 (Graphical 195 abstract). Concurrently, LCN10 has lower solute concentrations and conductivity (Fig. 3a, 3b, 3c, 3d). This fact can be 196 interpreted as further evidence that glaciers (subglacial and surrounding periglacial components) are the most important driver of changes in both SO42- concentrations and that the runoff in the non glaciated areas of the LCN10 sub-basin as 197 198 well as the possible groundwater discharge from headwater areas of the basin transports lower concentrations, thereby 199 diluting the LCN10 content. This interpretation is corroborated by the further evidence provided by lakes LCN68 and 200 LCN67. From Table SI1 we observe that the downstream lake (LCN67) has, among the other solutes, lower sulfate concentrations (733 µeq L⁻¹ respect to 926 µeq L⁻¹ for LCN68). Even in this case the discriminating feature is the 201 presence of a glacier lying within the upstream lake basin (see Fig. SI2 and Salerno et al.¹⁹- Fig. SI2d-). 202

203 Changes in glacier surface areas. Thakuri et al.¹⁰, considering glaciers larger than 1 km², reported a surface area loss of -8.7%, p<0.01 for the overall SNP during the 1992-2011 period (uncertainty $\pm 2.8\%$). Applying here the same remote 204 205 imagery and methods reported by these authors, but at glaciers located within the basins of the 20 survey lakes 206 (0.31±0.10 km² as median), we found a higher decrease in glacier area, but with an uncertainty connected with the 207 satellite resolution that increased by of an order of magnitude (-19.7%, p < 0.01, uncertainty $\pm 23.9\%$). The glacier within the LCN9 basin lost 14% (uncertainty $\pm 22\%$) of its surface area. According to many authors^{10,35,36,37}, even in SNP and 208 209 in the Himalayas in general, the main losses in area over the last decades have been observed for smaller glaciers 210 (glacier size vs glacier area change r=0.58, p<0.01, Fig. SI4). Therefore, the survey lake basins are subject to a glacier shrinkage twice that observed for the glacial masses in the SNP region. However, the reduced size of these glaciers causes the area loss estimation to be affected by a large uncertainty that is even higher than its magnitude, and the estimation is thus too rough to be directly compared with the chemical variations observed in the relevant downstream lakes. Therefore, as described above, we conducted two parallel analyses: at the SNP level and at the scale of a survey lake basin.

216 At a larger scale, considering the data from Thakuri et al.¹⁰, we analyzed which morphometric factors were able to 217 cause changes in surface areas in the last twenty years (Fig. 5a). The correlation matrix among all of these variables is 218 presented in Fig. SI4. From the PCA analysis shown in Fig. 5a, we can see that the downstream slope (r=-0.66, 219 p < 0.001) (Fig. 5c) and the relevant aspect (r=-0.75, p < 0.001) of glaciers are the main factors responsible for the surface 220 area losses, whereas elevation seems to have less influence. These findings agree with the few other studies investigating these relationships^{19,35,36,37}. Whereas elevation is a proxy for temperature^{35,37}, aspect is a proxy for 221 222 insolation³⁸, and slope is the key factor responsible for the gravitational driving stresses to which glaciers are subjected³. 223 Lower slopes are supposed to induce lower glacier ice flow, thus allowing the development of stagnant ice conditions and consequently lower terminus retreat rates^{3,39,40} and the development of supraglacial lakes^{19,41}. In contrast, as we 224 225 found here, under the climate warming pressure observed in the region for the last decades⁸, higher slopes favor the 226 shrinkage of glaciers because they are supposed to contribute to the higher glacier velocity⁴².

Moreover, we observed that even glacier aspect is indirectly correlated with surface area losses, i.e., the largest 227 228 shrinkages have been experienced mainly for E and W facing glaciers, and the shrinkages are decidedly smaller for S 229 facing ones (Fig. 5a). Considering that S oriented glaciers are supposed to receive more solar radiation³⁸, the observed 230 spatial shrinkage seems to be counterintuitive. However, we have to consider that, as their aspects deviates from S, the 231 glaciers become significantly smaller (glacier aspect vs glacier size r=-0.61, p< 0.001) and steeper (glacier aspect vs glacier downstream slope r=0.52, p< 0.01) (all correlations are shown in Fig. SI4) because they are located in valleys 232 233 perpendicular to the prevailing S-N direction of the monsoon. In our opinion, these valleys have been less hollowed, 234 and the current glaciers are thus small and steep. Therefore, the observed correlation between aspect and glacier surface 235 area losses indicates that gravitational stresses (conditioned by slopes) are higher for aspects far from S, and these, as 236 observed above, are key factors in defining the rate of retreat of glaciers under the climatic pressure.

Although affected by a wide uncertainty, even at the scale of 20 survey lake basins we analyzed which morphometric factors were able to cause changes in surface areas. The correlation matrix among all of these variables is presented in Fig. SI4. We can see that the glacier slope (r=-0.58, p<0.05) and the relevant aspect (r=-0.49, p<0.05) of glaciers are the main factors responsible for the surface area losses, though, for the effect of larger uncertainty, these correlations are weaker than what we found at large scale. 242 Changes in snow cover duration. During the 2002-2012 period, we observed a decrease of the seasonal snow cover 243 duration of -29% (p<0.01) within an elevation ranging from 4500 to 5500 m a.s.l. A similar behavior (even if we focus 244 on the analysis exclusively) was found within the 20 survey lake basins (-27%, p<0.01). We found that the largest 245 differences in snow cover duration occurred with W and S exposure and that the smallest occurred in the opposite 246 orientations. Aspect (r=0.23, p<0.001) and elevation (r=0.24, p<0.001) are the main factors responsible for changes in 247 snow cover duration and we did not find any strong relationship with slope (all correlations are shown in Fig. SI5). In 248 contrast to what we observed for glaciers, the snow cover duration at lower elevations decreased more due to higher 249 temperatures that were recordable downstream, and the snow cover duration on S and W slopes decreased more than on 250 N and E slopes, likely as a result of the higher solar irradiance received by these sides⁴³. In the next paragraph changes 251 observed in the seasonal snow cover duration are related to changes occurred in SO_4^{2-} and HCO_3^{-} concentrations of 20 252 survey lakes.

253 Changes in ionic concentrations. Once it is defined how variations recorded in glacier and snow cover duration have 254 been conditioned by morphometric factors, we analyzed which of these factors are mainly responsible for changes 255 observed in SO₄²⁻ and HCO₃⁻ concentrations in the survey lakes (Fig. 5b). Concerning the HCO₃⁻ concentrations, we did 256 not find any relationship with the selected morphometric boundary conditions (Fig. 5b; Fig. SI6). Differently, Figure 5b 257 reveals that glacier slope (r=0.81, p<0.001) (Fig. 5d) and relevant aspect (r=0.66, p<0.01) are well correlated with $SO_4^{2^-}$ 258 concentration changes, whereas the direct relationship with the glacier surface areas losses is significant, but weaker 259 (r=-0.49, p<0.05), probably affected by the high uncertainty connected with the measurements of small glaciers (all 260 correlations are shown in Fig. SI6). From Figure SI6 we can also observe that the SO_4^{2-} concentration changes are not 261 correlated with the basin slopes (r=0.18, p>0.10), as well as the direct relationship between glaciers slopes and basins slopes is weak (r=0.44, p<0.05) enforcing the causal link between the SO42- variations and the glacier slopes. As 262 263 indicated above, the glacier slope is also the main morphological feature able to explain the changes of the glacier 264 surface area at higher and more representative scale (Fig. 5a and Fig. 5b). Therefore, we can assert that lake basins with 265 steep glaciers oriented far from S have experienced higher glacier surface area losses and a concomitant SO₄²⁻ increases. 266 Differently, the main decreases in snow cover duration occurred at lower elevations and on land with a S orientation, 267 and they are less strictly dependent on the terrain slope (Fig. SI5 and Fig. SI6). Therefore, the changes observed in snow 268 cover duration seem not to be the main cause of the observed SO_4^{2-} increases.

LCN9 experienced a particularly evident increase in ionic concentrations, which was even 4-fold in the case of SO_4^{2-} concentration (Fig. 3a). In this case, the reason for this huge change can also be found in that the LCN9 glacier has the most divergent aspect (SE-E, 110°) from the south orientation and has the highest glacier slope (36°) among the 20 survey lakes (Table1). Moreover, a synchronous temporal pattern of solutes and conductivity can be observed in LCN9 and LCN10 (Fig. 3a, 3b, 3c): the increases for both lakes should be driven by the same source, i.e., the glacier locatedwithin the LCN9 basin.

To further investigate the processes, which could have led to the SO_4^{2-} increase, we analyzed the associations 275 276 between the main ions and how they changed during the last twenty years. We observed a highly significant positive relationship between SO_4^{2-} and $Ca^{2+} + Mg^{2+}$ in both in the early 1990s and in the early 2010s (r = 0.85 and 0.94, 277 278 p<0.001) (Fig. SI3). On the contrary, the relationship between HCO₃⁻ and Ca²⁺ + Mg²⁺ is weakly significant in the early 279 1990s (r=0.39, p<0.05) (Fig. SI3) and not significant in the early 2010s. The association between SO_4^{2-} and HCO_3^{-} was quite low in both periods (r = 0.45, p=0.04 and "not significant" in 2010s). The ratios of $(Ca^{2+} + Mg^{2+})/(Na^{+} + K^{+})$ was 280 281 7.2 in the early 1990s and 8.7 in the early 2010s, indicating the relative weakness of silicate dissolution. This is 282 confirmed also by the lack of correlation between Si and $Na^+ + K^+$ in the 2010s (r=0.17). These results make evident an 283 excess of SO_4^{2-} with respect to HCO_3^{-} in waters flowing in the drainage system of glaciers. This shift towards SO_4^{2-} 284 enriched waters is even more evident in LCN9 and LCN10, as shown by the comparison of the ionic composition of 285 lakes water in the 1990s and in the 2010s (Table SI1).

It is known that subglacial environments are dominated by carbonate dissolution and sulfide oxidation⁴⁴, often 286 microbially mediated^{45,46,47}. Through these processes, carbonate and sulfate concentrations in melt waters are greater 287 than might be expected from the bedrock lithology.^{48,49,50,51} A number of studies demonstrated that carbonation was 288 289 mainly responsible of solute acquisition in the quick-flow component of the glacier hydrological system, while the 290 enrichment in SO42- seems to be strictly associated to delayed flow waters.52 For these reasons, sulfate is considered an indicator of distributed type drainage beneath glaciers⁵³ and its increase, in most glacial environments, is an effect of 291 292 increasing residence times and rock/water interaction⁵⁴. In general, the oxidation of sulfide minerals and the dissolution 293 of carbonates are partially coupled, since the hydrogen ions, originated from the sulfide oxidations, fuel the carbonate 294 hydrolysis⁵³. Our results, showing the lack of any relationship between SO_4^{2-} and HCO_3^{-} , suggest that the two processes 295 are uncoupled in our system i.e. protons generated by sulfide oxidation do not dissolve carbonates. Furthermore, for 296 most lakes the c-ratio was at values lower than 0.5 in 2010s and, particularly for LCN9 and LCN10, reached 0.2 in 297 some years. These results suggest that other processes, likely in addition to carbonate dissolution and sulfide oxidation, 298 occurred in the selected lake basins.

A process consistent with the elevated SO_4^{2-} , Ca^{2+} and Mg^{2+} concentrations found recently in lake waters is the pyrite oxidation coupled to the preferential weathering of calcium-silicate minerals which presence is documented in the study area. As suggested by Williams et al.⁷ the Si produced from the weathering of calcium-silicate would be precipitated due to the effect of low temperature and this could explain the lack of correlation between SO_4^{2-} and Si, and the slight Si increase observed in the studied period. 304 The deglaciation observed in the study area has, first of all, increased the amount of meltwater causing a rising of conductivity of lake waters. The disproportional increase of SO_4^{2-} compared to HCO_3^{-} may be explained by i) a change 305 306 in flow-paths and residence time of the meltwaters, leading to more favorable conditions for the oxidation of sulfides, 307 and ii) the pyrite oxidation followed by the weathering of calcium-silicate. We cannot exclude a role of microbial populations at glacier beds if the conditions had become anoxic.45,46,47 Figure SI7 shows the significant correlation 308 309 between H⁺ differences and SO₄²⁻ differences (r=0.72, p<0.001) observed for the studied lakes, which can be interpreted 310 as further evidence of the importance of sulfide oxidation connected with the glacier retreat. At this regards it is interesting to observe that Andersen⁵⁵ wrote that sulfide oxidation contributes to solute fluxes from glaciers to an extent 311 312 greater than might be expected from mineral abundance.

313 Weathering and precipitation.

314 Regarding the weathering occurring on the non glaciated part of the basin (periglacial environment), we analyzed the behavior of the lakes (three) without glaciers within their basins. They showed a slight increase in SO_4^{2-} 315 316 concentration (+17% as median) (Fig. 2b), whereas higher increases were observed for lakes with glaciers within their 317 basins (+78% as median). These differences between the two groups were significant (p<0.01). The slight increase in SO42- concentration of lakes without glaciers could be attributed to the intensification of weathering rates caused by the 318 319 reduction of the snow cover duration⁵⁶, although we already discussed that this process is not the main driver of change. 320 Particularly, during the warmer season, on the sunny slopes, previously snow covered, an increase in soil temperature from 0°C to 25°C could be expected, which could intensify the weathering rates by an order of magnitude.⁵⁷ 321 322 Weathering could be further enhanced by wetter reactive surface areas of minerals⁵⁸, but the decreasing precipitation 323 trend, as observed above over the last twenty years, does not support this possibility.

Nonetheless, the decreasing precipitation trend could have influenced the solute concentration in a different way: lower but more concentrated runoff, could have slightly enriched the lakes⁵⁹. These solute variations were caused by enhanced weathering rates, or more probably by more concentrated runoff. Another mechanism considers that the possible dropping of water-table depth, due to decreased groundwater recharge, resulting in more rock exposed to air and enhanced mineral oxidation.⁶⁰ However they can be considered background increases that are common to all survey lakes with respect to the higher increases observed in conjunction with glacier shrinkages.

Annual chemical trends for two selected lakes. Sulfate was the ion showing the most pronounced change in LCN9 and LCN10 (Fig. SI1, Table SI1). Hence we tested the possible dependence of the SO_4^{2-} concentrations in LCN9 on monthly mean temperature, precipitation, and glacier melting, estimated by applying the T-index model²⁶. We found that SO_4^{2-} concentrations are only significantly correlated with the mean temperature in April (r=0.74, p<0.001). The temperature and precipitation of the other months did not show any significant influence on the observed SO_4^{2-} concentrations (Fig. SI8). As for estimated glacier melt, we found that the only month that was significantly correlated, even in this case, is April (r=0.74, p<0.001) (Fig. 6a, 6b), whereas March shows a weak correlation (r=0.45, p<0.1) (Fig. 6a).

338 As shown above, significant increases in mean air temperature have been experienced over the last twenty years for 339 the months of April, November and December.9 However, April is the only month in which the increase has been 340 effective, i.e., it has de facto favored the melting process. More precisely, the 0 °C isotherm in April shifted from 341 around the mean elevation of glacier fronts to the mean elevation of glaciers located in the survey lake basins (Fig. 1b). 342 In contrast, during the winter months, the temperature is so low that the 0 °C isotherm upward cannot have reached 343 these elevations, and it cannot have had an effective impact on the melting processes. As a result, the T-index model 344 estimates a significant increasing trend only for April (p<0.01). At this regards, the two main peaks of conductivity and 345 major ion concentrations in 1998 and 2010 observed in Figure 3 are caused by the high temperature registered for the 346 month of April of those years with subsequent high melting processes as estimated by the T-index model.

Previously, Lami et al.⁵ observed a significant relationship between the enhanced conductivity detected in LCN9 and LCN10 and the annual temperature until 2008. Here we confirm the observations of these authors until 2013 considering the temperature as the main climatic driver of change. Besides that, we found the temperature of April is the effective drive of the observed enhanced glacier melting process, which is considered here the main factor responsible for the observed increase of sulfate concentrations.

The chemical variations observed in lake water in the last two decades represent a response of these fragile ecosystems to climate change. In particular, glacier retreat proved to be the main driver of the solute increase affecting the lakes. Major ion concentrations, and sulfate at a higher extent, significantly rose in lake water. Even if these changes do not pose a direct and immediate threat to the biota, they occurred in a limited time span, and significantly modified the average chemical composition of lake water. For these reasons, the lakes and the main factors driving their variability should be regularly monitored in the future, also in relation to the lake role as ecosystem services.

358 SUPPORTING INFORMATION

A table showing the chemical and topographic features of 20 survey lakes. A figure showing the relative contribution of various ions to the total ionic content of LCN 9 in the early 1990s and in the early 2010s. ALOS 2008 imagery of LCN68 and LCN67 basins. Correlation matrixes among chemical species, land covers, and topographic boundary conditions. A figure showing the relationship between the SO4²⁻ and the H⁺ concentration changes observed in all

- survey lakes between 1990s and 2010s. A figure showing the PCAs among SO4²⁻ and HCO₃⁻ concentrations of LCN9,
- 364 mean temperature, and precipitation for each month of the year.

365 ACKNOWLEDGEMENTS

- 366 This work was supported by the MIUR through Ev-K2-CNR/SHARE and CNR-DTA/NEXTDATA project within the
- 367 framework of the Ev-K2-CNR and Nepal Academy of Science and Technology (NAST). Special thanks are due to the
- **368** Pyramid Laboratory's staff for its assistance in sampling and field work.

369 REFERENCES

- 1. Williamson, C. E.; Saros, J. E.; Schindler, D. W. Sentinels of change. *Science* 2009, 323, 887-887.
- 2. Adrian, R.; O'Reilly, C. M.; Zagarese, H.; Baines, S. B.; Hessen, D. O.; Keller, W.; Livingstone, D. M.;
- 372 Sommaruga, R.; Straile, D.; van Donk, E.; Weyhenmeyer, G. A.; Winderl, M. Lakes as sentinels of climate change.
- **373** *Limnol. Oceanogr.* **2009**, 54, 2283–2297.
- Scherler, D.; Bookhagen, B.; Strecker, M. R. Spatially variable response of Himalayan glaciers to climate change
 affected by debris coverage. *Nat. Geosci.* 2011, 4, 156–159.
- 4. Gardelle, J.; Arnaud, Y.; Berthier, E. Contrasted evolution of glacial lakes along the Hindu Kush Himalaya
 mountain range between 1990 and 2009. *Global Planet. Change* 2011, 75, 47–55
- 378 5. Lami, A.; Marchetto, A.; Musazzi, S.; Salerno, F.; Tartari, G.; Guilizzoni, P.; Rogora, M.; Tartari, G. A. Chemical
 379 and biological response of two small lakes in the Khumbu Valley, Himalayas (Nepal) to short-term variability and
- climatic change as detected by long-term monitoring and paleolimnological methods. *Hydrobiologia* 2010, 648,
 189–205.
- 382 6. Thies, H.; Nickus, U.; Mair, V.; Tessadri, R.; Tait, D.; Thaler, B.; Psenner, R. Unexpected response of high Alpine
 383 Lake waters to climate warming. *Environ. Sci. Technol.* 2007, 41, 7424–7429.
- 384 7. Williams, M. W; Knauf, M.; Caine, N.; Liu, F.; Verplanck, P. Geochemistry and source waters of rock glacier
 385 outflow, Colorado Front Range. *Permafrost Periglac.* 2006, 17(1), 13–33.
- 8. Mast, M. A.; Turk, J. T.; Clow, D. W.; Campbell D. H. Response of lake chemistry to changes in atmospheric
 deposition and climate in three high-elevation wilderness areas of Colorado. *Biogeochemistry* 2010, 98, 1-17.
- 388 9. Salerno, F.; Guyennon, N.; Thakuri, S.; Viviano, G.; Romano, E.; Vuillermoz, E.; Cristofanelli, P.; Stocchi, P.;
- 389 Agrillo, G.; Ma, Y.; Tartari, G. Weak precipitation, warm winters and springs impact glaciers of south slopes of Mt.
- Everest (central Himalaya) in the last 2 decades (1994–2013). *Cryosphere* **2015**, 9, 1229-1247.

- 10. Thakuri S.; Salerno, F.; Smiraglia, C.; Bolch, T.; D'Agata, C.; Viviano, G.; Tartari, G. Tracing glacier changes since
 the 1960s on the south slope of Mt. Everest (central Southern Himalaya) using optical satellite imagery. *Cryosphere*
- **393 2014**, 8(4), 1297-1315.
- 11. Tartari, G.; Salerno F.; Buraschi E.; Bruccoleri G.; Smiraglia C. Lake surface area variations in the North-Eastern
 sector of Sagarmatha National Park (Nepal) at the end of the 20th Century by comparison of historical maps. J. *Limnol.* 2008, 67, 139–154.
- 397 12. Gruber, S.; Fleiner, R.; Guegan, E.; Panday, P.; Schmid, M.-O.; Stumm, D.; Wester, P.; Zhang, Y.; Zhao, L. Review
 398 article: Inferring permafrost and permafrost thaw in the mountains of the Hindu Kush Himalaya region. *The* 399 *Cryosphere Discuss*. 2016, in review, doi:10.5194/tc-2016-104.
- 400 13. Caine, N. Recent hydrologic change in a Colorado alpine basin: an indicator of permafrost thaw? *Ann. Glaciol.*401 2010, 51(56), 130-134.
- 402 14. Leopold. M.; Lewis, G.; Dethier, D.; Caine N.; Williams M.W. Cryosphere: ice on Niwot Ridge and in the Green
 403 Lakes Valley, Colorado Front Range. *Plant Ecol. Divers.*2015, 8 (5-6), 625-638.
- 404 15. Amatya, L. K.; Cuccillato, E.; Haack, B.; Shadie, P.; Sattar, N.; Bajracharya, B.; Shrestha, B.; Caroli, P.; Panzeri,
 405 D.; Basani, M.; Schommer, B.; Flury, B.; Manfredi, E. C.; Salerno, F. Improving Communication for Management
 406 of Social-ecological Systems in High Mountain Areas. *Mt. Res. Dev.* 2010, 30, 69–79.
- 407 16. Salerno, F.; Viviano, G.; Manfredi, E. C.; Caroli, P.; Thakuri, S.; Tartari, G. Multiple Carrying Capacities from a
 408 management oriented perspective to operationalize sustainable tourism in protected area. *J. Environ. Manag.* 2013,
 409 128, 116–125.
- 410 17. Searle, M. P.; Simpson, R. L.; Law, R. D.; Parrish, R. R.; Waters, D. J. The structural geometry, metamorphic and
 411 magmatic evolution of the Everest massif, High Himalaya of Nepal– South Tibet. *J. Geol. Soc.* 2003, 160, 345–66.
- 412 18. Nepal, S. K.; Nepal S. A. Visitor Impacts on Trails in the Sagarmatha (Mt. Everest) National Park, Nepal. AMBIO,
 413 2004, 33(6), 334-340.
- 414 19. Salerno, F.; Thakuri, S.; D'Agata, C.; Smiraglia, C.; Manfredi, E. C.; Viviano, G.; Tartari, G. Glacial lake
 415 distribution in the Mount Everest region: Uncertainty of measurement and conditions of formation. *Global Planet*.
- 416 *Change* **2012**, 92–93, 30–39.
- 417 20. Tartari, G. A.; Tartari G.; Mosello, R. *Water chemistry of high altitude lakes in the Khumbu and Imja Kola valleys*
- 418 (*Nepalese Himalayas*); Lami, A.; Giussani G., Ed, Memorie dell'Istituto italiano di Idrobiologia: Verbania Pallanza,
 419 1998.
- 420 21. Ichiyanagi, K.; Yamanaka, M. D.; Muraji, Y.; Vaidya, B. K. Precipitation in Nepal between 1987 and 1996, *Int. J.*
- 421 *Climatol.* **2007**, 27, 1753–1762.

- 422 22. Pal, I.; Al-Tabbaa, A. Long-term changes and variability of monthly extreme temperatures in India. *Theor. Appl.*423 *Climatol.* 2010, 100, 45–56.
- 424 23. Yao, T.; Thompson L.; Yang W.; Yu W.; Gao Y.; Guo X.; Yang X.; Duan K.; Zhao H.; Xu B.; Pu J.; Lu A.;
- Xiang Y.; Kattel D. B.; Joswiak D. Different glacier status with atmospheric circulations in Tibetan Plateau and
 surroundings. *Nat. Clim. Change* 2012, 2(9), 663-667.
- 427 24. Fujita, K.; Sakai, A. Modelling runoff from a Himalayan debris-covered glacier, *Hydrol. Earth Syst. Sci.* 2014, 18,
 428 2679–2694.
- 429 25. Ueno, K., Endoh, N., Ohata, T., Yabuki, H., Koike, M., Zhang, Y. Characteristics of precipitation distribution in
 430 Tangula, Monsoon, 1993, B. Glaciol. Res., 12, 39–46, 1994.
- 431 26. Hock, R. Temperature index melt modeling in mountain areas. *J Hydrol.* 2003, 282, 104-115.
- 432 27. Kayastha, R. B.; Ageta, Y.; Nakawo, M. Positive degree-day factors for ablation on glaciers in the Nepalese
 433 Himalayas: case study on Glacier AX010 in Shorong Himal, Nepal. *Bull. Glaciol. Res.* 2000, 17, 1-10.
- 434 28. Salerno, F.; Gambelli, S.; Viviano, G.; Thakuri, S.; Guyennon, N.; D'Agata, C.; Diolaiuti, G.; Smiraglia, C.;
- 435 Stefani, F. ; Bocchiola, D. ; Tartari, G. High alpine ponds shift upwards as average temperatures increase: a case
- 436 study of the Ortles-Cevedale mountain group (Southern Alps, Italy) over the last 50 years. *Glob. Planet. Change*
- **437 2014**, 120, 81-91.
- 438 29. Venables, W. N.; Ripley, B. D. *Modern Applied Statistics with S*; Springer: New York, 2002.
- 439 30. Pohlert, T. The Pairwise Multiple Comparison of Mean Ranks Package (PMCMR). R package. 2014.
- 440 31. http://cran.r-project.org/web/packages/Kendall/Kendall.pdf. *Package Kendall*. R package. 2015.
- 32. Brown, G. H.; Tranter, M.; Sharp, M. Subglacial chemical erosion-seasonal variations in solute provenance, Haut
 Glacier d'Arolla, Switzerland. *Ann. Glaciol.* 1996, 22, 25–31.
- 33. Valsecchi, S.; Smiraglia, C.; Tartari, G.; Polesello, S. Chemical composition of monsoon deposition in the Everest
 region. *Sci. Total Environ.* 1999, 226, 187–99.
- 34. Balestrini, R.; Polesello, S.; Sacchi, E. Chemistry and isotopic composition of precipitation and surface waters in
 Khumbu valley (Nepal Himalaya): N dynamics of high elevation basins. *Sci. Total Environ.* 2014, 485-486, 681-
- **447** 692.
- 35. Salerno, F.; Buraschi, E.; Bruccoleri, G.; Tartari, G.; Smiraglia C. Glacier surface-area changes in Sagarmatha
 National Park, Nepal, in the second half of the 20th century, by comparison of historical maps. *J. Glaciol.* 2008, 54,
 738–752.
- 451 36. Loibl, D. M. ; Lehmkuhl, F. ; Grießinger, J. Reconstructing glacier retreat since the Little Ice Age in SE Tibet by
- 452 glacier mapping and equilibrium line altitude calculation. *Geomorphology* **2014**, 214, 22–39.

- 37. Racoviteanu, A. E. ; Arnaud, Y. ; Williams, M. W. ; Manley, W. F. Spatial patterns in glacier characteristics and
 area changes from 1962 to 2006 in the Kanchenjunga–Sikkim area, eastern Himalaya. *Cryosphere* 2015, 9, 505-523.
- 38. Oliphant, A. J.; Spronken-Smith, R. A.; Sturman A. P.; Owens, I. F. Spatial variability of surface radiation fluxes in
 mountainous region. *J. Appl. Meteorol.* 2003, 42, 113–128.
- 39. Bolch, T.; Buchroithner, M.; Pieczonka, T.; Kunert, A. Planimetric and volumetric glacier changes in the Khumbu
 Himal, Nepal, since 1962 using Corona, Landsat TM and ASTER data. *J. Glaciol.* 2008 54, 592–600.
- 459 40. Quincey, D. J.; Luckman, A.; Benn, D. Quantification of Everest region glacier velocities between 1992 and 2002,
 460 using satellite radar interferometry and feature tracking. *J. Glaciol.*2009, 55, 596–606.
- 461 41. Thakuri, S.; Salerno, F.; Bolch, T.; Guyennon, N.; Tartari, G. Factors controlling the accelerated expansion of Imja
 462 Lake, Mount Everest region, Nepal. *Ann. Glac.* 2016, 57(71), 245-257.
- 463 42. Cuffey, K. M.; Paterson, W. S. B. The Physics of Glaciers, 4th edition. Elsevier, Inc.. 2010.
- 464 43. Holst, T.; Rost, J.; Mayer, H. Net radiation balance for two forested slopes on opposite sides of a valley. *Int J.*465 *Biometeorol.* 2005, 49, 275–284.
- 466 44. Bottrell, S. H.; Tranter, M. Sulphide oxidation under partially anoxic conditions at the bed of the Haut Glacier
 467 d'Arolla, Switzerlan. *Hydrol. Process.* 2002, 16, 2363–2368.
- 468 45. Skidmore, M.; Anderson, S. P.; Sharp, M.; Foght, J.; Lanoil, B. D. Comparison of microbial community
 469 compositions of two subglacial environments reveals a possible role for microbes in chemical weathering processes.
- 470 *Appl. Environ. Microbiol.* **2005**, 71,6986–6997.
- 46. Hindshaw, R. S.; Heaton, T. H. E.; Boyd, E. S.; Lindsay, M. R.; Tipper, E. T. Influence of glaciation on mechanisms
 of mineral weathering in two high Arctic catchments. *Chem. Geol.* 2016. 420,037-50.
- 473 47. Wadham, J. L.; Tranter, M.; Skidmore, M.; Hodson, A. D.; Priscu, J.; Lyons, W. B.; Sharp, M.; Wynn, P.; Jackson,
- 474 M. Biogeochemical weathering under ice: Size matter. *Global Biogeochem. Cycles*, **2010** 24, GB3025, 1-11.
- 475 48. Sharp, M.; Tranter, M.; Brown, G. H.; Skidmore, M. Rates of chemical denudation and CO2 drawdown in a glacier476 covered alpine catchment. *Geology*, 1995, 23, 61–64.
- 477 49. Tranter, M., Sharp, M. J.; Lamb, H. R.; Brown, G. H.; Hubbard, B. P.; Willis, I. C. Geochemical weathering at the
- 478 bed of Haut Glacier d'Arolla, Switzerland—a new model. *Hydrol. Process.* **2002**, 16, 959–993.
- 479 50. Anderson, S.P. Biogeochemistry of glacial landscape system. Annu. Rev. Earth Planet Sci., 2007 35, 375-399.
- 480 51. Brown, G. H. Glacier meltwater hydrochemistry. *Appl. Geochem.* 2002, 17, 855–883.
- 481 52. Tranter, M.; Brown, G. H.; Raiswell, R.; Sharp, M.J.; Gurnell, A. M. A conceptual model of solute acquisition by
- 482 Alpine glacial meltwaters. J. Glaciol. 1993, 39, 573–581.

- 483 53. Wadham, J. L.; Tranter, M.; Hodson, A. J.; Hodgkins, R.; Bottrell, S.; Cooper, R., Raiswell, R. Hydro484 biogeochemical coupling beneath a large polythermal Arctic glacier: Implications for subice sheet biogeochemistry.
 485 *J. Geoph. Res.* 2010, 115, F04017.
- 54. Cooper, R. J.; Wadham, J. L.; Tranter, M.; Hodgkins, R.; Peters N. Groundwater hydrochemistry in the active layer
 of the proglacial zone, Finsterwalderbreen, Svalbard. *J. Hydrol.* 2002, 269, 208–223.
- 488 55. Anderson, S. P. Biogeochemistry of Glacial Landscape Systems. *Annu. Rev. Earth Planet. Sci.* 2007, 35, 375489 399.
- 490 56. Rogora, M.; Mosello, R., Arisci S. The effect of climate warming on the hydrochemistry of alpine lakes. *Water Air*491 *Soil Poll.* 2003. 148: 347-361.
- 492 57. Sommaruga-Wögrath, S.; Koinig, K. A.; Schmidt, R.; Sommaruga, R.; Tessadri, R.; Psenner, R. Temperature effects
 493 on the acidity of remote alpine lakes. *Nature* 1997, 387, 64-67.
- 494 58. Egli, M.; Lessovaia, S.; Chistyakov, K.; Inozemzev, S.; Polekhovsky, Y.; Ganyushkin, D. Microclimate affects soil
- 495 chemical and mineralogical properties of cold-alpine soils of the Altai mountains (Russia). J. Soils Sediments 2014,
- **496** 15, 1420-1436.
- 497 59. Grinsted, A.; Moore, J. C.; Pohjola, V.; Martma, T.; Isaksson, E. Svalbard summer melting, continentality, and sea
 498 ice extent from the Lomonosovfonna ice core. *J Geophys. Res.* 2006, 111, D07110.
- 499 60. Manning, A. H.; Verplanck, P. L.; Caine, J. S.; Todd, A. S. Links between climate change, water-table depth, and
- 500 water chemistry in a mineralized mountain watershed. *Appl. Geochem.* **2013**, 37, 64-78.

501 TABLES

502 TABLE 1. Mean morphometric features and land cover for the 20 selected lakes (reference year 1992). For LCN 10503 data are referred to the sub-basin draining directly into the lake (see graphical abstract).

504

	20 11		
Morphometry	20 survey lakes Median (range)	LCN9	LCN10
Lake elevation (m a.s.l.)	5010(4466-5427)	5209	5058
Lake area (km ²)	0.02(0.003-0.581)	0.005	0.013
Basin area (km^2)	0.8(0.1-41)	0.66	0.32
Basin slope (°)	29(9-35)	35	25
Basin aspect (°)	160(120-231)	120	120
Basin mean elevation (m a.s.l.)	5204(4843-5539)	5403	5102
Glacier area (km ²)	0.31(0.03-6.4)	0.11	-
Glacier slope (°)	26(15-36)	36	-
Glacier aspect (°)	151(110-221)	110	-
Glacier mean elevation (m a.s.l.)	5326(5087-5581)	5463	-
Land cover	20 survey lakes Mean (range)	LCN9	LCN10
Glacier (%)	18(0-50)	17	0
Bare rock (%)	28(0-66)	30	25
Bare soil (%)	40(9-74)	53	75
Shrub vegetation (%)	14(0-63)	0	0

505

506 TABLE 2. Median values of conductivity, selected ionic concentrations, and c-ratio (see the text for the explanation) of

507 the survey lakes sampled during the early 1990s and during the early 2010s. In brackets the standard deviation.

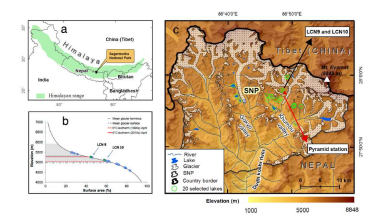
508

Feature	Early 1990s	Early 2010s	Difference (%)
Conductivity (µS cm ⁻¹ , 20 °C)	24.7 (16)	36.4 (27)	+47 (p<0.001)
SO42- (µeq L ⁻¹)	80 (129)	126 (245)	+57 (p<0.001)
HCO3- (μ eq L ⁻¹)	128 (54)	141 (61)	+11 (p<0.001)
Ca2++Mg2+ (µeq L ⁻¹)	221 (124)	331 (224)	+41 (p<0.001)
$H+(\mu eq L^{-1})$	0.03 (0.02)	0.07 (0.02)	+ 126% (p<0.01)
c-ratio	0.66 (0.22)	0.55 (0.26)	-18 (p<0.01)

509

510 FIGURE

511 FIGURE 1. a) Location of the study area in the Himalayas. b) Hypsometric curve of SNP and altitudinal glacier 512 distribution. Along this curve, the location of 20 selected lakes is shown. The 0 °C isotherms corresponding to the mean 513 monthly temperature in April are plotted for the years 1994 and 2013 according to the observed T trends and lapse 514 rates⁹. c) Focused map on the spatial distribution of lakes and glaciers in Sagarmatha National Park (SNP) and the 515 location of the Pyramid meteorological station.





517

518

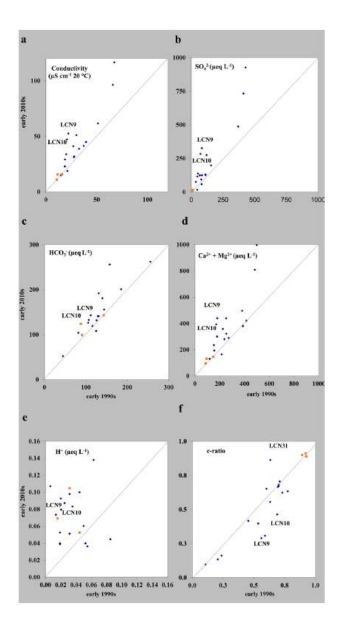
519

520

521

522

524 FIGURE 2. Scatter-plots between a) conductivity, b) SO_4^{2-} , c) HCO_3^{-} , d) $Ca^{2+}+Mg^{2+}$, e) H⁺, and f) c-ratio of the survey 525 lakes sampled during the early 1990s and the early 2010s. The orange points represent lakes without glaciers within the 526 basin.



533 FIGURE 3. Long-term trends of the annual values of a) conductivity, b) SO_4^{2-} , c) HCO_3^{-} d) $Ca^{2+}+Mg^{2+}$, e) H^+ 534 concentrations, and f) c-ratio in LCN9 (red) and LCN10 (blue).

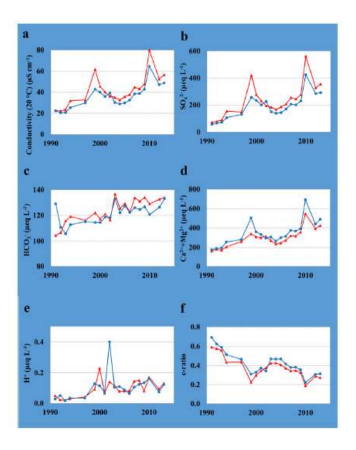


FIGURE 4. a) PCA among SO4²⁻, *HCO3⁻* concentrations of the survey lakes (early 1990s), land cover, and snow cover
duration (*Glacier*: glacier coverage during the early1990s; *Baresoil, Barerock, Shrubs*: coverage for 2008; *Snow*: mean
snow cover duration for each basin during the 2010s). b) The most significant correlation that emerged from the above
PCA.

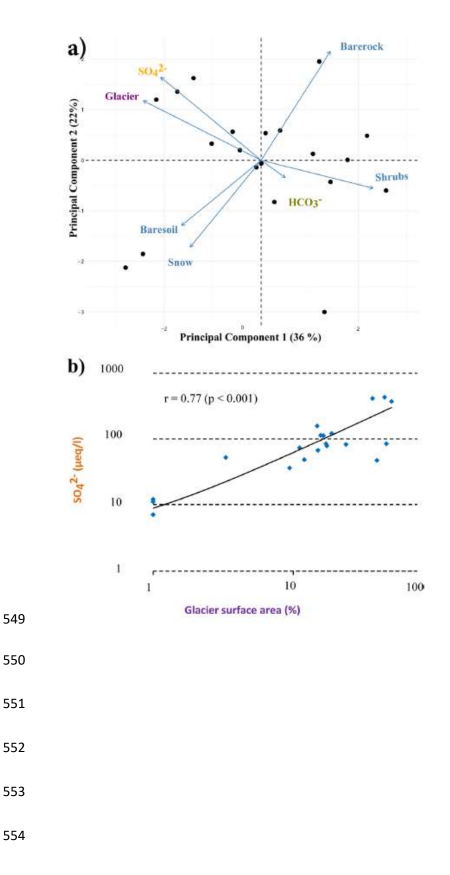
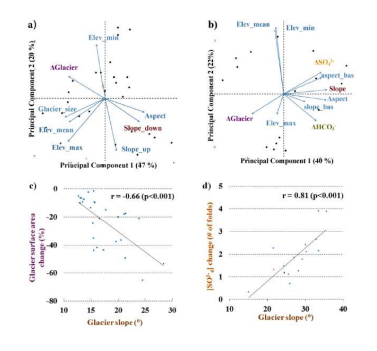


FIGURE 5. PCAs among morphometric parameters and a) glacier surface area variations ($\Delta Glacier$) that occurred in the last twenty years in the overall SNP (data from Thakuri et al.¹⁰); b) relative SO₄²⁻ and HCO₃⁻ variations (Δ SO₄²⁻, ΔHCO_3^-) that occurred in the same period for the 20 survey lakes; (*Glacier_size=size* of glaciers; *Elev_max, Elev_mean, Elev_min*: maximum, mean, and minimum glacier elevation, respectively; *Slope, Slope_down, Slope_up*: mean, downstream, and upstream glacier slope, respectively; *Aspect*: mean glacier aspect). The lower graphs (c and d) show the most significant correlations that emerged from the relevant PCAs shown above.



- 562 563
- 564
- 565
- 566
- 567
- 568
- _ _ _
- 569
- 570

571 FIGURE 6. a) PCA among SO4²⁻ and HCO₃⁻ concentrations of LCN9 sampled annually in late September or October 572 ($LCN9_SO4^{2-}$, $LCN9_HCO_3^{-}$, respectively) and the cumulated glacier melt (*melt_i*) modeled for each month (*i*) of the 573 year. b) The lower graph shows the most significant correlation that emerged from the PCA: $SO4^{2-}_LCN9$ vs *melt4*.

574

