



1 Leaf habit and nutrient availability drive leaf nutrient resorption globally

² Gabriela Sophia^{1,2,3}, Silvia Caldararu⁴, Benjamin D. Stocker^{3,5}, Sönke Zaehle^{1,6}

3 [1] Max Planck for Biogeochemistry, Jena, Germany; [2] International Max Planck Research
4 School on Global Biogeochemical Cycles; [3] Geographisches Institut, Universität Bern,
5 Switzerland; [4] Discipline of Botany, School of Natural Sciences, Trinity College Dublin,
6 Dublin, Ireland; [5] Oeschger Center, Universität Bern, Switzerland; [6] Friedrich Schiller
7 Universität Jena, Jena, Germany; (gsophia@bgc-jena.mpg.de)

8

9 Abstract

10 Nutrient resorption from senescing leaves can significantly affect ecosystem nutrient cycling, 11 making it an essential process to better understand long-term plant productivity under 12 environmental change that affects the balance between nutrient availability and demand. 13 Although it is known that nutrient resorption rates vary strongly between different species 14 and across environmental gradients, the underlying driving factors are insufficiently 15 quantified. Here, we present an analysis of globally distributed observations of leaf nutrient 16 resorption to investigate the factors driving resorption efficiencies for nitrogen (NRE) and 17 phosphorus (PRE). Our results show that leaf structure and habit, together with indicators of 18 nutrient availability, are the two most important factors driving spatial variation in NRE. 19 Overall, we found higher NRE in deciduous plants (65.2% \pm 12.4%, n=400) than in 20 evergreen plants ($57.9\% \pm 11.4\%$, n=551), likely associated with a higher share of metabolic 21 N in leaves of deciduous plants. Tropical regions show the lowest resorption for N (NRE: 22 52.4% \pm 12.1%) and tundra ecosystems in polar regions show the highest (NRE: 69.6% \pm 23 12.8%), while the minimum PRE is in temperate regions (57.8% \pm 13.6%) increasing to 24 boreal regions ($67.3\% \pm 13.6\%$). Soil clay content, N and P atmospheric deposition - a 25 globally available proxy for soil fertility - and MAP played an important role in this pattern, 26 where we found higher NRE and PRE in high latitudes. The statistical relationships 27 developed in this analysis indicate an important role of leaf habit and type for nutrient cycling 28 and guide improved representations of plant-internal nutrient re-cycling and nutrient 29 conservation strategies in vegetation models.

30 Keywords: Leaf nutrient content; Leaf structure; Nitrogen and phosphorus resorption
 31 efficiency; Plant ecophysiology; Plant functional traits; Plant nutrient limitation.





32 1. Introduction

³³ Nutrient cycling plays an important role in shaping the global distribution of terrestrial ³⁴ primary productivity (Le Bauer et al., 2008; Zaehle, 2013; Du et al., 2020). Nitrogen (N) and ³⁵ phosphorus (P) are the main limiting nutrients for plant growth. N is needed to maintain and ³⁶ produce essential proteins for the biosynthesis; while P is an element of genetic material and ³⁷ plays a major role in the regeneration of the main receptor of carbon (C) assimilation, and in ³⁸ the production of energy that conducts many processes in living cells (Chapin, 1980; ³⁹ Güsewell, 2004). The anthropogenic increase in atmospheric CO₂ since the beginning of ⁴⁰ industrialization has the potential to enhance the terrestrial carbon sink through increasing ⁴¹ plant photosynthetic rates, a process known as CO₂ fertilization (Bazzaz, 1990). A potential ⁴² limitation to the fertilization effect is progressive nutrient limitation to growth (Luo et al., ⁴³ 2004) and associated plant strategies to deal with such limitations. Thus, understanding the ⁴⁴ ways in which nutrients circulate in ecosystems and are acquired, lost, and conserved by ⁴⁵ plants, is essential for simulating plant response to global changes.

46 Nutrient resorption - defined here as the translocation of nutrients from senescing leaves to 47 temporary storage tissues - is a plant strategy for nutrient conservation (Killingbeck, 1996; 48 Kobe et al., 2005). It allows plants to directly reuse nutrients, decreasing the dependence on 49 soil nutrient availability and the competition for these nutrients with other plants and 50 microbes, especially in nutrient-limited environments (Aerts, 1996; Aerts and Chapin, 1999). 51 The question that arises is then why do plants not all resorb the entirety of leaf nutrients for 52 being more efficient? The fact that they don't implies the existence of costs and limitations to 53 resorption. A quantitative understanding of nutrient resorption can yield insights into plant 54 strategies to cope with nutrient limitation (Aerts and Chapin, 1999; Chapin et al., 2011). This 55 is because the resorption process influences most other ecosystem processes that determine 56 plant growth, as it directly affects litter quality and therefore soil organic matter 57 decomposition and has indirect consequences for plant nutrient uptake, carbon cycling and 58 finally plant competition (Killingbeck, 1996; Berg and McClaugherty, 2008). The average 59 fraction of leaf nutrients resorbed before abscission is estimated to be ~62% for N and ~65% 60 for P (Vergutz et al., 2013). Cleveland et al. (2013) estimated that this corresponds to 31% of 61 a plant's annual demand for N and 40% of the annual demand for P, but with large 62 geographical and species variations.





63 However, despite advances in recent years, the drivers behind nutrient resorption and its 64 variation are still unclear: First, soil fertility has long been assumed to be a key driver for 65 variations in nutrient resorption, with increased resorption in infertile soils as the plant's main 66 strategy for nutrient conservation (Aerts and Chapin, 1999). Nonetheless, there is diverging 67 evidence established at different geographic scales, showing positive correlations (Aerts and 68 Chapin, 1999), negative correlations (Yuan and Chen, 2015; Xu et al., 2021), and even a lack 69 of correlation between soil fertility and resorption efficiency (Vergutz et al., 2013). Second, 70 climate factors are also considered to be important drivers for resorption, but the evidence is 71 equally conflicting: On the one hand, Yuan and Chen (2009) and Yan et al. (2017) suggested 72 NRE is decreasing with mean annual temperature (MAT) and precipitation (MAP), with the 73 opposite trend for PRE, arguing that colder regions tend to be more N-limited, while 74 P-limitation is observed more commonly in warmer environments. From low to high latitudes 75 globally, the role of N in limiting productivity tends to increase as the availability of N is 76 mainly determined by temperature-limited processes such as biological N fixation and 77 mineralization of soil organic matter (Cleveland et al., 2013; Fay et al., 2015; Deng et al., 78 2018), but the presence of N fixers in tropical forests introduces complexity to the pattern of 79 nutrient limitation between tropical and temperate zones (Hedin et al., 2009). Nevertheless, 80 the limited availability of P in the tropics due to highly weathered soils distinguishes low- to 81 mid-latitude environments (Elser et al., 2007). On the other hand, Vergutz et al. (2013) and 82 Xu et al., 2021 showed that NRE and PRE are both increasing with decreasing MAT and 83 MAP toward higher latitudes.

A third set of studies suggests plant functional types, leaf stoichiometry and plant nutrient st demand as drivers for nutrient resorption (Reed et al., 2012; Han et al., 2013; Tang et al., 2013; Brant and Chen, 2015; Du et al., 2020; Chen et al., 2021a; Sun et al., 2023). When found greater nutrient resorption in evergreen species, it is assumed to be a conservation strategy given their comparatively low leaf nutrient content and slow growth rate and predominant occurrence in nutrient-limited biomes (Killingbeck, 1996; Yan et al., 2017; Xu et al., 2021). The same argument has been used for interpreting differences between broad-leaves and needle-leaves, in which nutrient resorption is generally observed to be higher in needles as a strategy to acclimatize and survive in resource-limited environments (Aerts and Chapin, 1999; Yuan et al., 2005; Yan et al., 2017; Xu et al., 2021). Previous





95 compared to trees, due to their smaller leaves with shorter life cycles and for the need to 96 optimize nutrient use in resource-limited environments (Killingbeck, 1996; Yuan and Chen, 97 2009; Yan et al., 2017; Xu et al., 2021). However, Brant and Chen (2015) suggest that 98 deciduous plants are more dependent on nutrient resorption as their investment in green leaf 99 nutrients is higher to maintain their fast growth through high physiological activity during the 100 growing season. Plants with a slow growth strategy, such as evergreens and needle-leaves, 101 have lower photosynthetic nutrient use efficiency due to a higher allocation of C and N to leaf 102 structural rather than metabolic compounds (Reich et al., 2017). Onoda et al. (2017) 103 empirically supports this by showing that a greater allocation of nutrients to structural 104 compounds is associated with decreased specific leaf area (SLA) and increased diffusive 105 limitation to photosynthesis. Thus, variations in leaf traits and construction costs could 106 contribute to differences in resorption between plant functional types (PFTs). Nevertheless, 107 Drenovsky et al. (2010; 2019) suggested that resorption variability is influenced by an 108 interplay of the discussed drivers, that includes soil properties, climatic conditions, and plant 109 characteristics. Estiarte et al. (2023) support that leaf biochemistry of plants determine the 110 first limitation to nutrient resorption, with a secondary regulation in resorption by 111 environmental conditions, while the costs of leaf aging remain consistent.

112 The divergence of observed patterns highlights the need for further investigation into the 113 main drivers of variations in nutrient resorption, distinguishing the influence of plant types, 114 soil and climatic conditions. In this study, we present a meta-analysis that combines the 115 version 5.0 of TRY Plant Trait database (Kattge et al., 2020) with different ancillary datasets 116 for climate and soil factors to investigate global patterns of resorption efficiencies for 117 nitrogen (NRE) and phosphorus (PRE). We aim to extend woody species observations for 118 nutrient resorption and investigate the factors that explain observed patterns along three main 119 axes: climate, soil fertility and leaf properties.

120

121 2. Methods

122 2.1 Data collection

123 We assembled the dataset from the TRY Plant Trait database (https://www.try-db.org, Kattge 124 et al., 2020, version 5.0) containing field measurements of paired leaf and litter mass-based 125 tissue N and P concentrations ($N_{mass, leaf}$, $P_{mass, leaf}$, $N_{mass, litter}$, $P_{mass, litter}$) to derive the fractional 126 nutrient resorption (described in Sect. 2.2), and plant functional traits recorded in parallel





127 from the same species and same location to consider as biological predictors variables (Table 128 1). As additional predictors for nutrient resorption, we combined it with climate and soil input 129 data (Table 2). We processed the data using R statistical software (version 4.0.4), keeping the 130 data at species-level. To manipulate the extracted functional traits, we used the package 131 {rtry} (Lam et al., 2022) developed to support the preprocessing of TRY Database (version 132 1.0.0), and {tidyverse} package (Wickham et al., 2019) with its dependencies (version 1.3.2).
133 The data processing followed the quality control according to the published protocol of TRY 134 (Kattge et al., 2011; 2020).

135

136 Table 1. Traits extracted from TRY database to derive nutrient resorption.

Plant traits

| | Variable name | Unit |
|------------------------|--|----------------------------------|
| $N_{ m mass, \ leaf}$ | Leaf nitrogen (N) content per leaf dry mass | mg g |
| $P_{ m mass, leaf}$ | Leaf phosphorus (P) content per leaf dry mass | mg g |
| $N_{ m mass,\ litter}$ | Litter nitrogen (N) content per litter dry mass | mg g |
| $P_{ m mass,\ litter}$ | Litter phosphorus (P) content per litter dry mass | mg g |
| SLA | Leaf area per leaf dry mass: petiole, rhachis and midrib excluded | $mm^2 mg^{-1}$ |
| SLA | Leaf area per leaf dry mass: petiole excluded | mm ² mg ⁻¹ |
| SLA | Leaf area per leaf dry mass: petiole included | mm ² mg ⁻¹ |
| SLA | Leaf area per leaf dry mass: undefined if petiole is in- or excluded | mm ² mg ⁻¹ |
| | Leaf dry mass | mg |
| | Leaf senescent dry mass | mg |
| LML | Leaf Mass Loss | unitless |
| PFT | Plant functional type / growth form | unitless |
| KGC | Köppen Climate Classification | unitless |

137 138

139 As predictors, we used a set of climate variables, N and P deposition, vegetation type-related 140 variables, and soil data (Table 2) with a spatial resolution of $0.5^{\circ} \times 0.5^{\circ}$ to match that of the 141 lowest resolution dataset (P deposition). Mean annual temperature (MAT), mean annual 142 precipitation (MAP) and the seasonal temperature amplitude were derived from the global 143 climate database WorldClim (Fick and Hijmans, 2017). We extracted the Köppen climate 144 classification to represent different climate zones from the TRY database and filled data gaps





145 using the {Kgc} R package (Bryant et al., 2017), which provides the Köppen climate 146 classification for each latitude and longitude. We calculated mean annual evapotranspiration 147 (ET) and growing season length (GSL) from FLUXCOM (Jung et al., 2011), in which GSL 148 was based on the seasonal phasing of gross primary productivity (GPP) considering the time 149 period between 20% and 80% of maximum GPP in an average year for the period 2002-2015. 150 Total soil P concentrations were derived from Yang et al. 2013; soil clay content and soil pH 151 were extracted from the Harmonized World Soil Database (HWSD; Wieder et al., 2014). We 152 used atmospheric N deposition values from CESM-CMIP6 (Hegglin; Kinnison and 153 Lamarque, 2016) taking the year 2010 as a reference considering that the fields are relatively 154 smooth, summing the emissions and making the annual mean, and P deposition was extracted 155 from Brahney et al. (2015) and Chien et al. (2016). All variables used as predictors of global 156 N and P resorption are described in table 2.

157

| 158 Table 2. All J | possible predi | ictors for nutrien | t resorption. |
|--------------------|----------------|--------------------|---------------|
|--------------------|----------------|--------------------|---------------|

| | Variable name | Unit | Reference |
|-----------------------|---------------------------|----------------------------------|-------------------------------|
| MAT | Mean Annual | °C | Fick and Hijmans, 2017 |
| | Temperature | | |
| MAP | Mean Annual Precipitation | mm | Fick and Hijmans, 2017 |
| AmplT | Temperature amplitude | °C | Fick and Hijmans, 2017 |
| ET | Evapotranspiration | mm | Jung et al., 2011 |
| N_dep2010 | Nitrogen deposition | kgN ha yr | Hegglin; Kinnison and |
| | | | Lamarque, 2016 |
| P_dep | Phosphorus deposition | kgN ha yr | Brahney et al., 2015; Chien e |
| | | | al., 2016 |
| soilP_tot | Total soil P | g P/m ² | Yang et al., 2013 |
| Clay | Top soil clay content | % weight | Wieder et al., 2014 |
| pН | Top soil pH | -log(H+) | Wieder et al., 2014 |
| GSL | Growing season length | days | Jung et al., 2011 |
| SLA | Specific leaf area | mm ² mg ⁻¹ | Kattge et al., 2020 |
| LLS | Leaf Longevity | month | Kattge et al., 2020 |
| Leaf habit(phenology) | Deciduous/Evergreen | - | Kattge et al., 2020 |
| Leaf Type | Broadleaves/Needles | - | Kattge et al., 2020 |

159

160 2.2 Data derivation

161 We define nutrient resorption efficiency (NuRE) as the amount of nutrient resorbed during162 leaf senescence calculated as:

164
$$NuRE = \left(1 - \frac{Nu_{senesced}}{Nu_{green}} MLCF\right) \times 100$$
(1)



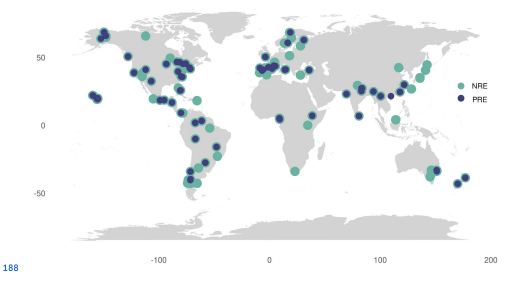


165 where Nu_{green} and Nu_{senesced} are nutrient (N or P) concentrations in dry green and senesced 166 leaves (mg g), respectively; MLCF (unitless) is the mass loss correction factor during 167 senescence to account for the loss of leaf mass when senescence occurs. Omitting MLCF 168 overestimates nutrient concentration in senescent leaves and underestimates resorption values 169 (Zhang et al., 2022). Zhang et al. (2022) showed a significant overall improvement when 170 considering MLCF, where both average of N and P resorption increased by ~9%, particularly 171 for cases with low resorption efficiencies. In the present study, not considering the MLCF 172 also underestimates the actual nutrient resorption efficiency when comparing the fraction of 173 resorption of four sub datasets from the final global dataset (Appendix A).

174 We calculated MLCF as the ratio between the dry mass of senesced and green leaves (van 175 Heerwaarden et al., 2003a), where it was not directly available as percentage leaf mass loss 176 (LML) in the data. We derived average values of MLCF per plant type from nutrient 177 resorption dataset to fill missing values: 0.712 for deciduous, 0.766 for evergreen, 0.69 for 178 conifers, and 0.75 for woody lianas, respectively. To fill in MLCF values for the remaining 179 leaf nutrient and litter data from TRY, we associated these means of MLCF with leaf habit, 180 leaf type and growth form information available on each species. For that, trees with needle 181 evergreen leaves were associated with conifers MLCF; deciduous trees/shrubs with 182 deciduous woody MLCF, and evergreen trees/shrubs with evergreen woody MLCF, 183 respectively. We grouped climbers and lianas with shrubs. In total we extracted data from 131 184 sites for NRE and 74 for PRE (Fig. 1), with more than one entry per site giving a total of 954 185 and 454 data points for NRE and PRE species-level, respectively. Temperate biomes were 186 most strongly represented in the dataset (518 entries), followed by tropical (180), boreal 187 (103), polar (102) and dry ecosystems (65).







189 Figure 1: Global distribution of data used for nitrogen resorption efficiency (NRE) and phosphorus resorption190 efficiency (PRE).

191

192 2.3 Statistical analysis

193 As the nutrient resorption data did not conform to a normal distribution (Shapiro–Wilk test), 194 we used the nonparametric Kruskal–Wallis one-way ANOVA test of variance to examine 195 differences of NRE and PRE among different climate zones, and Mann-Whitney Wilcoxon 196 test to evaluate differences between leaf habit, leaf type and growth form (deciduous vs 197 evergreen plants, broad-leaves vs needle-leaves, shrubs vs trees), using the {ggstatsplot} R 198 package (Patil, 2021). We applied Pearson correlation and linear regression to analyze the 199 relationship between nutrient resorption and the predictors described in Table 2. For MAP 200 and N deposition, we performed a log transformation prior to conducting the analysis to have 201 the distribution close to the normal. To find the best set of predictors for the variance in NRE 202 and PRE, we used multimodel inference (MMI; Burnham and Anderson, 2002) using the 203 Akaike's information criterion (AIC) and estimated the relative importance of each 204 explanatory variable. Different from setting only a single model based on AIC, multimodel 205 inference accounts for uncertainties in the model performance and in the considered 206 parameters. This approach involves modeling and evaluating all possible combinations of a 207 predetermined set of predictors. The evaluation is typically conducted using a criterion, such





208 as AIC or Bayesian information criterion (BIC), which favors simpler models and allows for 209 a comprehensive examination of all possible models and their respective performances. By 210 synthesizing the estimated coefficients of predictors across these models, MMI enables 211 inference regarding the overall importance of specific predictors. Before applying MMI, we 212 used generalized linear mixed effect models (GLMER) to fit different models after removing 213 drivers described in Table 2 that showed: (1) high collinearity between them ($R \ge 0.7$; Fig. 214 S5); (2) non-significant correlation with NRE (soil P) and PRE (MAP and SLA) (Fig. S5); 215 (3) a threshold of Variance Inflation Factor (VIF) higher than 10 (James et al. 2013). 216 Specifically, temperature amplitude, GSL and ET were not considered due to their high 217 correlation with MAT and MAP and due to high VIF. Based on ecological interactions, we 218 fitted the model considering interactions between climate variables MAT and MAP, as well as 219 between plant characteristics such as leaf structure, leaf habit and leaf type 220 (SLA:LeafPhenology:LeafType). If the ratio between the sample size and the number of 221 parameters considered was higher than 40, we fitted the model using Restricted Maximum 222 Likelihood REML and AICc (corrected for small sample sizes) to avoid bias. We selected the 223 model with lowest AIC and applied it into the 'dredge' function implemented in the 224 multimodal inference package {MuMIn} (Bartoń K, 2023) which generated a full submodel 225 set. A set of best-performing models for NRE and PRE was selected using a cut-off of Δ AIC 226 < 2, and based on these top models, the best model parameters were generated. Using 227 {MuMIn} package, we also calculated the relative importance of each predictor through the 228 sum of the Akaike weights across all models in which the respective parameter was being 229 considered, with a cut-off of 0.8 to distinguish between important and unimportant predictors 230 (Terrer et al., 2016). We performed all statistical analysis using p-value < 0.05 as statistically 231 significant.

232

233

3. Results

235 3.1 Global patterns of nutrient resorption between different climate zones

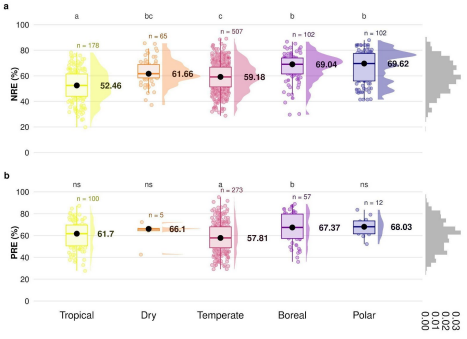
²³⁶ The global median of nutrient resorption for nitrogen (NRE) and phosphorus (PRE) is 60.0% ²³⁷ \pm 12.3% of standard deviation (n=954) and 61.2% \pm 13.6% (n=454), respectively. We find ²³⁸ differences for both NRE and PRE between the climate zones (Fig. 2). Tropical regions show ²³⁹ the lowest resorption for N (NRE: 52.4% \pm 12.1%) and tundra ecosystems in polar regions





240 show the highest (NRE: 69.6% \pm 12.8%) (Fig. 2a). PRE in temperate regions shows the 241 lowest values (57.8% \pm 13.6%). PRE increases towards the higher latitude with significant 242 difference of P resorption from temperate to boreal regions (67.3% \pm 13.6%) (Fig. 2b). In 243 contrast to NRE, the difference of PRE between tropical and other climate zones, as well as 244 polar regions, is not statistically significant (P > 0.05). NRE in dry regions (61.6% \pm 9.7%) is 245 statistically different from tropical and polar regions, while for PRE, the difference is not 246 significant between climate zones. However, the sample for this zone is substantially smaller. 247 Details of minimum, maximum, and median values can be found in Table B1.





249

250 Figure 2: Difference in the resorption efficiency of nitrogen (NRE; a) and phosphorus (PRE; b) between climate
251 zones by Köppen climate classification. Different letters indicate the significant differences in nutrient
252 resorption between the climate zones, 'ns' means non significant, and 'n' represents the number of observations.
253

254 3.2 Patterns of nutrient resorption between plant functional types

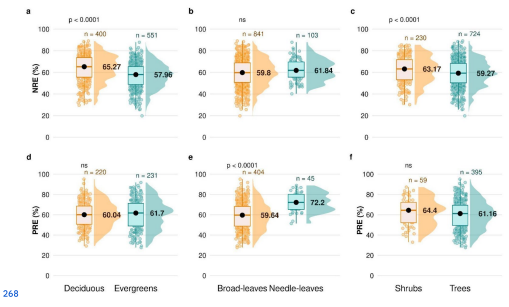
255 We explore the variation of nutrient resorption between plant functional groups. Deciduous 256 woody plants have a significantly higher NRE (65.2% \pm 12.4%, n=400) than evergreens 257 (57.9% \pm 11.4%, n=551) (P < 0.001) (Fig. 3a), and shrubs have a significantly higher NRE 258 (63.1% \pm 12.4%, n=230) than trees (59.2% \pm 12.1%, n=724) (P < 0.001) (Fig. 3c).





259 Conversely, there is no significant difference in NRE between broad- (59.8% ± 12.5%, 260 n=841) and needle-leaved plants (61.8% ± 9.9%, n=103) (P > 0.05) (Fig. 3b). PRE does 261 neither differ significantly between deciduous (60.0% ± 12.8%, n=220) and evergreen plants 262 (61.7% ± 14.4%, n=231) (P = 0.4) (Fig. 3d) nor between shrubs (64.4% ± 13.5%, n=59) and 263 trees (61.1% ± 13.6%, n=395) (P = 0.2) (Fig. 3f). However, PRE differs significantly between 264 leaf types, with needle-leaved showing higher resorption (72.2% ± 9.2%, n=45) than 265 broad-leaved plants (59.6% ± 13.5%, n=404) (P < 0.001) (Fig. 3e). Details of minimum, 266 maximum and median values can be found in Table B2.





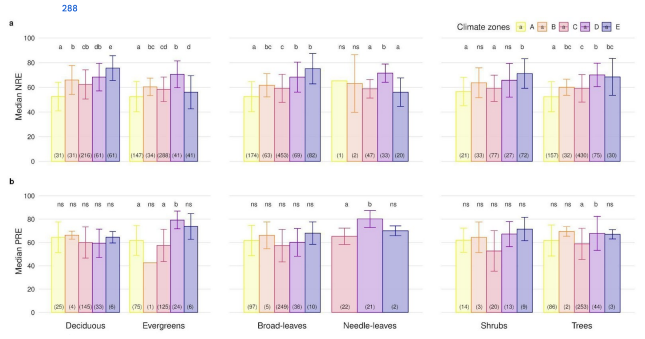
269 Figure 3: Difference in the nitrogen resorption efficiency (NRE) and phosphorus resorption efficiency (PRE)
270 between plant functional types (PFTs) on a global scale, comparing deciduous versus evergreens (a d),
271 broadleaved species versus needle leaves (b e), and shrubs versus trees (c f). 'n' represents the number of
272 observations, and 'p' indicates the significant difference of nutrient resorption between each PFT.
273

274 We next explore how climate zones affect NRE and PRE within plant functional groups. NRE 275 tends to increase from tropical to boreal climates (Fig. 4a) – a pattern seen among deciduous 276 and evergreen woody plants, among shrubs and trees, and among broadleaved, but not 277 needle-leaved plants. Also PRE increases from temperate to boreal and polar climates, but 278 declines from the tropics to temperate climates in evergreens (Fig. 4b). Apart from the overall 279 tendency, we observe a few statistical deviations from the general pattern that emerged across





280 all plants pooled: NRE is significantly lower in polar regions compared to boreal forests for 281 evergreens (NRE: 56.0% \pm 13.4%; NRE: 70.5% \pm 10.8%) and compared to needle leaved 282 plants (NRE: 56.0% \pm 11.5%; NRE: 51.5% \pm 7.3%) (P < 0.001); PRE shows the same pattern 283 deviation between these regions, but the pattern is not statistically significant (P > 0.05). 284 Also, we did not observe lower NRE for tropical regions in needle leaved plants because the 285 only observation of this plant type is in this climate zone. Details of minimum, maximum and 286 median values can be found in Table B3.



290 Figure 4: Median of nitrogen resorption efficiency (NRE; a) and phosphorus resorption efficiency (PRE; b)
291 between deciduous versus evergreens, broad- versus needle-leaves and shrubs versus trees in different climate
292 zones. Error bars are the standard deviations of the medians. Different letters indicate the significant differences
293 in nutrient resorption between the climate zones. Numbers in parentheses represent the number of observations.
294 Climate zones (A Tropical; B Dry; C Temperate; D Boreal; E Polar).

287

296 3.3 Main drivers of nutrient resorption

297 We investigate the main drivers for variation in nutrient resorption, considering biological, 298 climatic, and soil factors and using data from all PFTs and climate zones pooled. Dredge 299 model averaging based on a set of best-performing models with corrected AIC (see Methods 300 2.3) shows that the best model for NRE includes soil clay content, N deposition, MAP and

²⁹⁵





301 growth form (Table 3). The best combination of predictors for the PRE model includes N 302 deposition, leaf type, and MAT (Table 3). Sums of Akaike weights indicate that the order of 303 importance of predictors for NRE is N deposition (RI 0.99), MAP (RI 0.99), leaf habit (RI 304 0.98), followed by soil clay content (RI 0.97), growth form (RI 0.93) and leaf type (RI 0.87) 305 (Fig. 5a); while for PRE, the order is P deposition (RI 0.99), leaf type (RI 0.99), N deposition 306 (RI 0.94) followed by leaf habit (RI 0.89) (Fig. 5b). The criteria to fit the model selecting 307 and/or excluding predictors and interactions for the multimodel inference can be found in 308 Sect. 2.3. Correlations between all variables, as well as linear relationships with the 309 regression slope between nutrient resorption and all possible predictors can be found in Figs. 310 C1 and C2.

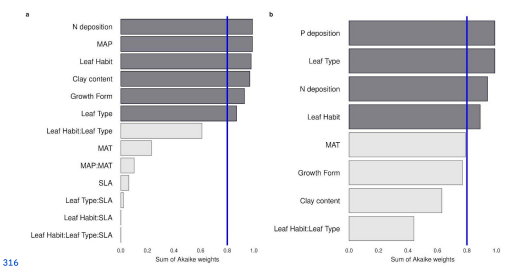
311

| 312 Table 3 Summarized results of dredge model averaging for nitrogen resorption efficiency (NRE) and |
|---|
| 313 phosphorus resorption efficiency (PRE). Significant codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1. SE |
| 314 means standard error |

| NRE | Estimate | SE | Adjusted SE | z value | Pr(> z) |
|----------------------|----------|-------|-------------|---------|------------|
| (Intercept) | 63.24 | 2.86 | 2.87 | 21.96 | <0.001 *** |
| Clay content | -0.33 | 0.09 | 0.09 | 3.54 | <0.001 *** |
| Growth Form | 2.57 | 1.11 | 1.12 | 2.30 | 0.02 * |
| Leaf habit | 2.02 | 2.32 | 2.33 | 0.86 | 0.38 |
| Leaf type | 0.66 | 2.51 | 2.52 | 0.26 | 0.79 |
| MAP | -5.07 | 1.58 | 1.58 | 3.19 | 0.001 ** |
| N deposition | 0.57 | 0.11 | 0.11 | 5.07 | <0.001 *** |
| Leaf habit:Leaf type | -0.51 | 2.69 | 2.70 | 0.19 | 0.84 |
| PRE | Estimate | SE | Adjusted SE | z value | Pr(> z) |
| (Intercept) | 78.28 | 9.45 | 9.56 | 8.18 | <0.001 *** |
| Clay content | -0.44 | 0.24 | 0.24 | 1.81 | 0.06 . |
| Growth Form | -1.35 | 2.99 | 3.03 | 0.44 | 0.65 |
| Leaf habit | 2.72 | 1.75 | 1.77 | 1.53 | 0.12 |
| Leaf type | -10.34 | 4.29 | 4.35 | 2.37 | 0.01 * |
| MAT | 1.08 | 0.49 | 0.49 | 2.18 | 0.02 * |
| N deposition | -1.77 | 0.54 | 0.54 | 3.23 | 0.001 ** |
| P deposition | -97.13 | 65.80 | 66.75 | 1.45 | 0.14 |







317 Figure 5: Importance of the abiotic and biotic predictors on nitrogen resorption efficiency (NRE; a) and **318** phosphorus resorption efficiency (PRE; b). The relative importance (RI) of each predictor is calculated through **319** the sum of the Akaike weights derived from multimodal inference selection, using corrected Akaike's **320** information criteria. The blue line distinguishes between important and unimportant predictors. Mean Annual **321** Precipitation (MAP); Mean Annual Temperature (MAT); SLA (Specific Leaf Area). Colon means interaction **322** between predictors. Leaf habit is represented as 'Leaf Phenology'. **323**

324

4. Discussion

Through an extensive global dataset of leaf nutrient resorption and a multifactorial analysis, we show that leaf habit and type are a strong driver of the spatial variation in nutrient resorption, with thicker, longer-lived leaves having lower resorption efficiencies. Climate, and soil-availability-related factors also emerge as strong drivers, in which we discuss a secondary regulation related to environmental conditions in space and time. Our study covers is significantly more woody species observations for nutrient resorption, especially for N, than previous studies (Yuan and Chen, 2009; Yan et al., 2017; Xu et al., 2021). We also account for variations in the mass loss of senescing leaves by deriving the MLCF when leaf mass loss at or leaf dry mass were available, and then apply the calculated average MLCF to the missing as data, rather than using a single average of MLCF from the literature per PFT (Yan et al., 2017; Xu et al., 2021), which may lead to a more correct estimate of nutrient resorption (see 37 Methods 2.2).





339 4.1 Nutrient resorption limited by leaf structure

The structural properties of leaves limit the efficiency of resorption along geographic and at climatic ranges. We find that the global mean for NRE is significantly higher in deciduous than evergreen plants, and is higher in shrubs than trees (discussed at the end of this section) at (Fig. 3a; 3c). This finding is in contrast to previous global studies that found decreasing at nutrient resorption with increasing green leaf nutrient content, implying that deciduous species, which generally have higher leaf N content than evergreen species, have higher at resorption (Yan et al., 2017; Xu et al., 2021). Nevertheless, our finding is in agreement with Vergutz et al (2013), who reported that deciduous woody species had higher NRE than at evergreen woody species and who found no significant differences for PRE.

We find that leaf habit is a strong driver for variation in resorption for both nutrients (Table 3; 550 Fig. 5). Fig. 3a shows that leaf habit is associated with clearly different mean NRE values for 551 evergreen and deciduous species, while the relationship of the average resorption is less clear 552 for PRE (Fig. 3d). This is likely the consequence of a dominance of evergreen species in the 553 tropics in our data set, but we cannot conclude that the lower amount of data for PRE is also a 554 drive of this pattern. The inconsistencies of patterns and significance in P resorption can be 555 related to high biochemical divergence in leaf P fractions compared to N, leading to varied 556 mobilization paths (Estiarte et al., 2023). The breakdown of proteins is the main way N 557 moves around as 75-80% of N is allocated in proteins, while P mobilization involves many 558 different catabolic pathways that lead to wider variety in P dynamics in leaves during leaf 559 development (Estiarte et al., 2023).

We observe no statistical difference between leaf types for NRE (Fig. 3). The higher PRE in a61 needle- than broad-leaves (Fig. 3e) is likely a species effect since almost all needle a62 observations for PRE are plants of the same family, *Pinaceae*. Nevertheless, leaf type is also a strong driver for variance in NRE and PRE (Table 3; Fig. 5). This finding goes together a64 with the view of thicker, longer-lived leaves - such as evergreens and needle-leaves - having a65 lower resorption efficiencies. One possible explanation for this global leaf habit and type a66 pattern is that thicker leaves from evergreens plants, i.e. those with low specific leaf area a67 (SLA), have more N allocated to structural leaf compartments, which means it is harder to a68 break down and resorb nutrients back, leading to less resorption. This is different to a69 deciduous plants, in which leaves are characterized by a higher SLA and a larger N a70 investment into metabolic compounds (Onoda et al., 2017).





371 The leaf economics spectrum (LES) distinguishes "fast" and "slow" economic strategies 372 found globally and existing independent of climate (Wright et al., 2004). A rapid return on 373 investments, or "fast" economic strategy, is typically associated with deciduous plants and 374 achieved through a combination of traits such as shorter leaf longevity, higher nutrient 375 concentrations, and thinner leaves (high specific leaf area SLA), resulting in higher gas 376 exchange rates per unit mass/area (Reich et al., 1992, 1997; Wright et al., 2004). Conversely, 377 a slow return on investments is associated with the opposite set of traits and typically found 378 in evergreen plants (Reich et al., 1992, 1997; Wright et al., 2004). The low SLA of long-lived 379 leaves is associated with low photosynthetic N-use efficiency, but with nutrient investment 380 spread over a longer period. The low photosynthetic N-use efficiency can be attributed to a 381 higher proportion of C and N being allocated to structural rather than metabolic components 382 of the leaf (Reich et al., 2017), which aligns with the theory on leaf carbon optimization 383 proposed by Kikuzawa (1995) and posits that shorter leaf longevity is associated with higher 384 photosynthetic rates or lower costs of leaf construction.

385 Here, we found that plants with a conservative nutrient resorption strategy are located at the 386 non-conservative end of the LES, that is, in the "fast" economic strategy. The discussion that 387 revolves around the LES is determined by a combination of trade-offs between investments 388 in structural and metabolic components, as well as trade-offs over time in the expected 389 returns on those investments (Reich et al., 2017). The non-transferable and possibly 390 transferable nutrients depend on where they are located in the cell and their biochemistry 391 (Estiarte et al., 2023). Metabolic fractions are considered to be fully accessible for resorption ³⁹² while structural fractions have been considered non-degradable (Estiarte et al., 2023). Wang 393 et al. (2023) brings the worldwide pattern of high leaf lifespan (LLS) in plants with low SLA 394 as a natural selection response to maximize carbon gain during leaf development, with 395 variations in SLA in deciduous and evergreen species being determined by microclimate 396 conditions. This pattern scales up from the organ level to a broader perspective that 397 encompasses the trade-off between growth and survival at the plant level (Kikuzawa and 398 Lechowicz, 2011). We found higher NRE in shrubs than trees as observed in previous studies 399 (Yuan and Chen, 2009; Yan et al., 2017; Xu et al., 2021), which is also reflected in the 400 identification of plant growth form as one of the main driving factors for NRE in the 401 multimodel inference analysis (Table 3; Fig. 5a). Compared to trees, shrubs typically have 402 smaller leaves and shorter leaf-lifespans. With that they need to be more resourceful with the





403 nutrients available and prioritize nutrient resorption as a way to optimize nutrient usage for 404 growth.

405 Resorption is an internal plant process that aims to maintain the balance of soil-plant 406 interactions in the acquisition and conservation of nutrients, considering which process is less 407 costly for the plant. The efficiency in nutrient-use by plants is determined mainly by the 408 nutrient residence time in the plant, in which they can access through the leaf longevity 409 maintaining the nutrients or through resorption before leaf abscission (Veneklaas, 2022). Our 410 results support the concept that nutrient resorption is mainly driven by the share of metabolic 411 vs total leaf N (P), which co-varies with SLA (proxy for construction costs). Therefore, 412 higher resorption in deciduous trees may be an important conservation strategy as this 413 process is less energetically costly than new growth. Brant and Chen (2015) discuss the 414 dependence of deciduous trees on nutrient resorption efficiency as their investment in green 415 leaf nutrients is higher to keep fast physiological activity during growing season, or the entire 416 nutrient economy is compromised. With that, we can argue that leaf longevity may be an 417 important strategy for evergreen plants to conserve their lower leaf nutrient content, as the 418 nutrient residence time is higher in evergreens. These plants retain nutrients for as long as 419 possible, because once the nutrients are transferred to the soil through litterfall, they are 420 partially lost from the system.

421

422 4.2 Effects of climate factors

423 Our global dataset shows that NRE significantly increases from tropical to polar zones (Fig. 424 2a), while PRE is lowest in temperate zones and significantly increases toward the poles (Fig. 425 2b). This suggests that the resorption of both nutrients is governed to some extent by a 426 comparable dependency on climate, possibly related to slowed soil organic matter 427 decomposition at lower mean annual temperatures, which reduces the net rate of 428 mineralization and in turn, limits the availability of nutrients for plant uptake from the soil 429 (Sharma and Kumar 2023). MAT emerges as one of the main drivers for PRE but not for 430 NRE (Table 3). This result may be the outcome of the overall distribution of deciduous and 431 evergreen species across climate zones, suggesting that global variations in N and P 432 resorption along climatic gradients may arise primarily from global patterns in deciduous vs. 433 evergreen and needle-leaved vs. broadleaved plants. This statement is important in the 434 context of projecting nutrient cycling under altered climate and indicates limited responses in





435 resorption to temporal changes in climate at decadal time scales – before the global436 distribution of leaf habit and type changes as a result of shifts in species composition.

437 MAP emerged as an important driver for NRE (Table 3; Fig. 5). One explanation is that low 438 MAP leads to soil moisture, constraining nutrient mobility and increasing the carbon cost for 439 plants to take up nutrients (Gill and Penuelas, 2016). Therefore, together with limited N 440 resorption mobility in leaf tissues discussed above (Estiarte and Penuelas, 2015), soil 441 moisture constrains N mobilization during the mineralization process (Thamdrup, 2012). Liu 442 et al. (2016) analyzed the relation between soil N mineralization and temperature sensitivity 443 on a global scale, and showed largest N mineralization rates at tropical latitudes and a general 444 poleward decrease. We can observe a similar pattern of NRE with latitude (Fig. C3). Deng et 445 al. (2018) observed a negative relationship between NRE and mineralisation rate, which 446 suggests a reciprocal causal relationship where systems emerge exhibiting either 447 simultaneously low mineralization and high resorption rates. The strong link found here 448 between NRE and leaf habit and leaf type - traits that are immutable within a given species -449 indicates that the variations we observe in resorption might be a possible reflection of species 450 composition with direct consequence for N cycling. It suggests that a positive feedback 451 mechanism exists that leads ecosystems to be characterized by high resorption and a slower 452 soil cycling, or vice versa (Phillips et al. (2013). For example, species adapted to low soil N 453 are favored in N-limited environments, but they also produce low-N litter that decreases 454 mineralisation and further favors their competitiveness (Chapin et al., 2011).

455 In addition, we found a negative correlation between resorption and growing season length 456 (Figs. C1). Plant strategies in regions with short growing seasons (e,g. high latitudes or 457 seasonally dry subtropical regions) are focused on nutrient conservation to maximize growth 458 during the favorable period, despite nutrient availability. In very cold and seasonal 459 environments, as seen in grassy tundra vegetation, soil nutrients are often not available 460 concurrently with plant demand (Lacroix et al., 2022), implying that it may be more 461 advantageous for plants to retain their nutrients. While we did not include growing season 462 length in the multimodel inference analysis due to its high collinearity with MAT, this aspect 463 is partially reflected in leaf habit.

464 When we separate the global patterns for different climate zones in plant functional types 465 (PFTs), our results show that the major climatic pattern is consistent across the growth forms 466 and leaf types and leaf habit (Fig. 4), in which NRE and PRE increases towards higher





467 latitudes and PRE shows a minimum at mid-latitudes. Our findings support that maximum 468 NRE and PRE may be firstly constrained by leaf properties, with secondary effects from 469 climate and soil texture (discussed below). Estiarte et al. (2023) suggest that a plant's leaf 470 biochemistry (biochemical and subcellular fractions of N and P) is the primary factor in 471 limiting nutrient resorption, followed by secondary regulation related to environmental 472 conditions in space and time. They present that resorption efficiency declines when soil 473 nutrient availability rises, as plant uptake becomes less costly in more fertile soil. However, 474 the expenses linked to aging leaves remain constant (Estiarte et al., 2023).

475

476 4.3 Effect of soil nutrient availability

477 N and P deposition and clay content emerged as important predictors for both PRE and NRE 478 (Table 3; Fig. 5). This reflects likely the influence of soil N and P availability for NRE and 479 PRE. Clay content is an important factor determining the nutrient retention capacity and 480 cation exchange capacity in soils (Chapin et al., 2011). Chronic N deposition has increased 481 soil N availability (Galloway et al., 2004) and leaf nutrient content (Chapin et al., 2011) over 482 the 20th century, and likely affected plant internal recycling and resorption as indicated by our 483 spatial results. In a fertilization experiment, higher P input had a negative effect on both NRE 484 and PRE (Yuan & Chen, 2015), suggesting that increased P deposition may reduce the plant 485 internal recycling and thus resorption. The cycling and accessibility of soil P are influenced 486 by N deposition (Marklein and Houlton, 2012) through various mechanisms including 487 changes in plant P use strategies (Dalling et al., 2016; Wu et al., 2020a). Higher N deposition 488 tends to reduce total soil P content (Sardans et al., 2016) so plants would need to increase 489 PRE to compensate for the high soil N:P stoichiometry and P limitation. Jonard et al. (2014) 490 suggests that forest ecosystems are becoming less efficient at recycling P due to excessive N 491 input and climatic stress. This observation likely contributes to our finding that N and P 492 deposition emerge as a stronger driver in a negative correlation with PRE (Fig. 5; Figs. C1). 493 The lack of effect by total soil P on NRE and PRE may result from the fact that this variable 494 does not represent the actual fraction of P available for plant uptake.

495 Another soil factor found to be important for nutrient resorption is the clay content (Table 3).
496 Clay minerals are formed during soil weathering and have high surface area that influences
497 the soil's water retention capacity, and a negative charge that enables nutrients retention and
498 exchange with plant roots (Chapin et al., 2011). High-latitude soils that are younger and





499 experience slow rates of chemical weathering usually have low clay content and therefore, 500 less potential for mineral nutrient storage, which may affect their availability for plant uptake 501 (Chapin et al., 2011). As a result, plants in these environments need to invest more in 502 resorption. Thus, together with MAP and MAT, soil clay content is also closely related to soil ⁵⁰³ nutrient supply on a global scale, which is reflected in its role as driving resorption (Table 3; 504 Fig. 5), as well as in the negative correlation between clay content and nutrient resorption 505 (Figs. C1). In the context of an important effect on nutrient resorption found for leaf 506 properties together with climate, soil texture and soil fertility - previously suggested to be 507 important (Aerts and Chapin, 1999; Yuan and Chen, 2015; Xu et al., 2021) - may indicate that 508 biological and environmental factors are not fully independent, as it is also determined by 509 multiple elements such as litter quality, precipitation, parental materials and soil texture. For 510 example, P availability is geologically and pedologically limited in warm environments, 511 which means mainly determined by soil parent materials (Augusto et al., 2017), and 512 therefore, soil texture becomes an important factor for P limitation in tropical regions. Also, 513 the role of P deposition in relation to plant demand is high for tropical forests (Van 514 Langenhove et al., 2020) but low worldwide (Cleveland et al., 2013). PRE in the tropics did 515 not differ statistically from other climate zones although we observe an increase of PRE from 516 mid to low latitudes (Figs. B1b and C3), which could indicate data limitation for PRE. The 517 combination of plant properties with an underlying soil and climate control as driving factors 518 for resorption variation is also supported by Drenovsky et al. (2010; 2019), who suggested a 519 combination of soil properties, climatic factors, and plant morphology to explain changes in 520 nutrient resorption.

521

522 4.4 Data uncertainties and implications

523 Our study contributes to the existing research on nutrient resorption by using a 524 comprehensive approach to derive resorption values from the TRY database. However, we 525 encountered limitations in this derivation due to lack or limited quality of data. The absence 526 of co-located nutrient measurements in leaf and litter led to a shortage of suitable data pairs, 527 mainly for PRE, in which the robustness of the model selection raised concerns about its 528 reliability. While our approach of accounting for the MLCF improved estimates of resorption 529 (Appendix A), we could not estimate the MLCF for all data pairs, and could not fill all gaps 530 using average functional type characteristics due to lacking trait attributes in the TRY





531 database. These two factors reduced the number of data points available for statistical 532 analysis using multi-model inference. In addition, although recognized the importance of leaf 533 lifespan (LLS), it was not possible to analyze the relationship between resorption and LLS 534 due to the few measurements of this functional trait. Nevertheless, applying the available 535 statistical methods to analyze the drivers behind NRE and PRE, we found consistent patterns 536 for the key gradients of climate, soil and plant functional type, that are informative for other 537 studies despite remaining unexplained variance. In order to improve the depth of resorption 538 investigation, we encourage researchers in field work to perform concurrent measurements of 539 litter nutrient content as well as leaf and litter dry mass.

The statistical analysis of dredge multi-model inference is dependent on the specific factors used in the analysis. We removed highly collinear variables and tested the impact of different factors of factors. Although such a change in factors affected the exact number of data points used in each multi-model inference, the overall identification of important and less important factors for NRE and PRE was robust, especially for PFTs.

545 By quantifying these trends that we have found, we can delve deeper into ecosystem models 546 by improving model parametrization and developing a dynamic nutrient resorption concept. 547 Studies that utilize data to infer nutrient cycling frequently simplify resorption making 548 general assumptions (Finzi et al., 2007; Cleveland et al., 2013), or simply representing this 549 process as a fixed value of 50% (Vergutz et al., 2013; Zaehle et al. 2014), which may cause 550 inaccuracies in their findings on nutrient cycling. The flow of recycling nutrients in land 551 surface models is a factor that determines how strong the soil nutrient availability controls 552 plant production. N resorption and N uptake in the FUN model (Fisher et al., 2010), for 553 example, is defined by the relative acquisition cost of the two sources. They discuss that the 554 cost of resorption assumes a constant based on global observations, but it may require a 555 clearer connection to leaf physiology. Here, we provide a start for a statistical model that can 556 connect resorption and plant properties and restrict how much plants could actually resorb 557 nutrients, as well as the dataset to test the predictions of a physiological model. In addition, 558 environmental drivers that have been shown to influence the overall patterns, such as soil 559 texture and climate, could be considered to influence the resorption efficiency after primary 560 leaf physiology limitation. Such information is essential when estimating how it can constrain 561 carbon assimilation in face of global changes (Galloway et al., 2008), and therefore, essential





562 to predict future plant growth and the capacity of the forest to act as a carbon sink (Thornton 563 et al., 2007; Arora et al., 2022).

564

565 5. Conclusions

566 Our analysis of the global plant trait database indicates that variations of NRE and PRE are 567 driven by the combination of plant properties with an additional soil and climate control. 568 Systematic variations of NRE across leaf habit and type indicate that these traits are linked to 569 plant nutrient use and conservation strategies and that leaf structure plays an important role in 570 determining the proportion of nutrients that can be resorbed. Different metrics of soil fertility 571 and soil-related variables were tested and found to have an influence on NRE and PRE 572 together with climatic variables and leaf structure and habit. Clay content, N and P deposition 573 had strong influence with a negative relationship - possibly an expression of its role in 574 nutrient retention - as well as MAP. These trends provide a target to benchmark the 575 simulation of nutrient recycling in global nutrient-enabled models. A focus on considering 576 the links between leaf structure and nutrient resorption efficiency should enable a more 577 realistic consideration of ecological and environmental controls on nutrient cycling and 578 limitation than the current state-of-the-art. The importance of intrinsic plant properties raises 579 important questions about the flexibility of leaf resorption under future changes in climate, 580 CO₂ concentrations and atmospheric deposition.

581

582 Acknowledgments

583 This work was supported by the European Research Council (ERC) under the European 584 Union's Horizon 2020 research and innovation programme (QUINCY; grant no. 647204). 585 BDS was funded by the Swiss National Science Foundation grant PCEFP2_181115. We 586 extend our thanks to our external reviewer Katrin Fleisher, for her helpful comments on the 587 manuscript.

588 Author contributions

589 GS, SC and SZ designed the study. GS performed the analysis. All authors contributed to 590 interpreting the results. GS drafted the manuscripts; all authors contributed to writing and 591 editing the manuscript.





592 Data Availability Statement

593 All data used in this study is publicly available through the TRY database 594 https://www.try-db.org/.

595 **Conflict of Interests**

596 SZ is a member of the editorial board of Biogeosciences.

597 **References**

598 Aerts, R.: Nutrient Resorption from Senescing Leaves of Perennials: Are there General 599 Patterns?, Journal of Ecology, 84, 1996.

600

601 Aerts, R. and Chapin, F. S.: The Mineral Nutrition of Wild Plants Revisited: A Re-evaluation602 of Processes and Patterns, in: Advances in Ecological Research, vol. 30, edited by: Fitter, A.603 H. and Raffaelli, D. G., Academic Press, 1–67, 1999.

604

605 Arora, V. K., Seiler, C., Wang, L., and Kou-Giesbrecht, S.: Towards an ensemble-based 606 evaluation of land surface models in light of uncertain forcings and observations, EGUsphere, 607 https://doi.org/10.5194/egusphere-2022-641, 2022.

608

609 Augusto, L., Achat, D. L., Jonard, M., Vidal, D., and Ringeval, B.: Soil parent material-A 610 major driver of plant nutrient limitations in terrestrial ecosystems, Glob. Chang. Biol., 23, 611 3808–3824, 2017.

612

613 Barton, K.: MuMIn : multi-model inference, R package version 0.12.0, 2009.

614

615 Bazzaz, F. A.: The Response of Natural Ecosystems to the Rising Global CO₂ Levels, Annu.
616 Rev. Ecol. Syst., 21, 167–196, 1990.

617

618 Berg, B. and McClaugherty, C. A.: Plant Litter. Decomposition, Humus Formation, Carbon 619 Sequestration, Springer Verlag, 2014.

620

621

622 Brahney, J., Mahowald, N., Ward, D. S., Ballantyne, A. P., and Neff, J. C.: Is atmospheric623 phosphorus pollution altering global alpine Lake stoichiometry?, Global Biogeochem.624 Cycles, 29, 1369–1383, 2015.

625

626 Brant, A. N. and Chen, H. Y. H.: Patterns and Mechanisms of Nutrient Resorption in Plants, 627 CRC Crit. Rev. Plant Sci., 34, 471–486, 2015.





629 Bryant, C., Wheeler, N. R., Rubel, F., French, R. H.: kgc: Koeppen-Geiger Climatic Zones, R 630 package version 1.0.0.2, 2017.

631

632 Burnham, K. P. and Anderson, D. R.: Model Selection and Inference, Springer New York, 20 633 pp., n.d.

634

635 Chapin, F. S.: The Mineral Nutrition of Wild Plants, Annu. Rev. Ecol. Syst., 11, 233–260, 636 1980.

637

638 ChapinIII, F. S., Matson, P. A., and Vitousek, P. M.: Principles of Terrestrial Ecosystem 639 Ecology, Springer, New York, NY, 2011.

640

641 Chen, H., Reed, S. C., Lü, X., Xiao, K., Wang, K., and Li, D.: Coexistence of multiple leaf
642 nutrient resorption strategies in a single ecosystem, Sci. Total Environ., 772, 144951, 2021.
643

644 Chien, C.-T., Mackey, K. R. M., Dutkiewicz, S., Mahowald, N. M., Prospero, J. M., and 645 Paytan, A.: Effects of African dust deposition on phytoplankton in the western tropical 646 Atlantic Ocean off Barbados, Global Biogeochem. Cycles, 30, 716–734, 2016.

647

648 Cleveland, C. C., Houlton, B. Z., Smith, W. K., Marklein, A. R., Reed, S. C., Parton, W., Del 649 Grosso, S. J., and Running, S. W.: Patterns of new versus recycled primary production in the 650 terrestrial biosphere, Proc. Natl. Acad. Sci. U. S. A., 110, 12733–12737, 2013.

651

652 Dalling, J. W., Heineman, K., Lopez, O. R., Wright, S. J., and Turner, B. L.: Nutrient
653 Availability in Tropical Rain Forests: The Paradigm of Phosphorus Limitation, in: Tropical
654 Tree Physiology: Adaptations and Responses in a Changing Environment, edited by:
655 Goldstein, G. and Santiago, L. S., Springer International Publishing, Cham, 261–273, 2016.
656

657 Deng, M., Liu, L., Jiang, L., Liu, W., Wang, X., Li, S., Yang, S., and Wang, B.: Ecosystem 658 scale trade-off in nitrogen acquisition pathways, Nat Ecol Evol, 2, 1724–1734, 2018.

659

660 Drenovsky, R. E., James, J. J., and Richards, J. H.: Variation in nutrient resorption by desert 661 shrubs, J. Arid Environ., 74, 1564–1568, 2010.

662

663 Drenovsky, R. E., Pietrasiak, N., and Short, T. H.: Global temporal patterns in plant nutrient 664 resorption plasticity, Glob. Ecol. Biogeogr., 28, 728–743, 2019.

665

666 Du, E., Terrer, C., Pellegrini, A. F. A., Ahlström, A., van Lissa, C. J., Zhao, X., Xia, N., Wu, 667 X., and Jackson, R. B.: Global patterns of terrestrial nitrogen and phosphorus limitation, 668 https://doi.org/10.1038/s41561-019-0530-4, 2020.





670 Elser, J. J., Bracken, M. E. S., Cleland, E. E., Gruner, D. S., Harpole, W. S., Hillebrand, H.,
671 Ngai, J. T., Seabloom, E. W., Shurin, J. B., and Smith, J. E.: Global analysis of nitrogen and
672 phosphorus limitation of primary producers in freshwater, marine and terrestrial ecosystems,
673 Ecol. Lett., 10, 1135–1142, 2007.

674

675 Estiarte, M., Campioli, M., Mayol, M., and Penuelas, J.: Variability and limits of nitrogen and676 phosphorusresorptionduringfoliarsenescence,PlantComm,4,677 https://doi.org/10.1016/j.xplc.2022.100503, 2023.

678

679 Fay, P. A., Prober, S. M., Harpole, W. S., Knops, J. M. H., Bakker, J. D., Borer, E. T., Lind, E.
680 M., MacDougall, A. S., Seabloom, E. W., Wragg, P. D., Adler, P. B., Blumenthal, D. M.,
681 Buckley, Y. M., Chu, C., Cleland, E. E., Collins, S. L., Davies, K. F., Du, G., Feng, X., Firn,
682 J., Gruner, D. S., Hagenah, N., Hautier, Y., Heckman, R. W., Jin, V. L., Kirkman, K. P., Klein,
683 J., Ladwig, L. M., Li, Q., McCulley, R. L., Melbourne, B. A., Mitchell, C. E., Moore, J. L.,
684 Morgan, J. W., Risch, A. C., Schütz, M., Stevens, C. J., Wedin, D. A., and Yang, L. H.:
685 Grassland productivity limited by multiple nutrients, Nat Plants, 1, 15080, 2015.

686

687 Fick, S. E. and Hijmans, R. J.: WorldClim 2: new 1-km spatial resolution climate surfaces for 688 global land areas, Int. J. Climatol., 37, 4302–4315, 2017.

689

690 Finzi, A. C., Norby, R. J., Calfapietra, C., Gallet-Budynek, A., Gielen, B., Holmes, W. E., 691 Hoosbeek, M. R., Iversen, C. M., Jackson, R. B., Kubiske, M. E., Ledford, J., Liberloo, M., 692 Oren, R., Polle, A., Pritchard, S., Zak, D. R., Schlesinger, W. H., and Ceulemans, R.: 693 Increases in nitrogen uptake rather than nitrogen-use efficiency support higher rates of 694 temperate forest productivity under elevated CO₂, Proc. Natl. Acad. Sci. U. S. A., 104, 695 14014–14019, 2007.

696

697 Fisher, J. B., Sitch, S., Malhi, Y., Fisher, R. A., Huntingford, C., and Tan, S.-Y.: Carbon cost 698 of plant nitrogen acquisition: A mechanistic, globally applicable model of plant nitrogen 699 uptake, retranslocation, and fixation, Global Biogeochem. Cycles, 24, 700 https://doi.org/10.1029/2009gb003621, 2010.

701

702 Galloway, J. N., Dentener, F. J., Capone, D. G., Boyer, E. W., Howarth, R. W., Seitzinger, S.
703 P., Asner, G. P., Cleveland, C. C., Green, P. A., Holland, E. A., Karl, D. M., Michaels, A. F.,
704 Porter, J. H., Townsend, A. R., and Vöosmarty, C. J.: Nitrogen Cycles: Past, Present, and
705 Future, Biogeochemistry, 70, 153–226, 2004.

706

707 Galloway, J. N., Townsend, A. R., Erisman, J. W., Bekunda, M., Cai, Z., Freney, J. R.,
708 Martinelli, L. A., Seitzinger, S. P., and Sutton, M. A.: Transformation of the nitrogen cycle:
709 recent trends, questions, and potential solutions, Science, 320, 889–892, 2008.





711 Güsewell, S.: N : P ratios in terrestrial plants: variation and functional significance, New **712** Phytol., 164, 243–266, 2004.



714 Han, W., Tang, L., Chen, Y., and Fang, J.: Relationship between the relative limitation and
715 resorption efficiency of nitrogen vs phosphorus in woody plants, PLoS One, 8, e83366, 2013.
716

717 Hedin, L. O., Brookshire, E. N. J., Menge, D. N. L., and Barron, A. R.: The Nitrogen Paradox718 in Tropical Forest Ecosystems, Annu. Rev. Ecol. Evol. Syst., 40, 613–635, 2009.

719

720 James, G., Witten, D., Hastie, T., and Tibshirani, R.: An Introduction to Statistical Learning,721 Springer US, 15 pp., n.d.

722

723 Jonard, M., Fürst, A., Verstraeten, A., Thimonier, A., Timmermann, V., Potočić, N., Waldner,
724 P., Benham, S., Hansen, K., Merilä, P., Ponette, Q., de la Cruz, A. C., Roskams, P., Nicolas,
725 M., Croisé, L., Ingerslev, M., Matteucci, G., Decinti, B., Bascietto, M., and Rautio, P.: Tree
726 mineral nutrition is deteriorating in Europe, Glob. Chang. Biol., 21, 418–430, 2015.

727

Joswig, J. S., Wirth, C., Schuman, M. C., Kattge, J., Reu, B., Wright, I. J., Sippel, S. D.,
Rüger, N., Richter, R., Schaepman, M. E., van Bodegom, P. M., Cornelissen, J. H. C., Díaz,
S., Hattingh, W. N., Kramer, K., Lens, F., Niinemets, Ü., Reich, P. B., Reichstein, M.,
Römermann, C., Schrodt, F., Anand, M., Bahn, M., Byun, C., Campetella, G., Cerabolini, B.
E. L., Craine, J. M., Gonzalez-Melo, A., Gutiérrez, A. G., He, T., Higuchi, P., Jactel, H.,
Kraft, N. J. B., Minden, V., Onipchenko, V., Peñuelas, J., Pillar, V. D., Sosinski, Ê.,
Soudzilovskaia, N. A., Weiher, E., and Mahecha, M. D.: Climatic and soil factors explain the
two-dimensional spectrum of global plant trait variation, Nat Ecol Evol, 6, 36–50, 2022.

737 Jung, M., Reichstein, M., Margolis, H. A., Cescatti, A., Richardson, A. D., Arain, M. A.,
738 Arneth, A., Bernhofer, C., Bonal, D., Chen, J., Gianelle, D., Gobron, N., Kiely, G., Kutsch,
739 W., Lasslop, G., Law, B. E., Lindroth, A., Merbold, L., Montagnani, L., Moors, E. J., Papale,
740 D., Sottocornola, M., Vaccari, F., and Williams, C.: Global patterns of land-atmosphere fluxes
741 of carbon dioxide, latent heat, and sensible heat derived from eddy covariance, satellite, and
742 meteorological observations, J. Geophys. Res., 116, https://doi.org/10.1029/2010jg001566,
743 2011.

745 Kattge, J., Díaz, S., Lavorel, S., Prentice, I. C., Leadley, P., Bönisch, G., Garnier, E., Westoby,
746 M., Reich, P. B., Wright, I. J., Cornelissen, J. H. C., Violle, C., Harrison, S. P., Van
747 BODEGOM, P. M., Reichstein, M., Enquist, B. J., Soudzilovskaia, N. A., Ackerly, D. D.,
748 Anand, M., Atkin, O., Bahn, M., Baker, T. R., Baldocchi, D., Bekker, R., Blanco, C. C.,
749 Blonder, B., Bond, W. J., Bradstock, R., Bunker, D. E., Casanoves, F., Cavender-Bares, J.,
750 Chambers, J. Q., Chapin, F. S., Iii, Chave, J., Coomes, D., Cornwell, W. K., Craine, J. M.,
751 Dobrin, B. H., Duarte, L., Durka, W., Elser, J., Esser, G., Estiarte, M., Fagan, W. F., Fang, J.,

⁷⁴⁴





752 Fernández-Méndez, F., Fidelis, A., Finegan, B., Flores, O., Ford, H., Frank, D., Freschet, G.
753 T., Fyllas, N. M., Gallagher, R. V., Green, W. A., Gutierrez, A. G., Hickler, T., Higgins, S. I.,
754 Hodgson, J. G., Jalili, A., Jansen, S., Joly, C. A., Kerkhoff, A. J., Kirkup, D., Kitajima, K.,
755 Kleyer, M., Klotz, S., Knops, J. M. H., Kramer, K., Kühn, I., Kurokawa, H., Laughlin, D.,
756 Lee, T. D., Leishman, M., Lens, F., Lenz, T., Lewis, S. L., Lloyd, J., Llusià, J., Louault, F.,
757 Ma, S., Mahecha, M. D., Manning, P., Massad, T., Medlyn, B. E., Messier, J., Moles, A. T.,
758 Müller, S. C., Nadrowski, K., Naeem, S., Niinemets, Ü., Nöllert, S., Nüske, A., Ogaya, R.,
759 Oleksyn, J., Onipchenko, V. G., Onoda, Y., Ordoñez, J., Overbeck, G., et al.: TRY - a global
760 database of plant traits, Glob. Chang. Biol., 17, 2905–2935, 2011.

762 Kattge, J., Bönisch, G., Díaz, S., Lavorel, S., Prentice, I. C., Leadley, P., Tautenhahn, S., 763 Werner, G. D. A., Aakala, T., Abedi, M., Acosta, A. T. R., Adamidis, G. C., Adamson, K., 764 Aiba, M., Albert, C. H., Alcántara, J. M., Alcázar C, C., Aleixo, I., Ali, H., Amiaud, B., 765 Ammer, C., Amoroso, M. M., Anand, M., Anderson, C., Anten, N., Antos, J., Apgaua, D. M. 766 G., Ashman, T.-L., Asmara, D. H., Asner, G. P., Aspinwall, M., Atkin, O., Aubin, I., 767 Baastrup-Spohr, L., Bahalkeh, K., Bahn, M., Baker, T., Baker, W. J., Bakker, J. P., Baldocchi, 768 D., Baltzer, J., Banerjee, A., Baranger, A., Barlow, J., Barneche, D. R., Baruch, Z., 769 Bastianelli, D., Battles, J., Bauerle, W., Bauters, M., Bazzato, E., Beckmann, M., Beeckman, 770 H., Beierkuhnlein, C., Bekker, R., Belfry, G., Belluau, M., Beloiu, M., Benavides, R., 771 Benomar, L., Berdugo-Lattke, M. L., Berenguer, E., Bergamin, R., Bergmann, J., Bergmann 772 Carlucci, M., Berner, L., Bernhardt-Römermann, M., Bigler, C., Bjorkman, A. D., Blackman, 773 C., Blanco, C., Blonder, B., Blumenthal, D., Bocanegra-González, K. T., Boeckx, P., 774 Bohlman, S., Böhning-Gaese, K., Boisvert-Marsh, L., Bond, W., Bond-Lamberty, B., Boom, 775 A., Boonman, C. C. F., Bordin, K., Boughton, E. H., Boukili, V., Bowman, D. M. J. S., 776 Bravo, S., Brendel, M. R., Broadley, M. R., Brown, K. A., Bruelheide, H., Brumnich, F., 777 Bruun, H. H., Bruy, D., Buchanan, S. W., Bucher, S. F., Buchmann, N., Buitenwerf, R., 778 Bunker, D. E., et al.: TRY plant trait database - enhanced coverage and open access, Glob. 779 Chang. Biol., 26, 119–188, 2020.

780

781 Kikuzawa, K.: Leaf phenology as an optimal strategy for carbon gain in plants, Can. J. Bot.,782 https://doi.org/10.1139/b95-019, 1995.

783

784 Kikuzawa, K. and Lechowicz, M. J.: Ecology of leaf longevity, 2011th ed., Springer, Tokyo, 785 Japan, 147 pp., 2011.

786

787 Killingbeck, K. T.: Nutrients in senesced leaves: Keys to the search for potential resorption 788 and resorption proficiency, Ecology, 77, 1716–1727, 1996.

789

790 Kobe, R. K., Lepczyk, C. A., and Iyer, M.: Resorption efficiency decreases with increasing 791 green leaf nutrients in a global data set, Ecology, 86, 2780–2792, 2005.





793 Lacroix, F., Zaehle, S., Caldararu, S., Schaller, J., Stimmler, P., Holl, D., Kutzbach, L., and
794 Goeckede, M.: Decoupling of permafrost thaw and vegetation growth could mean both
795 ongoing nutrient limitation and an emergent source of N2O in high latitudes, Earth and Space
796 Science Open Archive, https://doi.org/10.1002/essoar.10510605.1, 2022.

798 Lam, O. H. Y., Tautenhahn, S., Walther, G., Boenisch, G., Baddam, P., and Kattge, J.: The799 "rtry"Rpackageforpreprocessingplanttraitdata,800 https://doi.org/10.5194/egusphere-egu22-13251, 2022.

801

802 Lang, F., Bauhus, J., Frossard, E., George, E., Kaiser, K., Kaupenjohann, M., Krüger, J.,
803 Matzner, E., Polle, A., Prietzel, J., Rennenberg, H., and Wellbrock, N.: Phosphorus in forest
804 ecosystems: New insights from an ecosystem nutrition perspective, J. Plant Nutr. Soil Sci.,
805 179, 129–135, 2016.

806

807 LeBauer, D. S. and Treseder, K. K.: Nitrogen limitation of net primary productivity in808 terrestrial ecosystems is globally distributed, Ecology, 89, 371–379, 2008.

809

810 Liu, Y., Wang, C., He, N., Wen, X., Gao, Y., Li, S., Niu, S., Butterbach-Bahl, K., Luo, Y., and
811 Yu, G.: A global synthesis of the rate and temperature sensitivity of soil nitrogen
812 mineralization: latitudinal patterns and mechanisms, Glob. Chang. Biol., 23, 455–464, 2017.
813

814 Luo, Y., Su, B., Currie, W. S., Dukes, J. S., Finzi, A., Hartwig, U., Hungate, B., McMurtrie,
815 R. E., Oren, R., Parton, W. J., Pataki, D. E., Shaw, R. M., Zak, D. R., and Field, C. B.:
816 Progressive Nitrogen Limitation of Ecosystem Responses to Rising Atmospheric Carbon
817 Dioxide, Bioscience, 54, 731–739, 2004.

819 Marklein, A. R. and Houlton, B. Z.: Nitrogen inputs accelerate phosphorus cycling rates
820 across a wide variety of terrestrial ecosystems, New Phytol., 193, 696–704, 2012.
821

822 Onoda, Y., Wright, I. J., Evans, J. R., Hikosaka, K., Kitajima, K., Niinemets, Ü., Poorter, H., 823 Tosens, T., and Westoby, M.: Physiological and structural tradeoffs underlying the leaf 824 economics spectrum, New Phytol., 214, 1447–1463, 2017.

825

826 Patil, I.: Visualizations with statistical details: The "ggstatsplot" approach, J. Open Source 827 Softw., 6, 3167, 2021.

828

829 Phillips, R. P., Brzostek, E., and Midgley, M. G.: The mycorrhizal-associated nutrient 830 economy: a new framework for predicting carbon-nutrient couplings in temperate forests, 831 New Phytol., 199, 41–51, 2013.

⁸¹⁸





833 Reed, S. C., Townsend, A. R., Davidson, E. A., and Cleveland, C. C.: Stoichiometric patterns 834 in foliar nutrient resorption across multiple scales, New Phytol., 196, 173–180, 2012. 835 836 Reich, P. B. and Flores-Moreno, H.: Peeking beneath the hood of the leaf economics 837 spectrum, New Phytol., 214, 1395–1397, 2017. 838 839 Reich, P. B., Walters, M. B., and Ellsworth, D. S.: Leaf Life-Span in Relation to Leaf, Plant, 840 and Stand Characteristics among Diverse Ecosystems, Ecol. Monogr., 62, 365–392, 1992. 841 842 Reich, P. B., Walters, M. B., and Ellsworth, D. S.: From tropics to tundra: global convergence 843 in plant functioning, Proc. Natl. Acad. Sci. U. S. A., 94, 13730–13734, 1997. 844 845 Reich, P. B., Rich, R. L., Lu, X., Wang, Y.-P., and Oleksyn, J.: Biogeographic variation in 846 evergreen conifer needle longevity and impacts on boreal forest carbon cycle projections, 847 Proc. Natl. Acad. Sci. U. S. A., 111, 13703-13708, 2014. 848 849 Sardans, J., Alonso, R., Janssens, I. A., Carnicer, J., Vereseglou, S., Rillig, M. C., 850 Fernández-Martínez, M., Sanders, T. G. M., and Peñuelas, J.: Foliar and soil concentrations 851 and stoichiometry of nitrogen and phosphorous across E uropean P inus sylvestris forests: 852 relationships with climate, N deposition and tree growth, Funct. Ecol., 30, 676–689, 2016. 853 854 Sharma, P. K. and Kumar, S.: Soil Temperature and Plant Growth, in: Soil Physical 855 Environment and Plant Growth: Evaluation and Management, edited by: Sharma, P. K. and 856 Kumar, S., Springer International Publishing, Cham, 175–204, 2023. 857 858 Sun, X., Li, D., Lü, X., Fang, Y., Ma, Z., Wang, Z., Chu, C., Li, M., and Chen, H.: 859 Widespread controls of leaf nutrient resorption by nutrient limitation and stoichiometry, 860 Funct. Ecol., 37, 1653-1662, 2023. 861 862 Tang, L., Han, W., Chen, Y., and Fang, J.: Resorption proficiency and efficiency of leaf 863 nutrients in woody plants in eastern China, J Plant Ecol, 6, 408–417, 2013. 864 865 Terrer, C., Vicca, S., Hungate, B. A., Phillips, R. P., and Prentice, I. C.: Mycorrhizal 866 association as a primary control of the CO₂ fertilization effect, Science, 353, 72–74, 2016. 867 868 Thornton, P. E., Lamarque, J.-F., Rosenbloom, N. A., and Mahowald, N. M.: Influence of 869 carbon-nitrogen cycle coupling on land model response to CO2fertilization and climate 870 variability, Global Biogeochem. Cycles, 21, https://doi.org/10.1029/2006gb002868, 2007.





872 Van Heerwaarden, L. M., Toet, S., and Aerts, R.: Current measures of nutrient resorption 873 efficiency lead to a substantial underestimation of real resorption efficiency: facts and 874 solutions, Oikos, 101, 664–669, 2003.



876 Van Langenhove, L., Verryckt, L. T., Bréchet, L., Courtois, E. A., Stahl, C., Hofhansl, F.,
877 Bauters, M., Sardans, J., Boeckx, P., Fransen, E., Peñuelas, J., and Janssens, I. A.:
878 Atmospheric deposition of elements and its relevance for nutrient budgets of tropical forests,
879 Biogeochemistry, 149, 175–193, 2020.

880

881 Veneklaas, E. J.: Phosphorus resorption and tissue longevity of roots and leaves – importance 882 for phosphorus use efficiency and ecosystem phosphorus cycles, Plant Soil, 476, 627–637, 883 2022.

884

885 Vergutz, L., Manzoni, S., Porporato, A., Novais, R. F., and Jackson, R. B.: Global resorption
886 efficiencies and concentrations of carbon and nutrients in leaves of terrestrial plants, Ecol.
887 Monogr., 82, 205–220, 2012.

888

889 Wang, H., Prentice, I. C., Wright, I. J., Warton, D. I., Qiao, S., Xu, X., Zhou, J., Kikuzawa, 890 K., and Stenseth, N. C.: Leaf economics fundamentals explained by optimality principles, Sci 891 Adv, 9, eadd5667, 2023.

892

893 Wickham, H., Averick, M., Bryan, J., Chang, W., McGowan, L., François, R., Grolemund,
894 G., Hayes, A., Henry, L., Hester, J., Kuhn, M., Pedersen, T., Miller, E., Bache, S., Müller, K.,
895 Ooms, J., Robinson, D., Seidel, D., Spinu, V., Takahashi, K., Vaughan, D., Wilke, C., Woo,
896 K., and Yutani, H.: Welcome to the tidyverse, J. Open Source Softw., 4, 1686, 2019.

897

898 Wieder, W.: Regridded Harmonized World Soil Database v1.2, 899 https://doi.org/10.3334/ORNLDAAC/1247, 2014.

900

901 Wright, I. J., Reich, P. B., Westoby, M., Ackerly, D. D., Baruch, Z., Bongers, F.,
902 Cavender-Bares, J., Chapin, T., Cornelissen, J. H. C., Diemer, M., Flexas, J., Garnier, E.,
903 Groom, P. K., Gulias, J., Hikosaka, K., Lamont, B. B., Lee, T., Lee, W., Lusk, C., Midgley, J.
904 J., Navas, M.-L., Niinemets, U., Oleksyn, J., Osada, N., Poorter, H., Poot, P., Prior, L.,
905 Pyankov, V. I., Roumet, C., Thomas, S. C., Tjoelker, M. G., Veneklaas, E. J., and Villar, R.:
906 The worldwide leaf economics spectrum, Nature, 428, 821–827, 2004.

907

908 Wu, H., Xiang, W., Ouyang, S., Xiao, W., Li, S., Chen, L., Lei, P., Deng, X., Zeng, Y., Zeng, 909 L., and Peng, C.: Tree growth rate and soil nutrient status determine the shift in nutrient-use 910 strategy of Chinese fir plantations along a chronosequence, For. Ecol. Manage., 460, 117896, 911 2020.





913 Xu, M., Zhu, Y., Zhang, S., Feng, Y., Zhang, W., and Han, X.: Global scaling the leaf
914 nitrogen and phosphorus resorption of woody species: Revisiting some commonly held
915 views, Sci. Total Environ., 788, 147807, 2021.
916

917 Yan, T., Zhu, J., and Yang, K.: Leaf nitrogen and phosphorus resorption of woody species in
918 response to climatic conditions and soil nutrients: a meta-analysis,
919 https://doi.org/10.1007/s11676-017-0519-z, 2018.

920

921 Yang, X., Post, W. M., Thornton, P. E., and Jain, A.: The distribution of soil phosphorus for922 global biogeochemical modeling, Biogeosciences, 10, 2525–2537, 2013.

923

924 Yuan, Z. Y. and Chen, H. Y. H.: Global-scale patterns of nutrient resorption associated with 925 latitude, temperature and precipitation, Glob. Ecol. Biogeogr., 18, 11–18, 2009.

926

927 Yuan, Z. Y. and Chen, H. Y. H.: Negative effects of fertilization on plant nutrient resorption, 928 Ecology, 96, 373–380, 2015.

929

930 Yuan, Z.-Y., Li, L.-H., Han, X.-G., Huang, J.-H., Jiang, G.-M., Wan, S.-Q., Zhang, W.-H., and
931 Chen, Q.-S.: Nitrogen resorption from senescing leaves in 28 plant species in a semi-arid
932 region of northern China, J. Arid Environ., 63, 191–202, 2005.

933

934 Zaehle, S.: Terrestrial nitrogen-carbon cycle interactions at the global scale, Philos. Trans. R.935 Soc. Lond. B Biol. Sci., 368, 20130125, 2013.

936

937 Zaehle, S., Medlyn, B. E., De Kauwe, M. G., Walker, A. P., Dietze, M. C., Hickler, T., Luo,
938 Y., Wang, Y.-P., El-Masri, B., Thornton, P., Jain, A., Wang, S., Warlind, D., Weng, E., Parton,
939 W., Iversen, C. M., Gallet-Budynek, A., McCarthy, H., Finzi, A., Hanson, P. J., Prentice, I.
940 C., Oren, R., and Norby, R. J.: Evaluation of 11 terrestrial carbon-nitrogen cycle models
941 against observations from two temperate Free-Air CO2 Enrichment studies, New Phytol.,
942 202, 803–822, 2014.

943

944 Zhang, M., Luo, Y., Meng, Q., and Han, W.: Correction of leaf nutrient resorption efficiency 945 on the mass basis, J Plant Ecol, 15, 1125–1132, 2022.

- 948
- 949
- 950
- 951
- 952
- 953

⁹⁴⁶





954 Appendix A - Sensitivity study of the importance of MLCF

955 We assembled the global dataset from the gap-filled version of TRY Plant Trait database 956 (https://www.try-db.org, Kattge et al., 2020, version 5.0) containing field measurements of 957 paired leaf and litter mass-based tissue N and P concentrations ($N_{\text{mass, leaf}}$, $P_{\text{mass, litter}}$, 958 $P_{\text{mass, litter}}$) to derive the fractional nutrient resorption (described in Methods Sect. 2.1).

959 In order to understand the importance of considering MLCF in the formula to derive reliable960 nutrient resorption values, we compared four sub datasets from the final global dataset:

961 (a) we derived nutrient resorption from nutrient resorption database, in which MLCF was962 calculated directly from leaf dry mass or leaf mass loss measurements;

963 (b) the second dataset we derived nutrient resorption from nutrient resorption database as
964 well, but we filled the missing values of MLCF using the mean for each plant functional type:
965 0.712 for deciduous, 0.766 for evergreen, 0.69 for conifers, and 0.75 for woody lianas,
966 respectively.

967 (c) the third dataset we derived nutrient resorption using leaf nutrient and litter data from968 TRY traits, in which we did not include MLCF in the formula, calculated as:

969
$$NuRE = \left(1 - \frac{Nu_{senesced}}{Nu_{green}}\right) \times 100$$
(2)

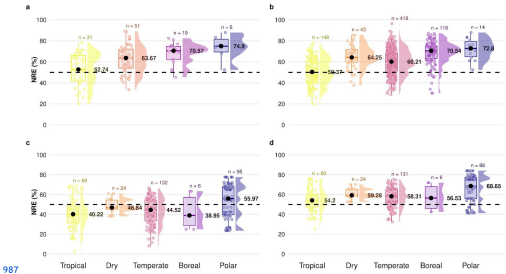
970 (d) the fourth dataset we derived nutrient resorption using leaf nutrient and litter data from 971 TRY, but here we filled MLCF with the mean per PFT calculated before, in which we 972 associated these means with leaf phenology, leaf type and growth form information. For that, 973 trees with needle evergreen leaves received conifers MLCF, deciduous trees/shrubs received 974 deciduous woody MLCF, and evergreen trees/shrubs received evergreen woody MLCF, 975 respectively.

976 Figure A1 shows nitrogen resorption efficiency (NRE) between different climate zones, 977 where we can see underestimated values of resorption only when we do not consider MLCF 978 in the formula (Fig. A1c), with values around or lower 50% of N resorption. We can see more 979 reliable resorption values around 60% when considering MLCF in the formula (Fig. A1a A1b 980 A1d). When applying the mean of MLCF for the table deriving NRE from TRY traits (Fig. 981 A1d), we could reproduce a similar pattern compared to the resorption database imported 982 from TRY (Fig. A1a). Figure A2 shows the distribution of NRE for each subset described 983 before, where we can see a clear difference in data distribution only when we do not consider





984 MLCF in the formula (Fig. A2c). For our final dataset, we then considered together the 985 dataset (b) and (d), in which are the most reliable data for nutrient resorption as it is providing 986 more data points for resorption, as well as it is considering MLCF in the formula.



988

989 Figure A1: Nitrogen resorption efficiency (NRE %) between climate zones by Köppen climate classification.
990 (a) nutrient resorption values derived directly from nutrient resorption dataset, with MLCF calculated from leaf
991 dry mass or leaf mass loss measurements; (b) nutrient resorption values derived directly from nutrient resorption
992 dataset, but with missing MLCF filled by the mean for each plant functional type; (c) nutrient resorption values
993 derived from TRY traits with no MLCF in the formula; (d) nutrient resorption values derived from TRY traits,
994 but with missing MLCF filled by the mean for each plant functional type.



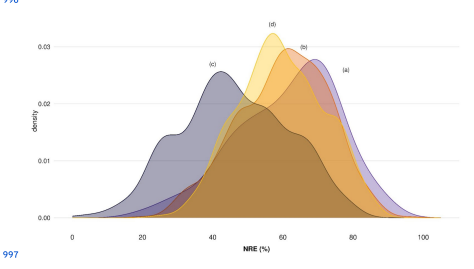






Figure A2: Distribution of Nitrogen resorption efficiency (NRE %) for all subsets: (a) nutrient resorption values derived directly from nutrient resorption dataset, with MLCF calculated from leaf dry mass or leaf mass loss measurements; (b) nutrient resorption values derived directly from nutrient resorption dataset, but with missing MLCF filled by the mean for each plant functional type; (c) nutrient resorption values derived from TRY traits with no MLCF in the formula; (d) nutrient resorption values derived from TRY traits, but with missing MLCF filled by the mean for each plant functional type.

1004

1005 Appendix B - Global patterns of nutrient resorption efficiency for N and P

1006 by PFTs and climate zones

1007 Table B1 | Summary of Nitrogen resorption efficiency (NRE; %) and Phosphorus resorption efficiency (PRE; 1008 %) in different climate zones. For each relationship, the number of observations (N), minimum (Min), maximum 1009 (Max), median, and standard deviation (SD) were reported. Letters in Significance show the statistical 1010 comparison between each climate zone.

| Resorption (%) | Climate zone | Ν | Min | Max | Median | SD | Significance |
|-------------------|--------------|-----|-------|-------|--------|-------|--------------|
| NRE | Tropical | 178 | 19.77 | 78.23 | 52.46 | 12.15 | а |
| | Dry | 65 | 37.17 | 85.48 | 61.66 | 9.72 | bc |
| | Temperate | 507 | 28.77 | 89.11 | 59.18 | 11.06 | С |
| | Boreal | 102 | 29.64 | 86.72 | 69.03 | 11.0 | b |
| | Polar | 102 | 41.42 | 87.89 | 69.62 | 12.84 | b |
| PRE | Tropical | 100 | 27.65 | 87.23 | 61.7 | 12.84 | ns |
| | Dry | 5 | 42.55 | 72.31 | 66.09 | 11.47 | ns |
| | Temperate | 273 | 29.14 | 95.11 | 57.80 | 13.65 | a |
| | Boreal | 57 | 35.92 | 88.88 | 67.36 | 13.65 | b |
| | Polar | 12 | 52.16 | 83.58 | 68.02 | 8.84 | ns |

1011

1012 Table B2 | Summary of Nitrogen resorption efficiency (NRE; %) and Phosphorus resorption efficiency (PRE; 1013 %) in different plant functional types (PFTs). For each relationship, the number of observations (N), minimum **1014** (Min) maximum (Max) median a value and standard deviation (ED) wave reported in value in distribution (ED) wave reported in the value in the v

1014 (Min), maximum (Max), median, p value and standard deviation (SD) were reported. 'p-value' < 0.05 indicates 1015 statistical significance.

| Resorption (%) | PFT | Ν | Min | Max | Median | p value | SD |
|----------------|---------------|-----|-------|-------|--------|---------|-------|
| NRE | Deciduous | 400 | 29.64 | 89.11 | 65.27 | | 12.48 |
| | Evergreens | 551 | 19.77 | 87.89 | 57.96 | <0.001 | 11.45 |
| | Broad-leaves | 841 | 19.77 | 89.11 | 59.8 | | 12.53 |
| | Needle-leaves | 103 | 40.19 | 87.89 | 61.84 | 0.05 | 9.97 |
| | Shrubs | 230 | 30.13 | 85.48 | 63.17 | | 12.48 |
| | Trees | 724 | 19.77 | 89.11 | 59.27 | <0.001 | 12.17 |
| PRE | Deciduous | 220 | 29.22 | 95.78 | 60.04 | | 12.86 |
| | Evergreens | 231 | 27.65 | 91.78 | 61.7 | 0.46 | 14.41 |





| Broad-leaves | 404 | 27.65 | 95.11 | 59.64 | | 13.50 |
|---------------|-----|-------|-------|-------|--------|-------|
| Needle-leaves | 45 | 51.35 | 88.88 | 72.2 | <0.001 | 9.23 |
| Shrubs | 59 | 32.97 | 87.23 | 64.4 | | 13.50 |
| Trees | 395 | 27.65 | 95.11 | 61.1 | 0.89 | 13.67 |

1016

1017

1018 Table B3 | Summary of Nitrogen resorption efficiency (NRE; %) and Phosphorus resorption efficiency (PRE; 1019 %) in different plant functional types (PFT) separated in different climate zones. For each relationship, the 1020 number of observations (N), minimum (Min), maximum (Max), median, and standard deviation (SD) were 1021 reported. Letters in Significance show the statistical comparison between each climate zone.
NRE

| NRE | | | | | | | | |
|---------------|---------------|-----|-------|-------|--------|-------|--------------|--|
| PFT | Climate zones | Ν | Min | Max | Median | SD | Significance | |
| Deciduous | Tropical | 31 | 31.97 | 71.80 | 52.53 | 11.64 | a | |
| | Dry | 31 | 37.17 | 85.48 | 65.95 | 11.68 | b | |
| | Temperate | 216 | 31.95 | 89.11 | 62.39 | 11.84 | cb | |
| | Boreal | 61 | 29.64 | 86.72 | 68.28 | 11.17 | db | |
| | Polar | 61 | 47.15 | 84.16 | 75.60 | 9.99 | е | |
| Evergreens | Tropical | 147 | 19.77 | 78.23 | 52.43 | 12.28 | а | |
| | Dry | 34 | 40.97 | 79.57 | 60.42 | 7.06 | bc | |
| | Temperate | 288 | 28.77 | 81.56 | 58.40 | 9.93 | cd | |
| | Boreal | 41 | 30.13 | 82.44 | 70.57 | 10.87 | b | |
| | Polar | 41 | 41.42 | 87.89 | 56.03 | 13.44 | d | |
| Broad-leaves | Tropical | 174 | 19.77 | 78.23 | 52.46 | 12.15 | а | |
| | Dry | 63 | 37.17 | 85.48 | 61.66 | 9.42 | bc | |
| | Temperate | 453 | 28.77 | 89.11 | 59.18 | 11.36 | С | |
| | Boreal | 69 | 29.64 | 86.72 | 68.28 | 12.13 | b | |
| | Polar | 82 | 41.42 | 84.16 | 75.10 | 12.34 | b | |
| Needle-leaves | Tropical | 1 | 65.25 | 65.25 | 65.25 | - | ns | |
| | Dry | 2 | 46.60 | 79.65 | 63.13 | 23.37 | ns | |
| | Temperate | 47 | 40.19 | 81.56 | 58.80 | 7.45 | а | |
| | Boreal | 33 | 51.02 | 82.44 | 71.52 | 7.33 | b | |
| | Polar | 20 | 46.76 | 87.89 | 56.03 | 11.58 | а | |
| Shrubs | Tropical | 21 | 33.81 | 74.33 | 59.60 | 11.45 | а | |
| | Dry | 33 | 37.17 | 85.48 | 63.72 | 12.08 | ns | |
| | Temperate | 77 | 31.29 | 80.96 | 59.16 | 10.63 | а | |
| | Boreal | 27 | 30.13 | 85.15 | 65.77 | 13.66 | ns | |
| | Polar | 72 | 41.42 | 84.16 | 71.16 | 11.92 | b | |
| Trees | Tropical | 157 | 19.77 | 78.23 | 52.35 | 12.18 | а | |
| | Dry | 32 | 47.10 | 76.26 | 60.08 | 6.59 | bc | |
| | Temperate | 430 | 28.77 | 89.11 | 59.18 | 11.13 | С | |
| | Boreal | 75 | 29.64 | 86.11 | 70.05 | 9.49 | b | |
| | Polar | 30 | 46.76 | 87.89 | 68.44 | 14.89 | bc | |



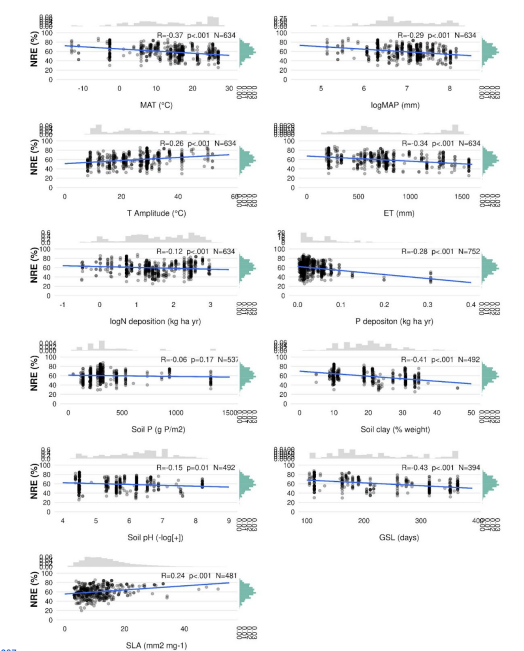


| | | | PRE | | | | |
|---------------|---------------|-----|-------|-------|--------|-------|--------------|
| PFT | Climate zones | Ν | Min | Max | Median | SD | Significance |
| Deciduous | Tropical | 25 | 35.92 | 76.26 | 64.40 | 13.14 | ns |
| | Dry | 4 | 64.40 | 72.31 | 66.29 | 3.44 | ns |
| | Temperate | 145 | 29.22 | 95.11 | 59.95 | 13.32 | ns |
| | Boreal | 33 | 35.92 | 84.33 | 59.31 | 12.18 | ns |
| | Polar | 6 | 59.31 | 71.52 | 64.51 | 4.90 | ns |
| Evergreens | Tropical | 75 | 27.65 | 87.23 | 61.70 | 12.81 | а |
| | Dry | 1 | 42.55 | 42.55 | 42.55 | - | ns |
| | Temperate | 125 | 29.14 | 91.78 | 57.44 | 13.85 | a |
| | Boreal | 24 | 61.38 | 88.88 | 79.26 | 7.58 | b |
| | Polar | 6 | 52.16 | 83.58 | 73.73 | 11.03 | ns |
| Broad-leaves | Tropical | 97 | 27.65 | 87.23 | 61.70 | 12.98 | ns |
| | Dry | 5 | 42.55 | 72.31 | 66.10 | 11.47 | ns |
| | Temperate | 249 | 29.14 | 95.11 | 57.28 | 13.93 | ns |
| | Boreal | 36 | 35.92 | 84.33 | 60.14 | 11.92 | ns |
| | Polar | 10 | 52.16 | 83.58 | 68.03 | 9.63 | ns |
| Needle-leaves | Temperate | 22 | 51.35 | 82.62 | 65.25 | 7.06 | a |
| | Boreal | 21 | 61.38 | 88.88 | 80.14 | 7.22 | b |
| | Polar | 2 | 67.02 | 73.00 | 70.01 | 4.22 | ns |
| Shrubs | Tropical | 14 | 47.85 | 79.97 | 61.95 | 10.39 | ns |
| | Dry | 3 | 42.55 | 66.09 | 64.40 | 13.13 | ns |
| | Temperate | 20 | 32.97 | 87.23 | 52.72 | 17.36 | ns |
| | Boreal | 13 | 46.60 | 82.20 | 67.17 | 10.70 | ns |
| | Polar | 9 | 52.16 | 83.58 | 71.52 | 10.0 | ns |
| Trees | Tropical | 86 | 27.65 | 87.23 | 61.70 | 13.24 | ns |
| | Dry | 2 | 66.49 | 72.31 | 69.40 | 4.11 | ns |
| | Temperate | 253 | 29.14 | 95.11 | 58.78 | 13.35 | а |
| | Boreal | 44 | 35.92 | 88.88 | 67.78 | 14.48 | b |
| | Polar | 3 | 61.11 | 68.68 | 67.03 | 3.97 | ns |



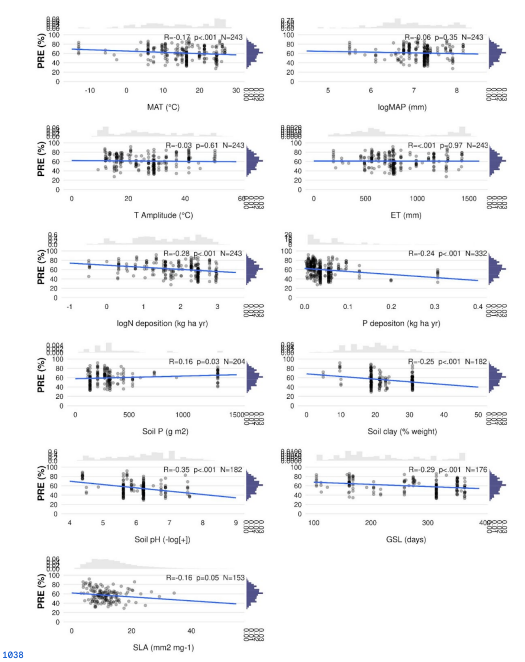


1035 Appendix C - Linear regressions of nutrient resorption with environmental **1036** and biological factors









1039 Figura C1. Linear regression of Nitrogen resorption efficiency (NRE; %) and Phosphorus resorption efficiency 1040 (PRE; %) with all possible predictor variables. Environmental predictors: Mean Annual Temperature (MAT), 1041 Mean Annual Precipitation (MAP), Evapotranspiration (ET), Temperature amplitude (T amplitude), Nitrogen 1042 deposition (N deposition), Phosphorus deposition (P deposition), total soil P (soil P) soil clay fraction (Soil

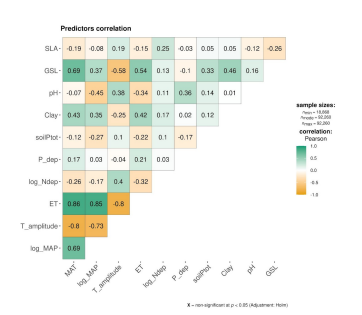
1046

1047

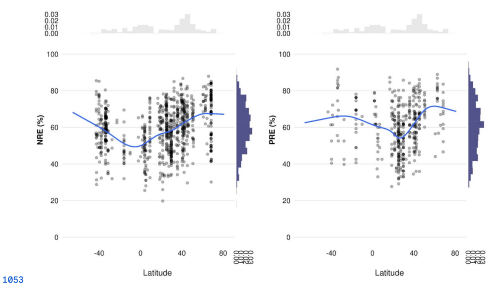




1043 Clay), soil pH. Biological predictors: Growing Season Length (GSL), Specific Leaf Area (SLA). R: Pearson 1044 correlation; p < 0.05 indicates statistical significance; N: number of observations. 1045



1048 Figure C2: Multiple Pearson correlation between all predictors. Mean Annual Temperature (MAT); Mean
1049 Annual Precipitation (MAP); Evapotranspiration (ET); Temperature amplitude (T amplitude); Nitrogen
1050 deposition (N deposition); Phosphorus deposition (P deposition); total soil P (soilPtot); soil clay fraction (Clay);
1051 soil pH; Growing Season Length (GSL); Specific Leaf Area (SLA).
1052



1054 Figure C3: Linear regression of nitrogen and phosphorus resorption efficiency (NRE %; PRE %;) with latitude.