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Elevation alters carbon and nutrient concentrations and stoichiometry in *Quercus aquifolioides* in southwestern China



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HIGHLIGHTS

- The concentration of non-structural carbohydrates (NSCs) in fine roots decreased with elevation in winter.
- Winter NSCs stored in roots transferred to aboveground tissues during winter and at the beginning of growing season.
- Both available soil P and plant tissue P concentrations decreased with increasing elevation.
- The upper limit of *Q. aquifolioides* on Balang Mountain may be co-determined by winter root NSC storage and P availability.

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GRAPHICAL ABSTRACT



ABSTRACT

Elevation is a complex environmental factor altering temperature, light, moisture and soil nutrient availability, and thus may affect plant growth and physiology. Such effects of elevation may also depend on seasons. Along an elevational gradient of the Balang Mountain, southwestern China, we sampled soil and 2-year old leaves, 2-year old shoots, stem sapwood and fine roots (diameter < 5 mm) of *Quercus aquifolioides* at 2843, 2978, 3159, 3327, 3441 and 3589 m a.s.l. in both summer and winter. In summer, the concentrations of tissue non-structural carbohydrates (NSC) did not decrease with increasing elevation, suggesting that the carbon supply is sufficient for plant growth at high altitude in the growing season. The concentration of NSC in fine roots decreased with elevation in winter, and the mean concentration of NSC across tissues in a whole plant showed no significant difference between the two sampling seasons, suggesting that the direction of NSC reallocation among plant tissues changed with season. During the growing season, NSC transferred from leaves to other tissues, and in winter NSC stored in roots transferred from roots to aboveground tissues. Available soil N increased with elevation, but total N concentrations in plant tissues did not show any clear elevational pattern. Both available soil P and total P concentrations in all plant tissues decreased with increasing elevation. Thus, tissue N:P ratio increased with

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elevation, suggesting that P may become a limiting element for plant growth at high elevation. The present study suggests that the upper limit of *Q. aquifolioides* on Balang Mountain may be co-determined by winter root NSC storage and P availability. Our results contribute to better understanding of the mechanisms for plants' upper limit formation.

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1. Introduction

Global warming is altering soil biogeochemistry in mountain systems around the world (Mayor et al., 2017), and there is increasing evidence that mountains are highly responsive to global climate change (Pauli et al., 2012). The global occurrence of mountains across all latitudes and the influential effect of declining temperature with increasing elevation, leading to the formation of treelines (Körner, 2012), offer unique opportunities to examine warming-induced phenomena in a natural context. As many environmental factors, such as temperature, precipitation, soil chemical and physical properties and growing season length, change simultaneously with elevation (White et al., 1999; Hughes, 2000; Körner, 2007), elevation can be used as a proxy for variation in biotic and abiotic characteristics to study and better understand how changes in climate may affect the growth and physiology of plants (Davis et al., 1991).

Environmental variation along elevational gradients may influence the functional and structural features of plants (Cordell et al., 1998; Peterson, 1998; Sexton et al., 2009), thus determining their growth (Li et al., 2003; Li and Yang, 2004) and distribution (Kessler, 2001; Vetaas and Grytnes, 2002; Grytnes, 2003). At an individual level, because lower temperature and shorter growing season negatively affect the carbon gain (photosynthesis), but results in a relatively greater decrease of carbon consumption or loss (e.g., respiration), the carbon balance of a plant in a year cycle may turn positive with increasing altitude (Ryan, 1991; Atkin and Tjoelker, 2003). In this sense, the carbon balance can be regarded as a proxy for plant responses to the total environment and an indicator of the cascading effects of warming temperature (Sundqvist et al., 2013). Current species distribution and function along an elevation gradient can be used to track range shifts of organisms in response to future climate change.

Individual-level carbon balance can be characterized by tissue concentrations of non-structural carbohydrates (NSC), which involves soluble sugars and starch (Mooney, 1972; Chapin et al., 1990; Li et al., 2002). Tissue NSC reflect the ratio of carbon source to sink activity (Hoch and Körner, 2003; Bansal and Germino, 2008; Li et al., 2008a). When carbon supply from photosynthesis exceeds plant's demand for growth and maintenance, a large amount of NSC will accumulate as reserves in plant tissues to support future growth (Chapin et al., 1990). By contrast, when source activity is insufficient and the carbon demand exceeds the carbon supply, little NSC will accumulate (Li et al., 2001, 2002). Nevertheless, in situ evidence for deficient carbon supply to meet growth demand of trees approaching the treeline is scarce (Bansal and Germino, 2008; Mayor et al., 2017). A large proportion (up to 70%) of carbon gained through photosynthesis can be lost through respiration, which is more sensitive to temperature than photosynthesis (Atkin et al., 2005).

Previous studies showed that, in the growing season, the NSC level in trees at higher elevation is not reduced (declined carbon supplied from photosynthesis), and sometimes even higher (restricted carbon demand due to a limited tissue formation), than that in trees at lower altitude (Körner, 2003; Hoch and Körner, 2003; Handa et al., 2005; Shi et al., 2006, 2008; Gruber et al., 2011; Sala et al., 2012; Wiley and Helliker, 2012; Dang et al., 2015). However, most of these studies compared NSC in trees only at two or three different elevations and across an altitude gradient of ~300 m, i.e., considering trees at the upper limit of their climate range and trees at one or two lower elevations (Körner, 2003; Li et al., 2008a; Fajardo and Piper, 2014). These results

detract from a broad understanding of elevational trend of NSC in plants, especially when a plant species is distributed across a wide elevational range. Furthermore, few studies have specifically considered the NSC status in the dormant season, since the alpine treeline is normally not accessible in winter due to snow cover (Hoch and Körner, 2003; Richardson et al., 2004). Thus, it is still unclear whether the elevational pattern of plant NSC in the dormant season (i.e., winter) is similar to that in the growing season (Li et al., 2008a; Sveinbjörnsson et al., 2010; Dang et al., 2015). Zhu et al. (2012a, 2012b) investigated the seasonal dynamics of tissue NSC in a treeline species across a year cycle, but this study compared only three elevations, which makes the altitudinal pattern of NSC levels difficult to interpret. Therefore, our capacity to predict how NSC status of trees changes with elevation is still limited, and additional research is needed to clarify whether the carbon balance of trees changes (increase or decrease) linearly or non-linearly with increasing elevation.

The concentrations of NSC in different plant tissues can respond to elevation differently (Li et al., 2008a, 2008b; Shi et al., 2006; Zhu et al., 2012a, 2012b), but the underlying driving factor for such differences is still unclear. Because low temperatures at high altitudes are likely to limit biogeochemical processes, soil organic carbon concentration commonly increases and the availability of soil nutrients decreases with increasing elevation (Sundqvist et al., 2013; He et al., 2016). Nitrogen (N) and phosphorus (P) are key nutrients for the photosynthesis process, which can directly affect the concentration of NSC in leaves, as well as in other tissues (Li et al., 2008b; Reich et al., 2008). Elser et al. (2003) found that different species at high elevation showed different responses to N and P concentrations in tissues. The relative importance of P versus N in limiting plant growth may vary with altitude due to differences in a range of factors, including N deposition, parent material, and vegetation type (Bowman and Hurry, 1993; Shaver and Chapin, 1995). Although N is commonly considered the main limiting nutrient for plant growth in temperate and cold environments (Vitousek and Howarth, 1991), the role of P limitation (and co-limitation of N and P) is increasingly recognized as of importance (Attiwill and Adams, 1993; Zotz, 2004; Reich et al., 2008). Although an increasing amount of evidence suggests that N versus P limitation varies among plant communities, relatively little is known about how the availability and the relative importance of plant N and P will be influenced by elevation in mountain systems, and whether elevational patterns of nutrients differ between plant tissues.

We sampled a common evergreen broad-leaved tree *Quercus aquifolioides* at six elevations in both summer and winter to examine the elevational trends of NSC, N, and P in different plant tissues and their relationships across seasons. Our hypotheses were (1) that tissue NSC concentration of trees changes with elevation, (2) that this elevational pattern is related to season associated with temperature, and (3) that this pattern is correlated with the elevational patterns of plant N and P status, because plant N and P are two key elements for the photosynthesis process that can directly affect the tissue NSC concentration and NSC balance (Li et al., 2008); Reich et al., 2008).

2. Materials and methods

2.1. Study site and climate conditions

The study was carried out on the sunny slope of the Balang Mt. in the Wolong Natural Reserve (102°57′ E, 30°53′ N), located at the eastern

edge of Qinghai-Tibetan Plateau in southwestern China. The study area has a subtropical inland mountain climate with sunny, dry winter (November to April) and warm, humid summer (June to September). According to long-term climate data collected at 1920 m a.s.l. (Wolong Nature Reserve Authority), the annual mean precipitation is 995 mm, with rainfall mainly occurring from May to September, and the annual mean temperature is 12.8 °C, with the monthly mean temperature of 17.0 °C in July and -1.7 °C in January. Soil on the study slope is mountain brown soil with a depth of <50 cm.

Along the elevational gradient on the sunny slope of the Balang Mt., three climate stations were established at 2800, 3200 and 3500 m a.s.l., respectively. Climate data were collected for 2012 and 2013.

2.2. Study species and sampling protocol

Quercus aquifolioides occupies a wide range of habitats and occurs at altitudes from 2000 to 4500 m a.s.l. (Zhou, 1992; Zhu et al., 2012a, 2012b). It is a dominant, late-successional and climax species of the sclerophyllous evergreen broad-leaved forests on sunny, south-facing slopes in the region of the Hengduan Mountains (93°18′–104°43′ E, 26°33′–31°55′ N), southwestern China. On the sunny (south) slope of the Balang Mt. pure *Q. aquifolioides* stand ranged from 2800 m to 3600 m a.s.l. (upper limit). This naturally generated shrub stand is 30–35 years old, and comprises multi-stemmed clumps. Since the nature reserve was established in 1976, the *Q. aquifolioides* stands had not been disturbed.

We randomly selected six plots $(5-10 \text{ m} \times 10 \text{ m})$ within the *Q. aquifolioides* stands at each of six elevations along the elevational gradient, i.e., 2843, 2978, 3159, 3327, 3441, and 3589 (± 5) m a.s.l. The distance between any two plots at the same elevation was at least 50 m apart. We recorded the environmental conditions of each plot, and measured the mean height (H), mean diameter at breast height (DBH) and number of main stems of *Q. aquifolioides* clumps in each plot. Growth characteristics of *Q. aquifolioides* are summarized in Table 1. Samples were taken on 18–19 July 2014 (hereafter referred to as summer, i.e., at the time of peak growth) and on 6–7 November 2014 (winter, at the dormant season), when the plots at the lowest elevation (2843 m) were already covered by a thin layer of snow.

In each plot, four types of Q. aquifolioides tissues (2-year old leaves, 2-year old shoots, stem sapwood, and fine roots with a diameter of <0.5 cm) were sampled from >5 randomly selected individuals on each of the two sampling dates (summer, winter). Samples of leaves and shoots were taken from non-shaded branches on the upslope side of plants, between 10:30 and 16:00, when all plots received direct sunshine (Graham et al., 2003). To collect fine roots, we first found coarse roots originating from each selected Q. aquifolioides plant, and then fine roots (<0.5 cm in diameter) attached to those coarse roots were manually excavated using mini-spade, mini-pick, and trowel. Only fine roots that were located within the 0-20 cm soil layer around the stump (<50 cm) and covered by soil were collected, i.e., we did not collect any deep roots from parent materials (soil depth was <20 cm on the study slope) and roots exposed to air or sunlight. Leaves, shoots and fine roots collected from all trees within a plot were pooled and mixed to get a sample for each tissue for each plot (n = 6 plots), respectively. All

Table 1

Growth (mean \pm 1SE, n = 6) of the sampling trees of *Quercus aquifoliodes* along the elevation gradient on the sunny slope of the Balang Mt.

-				
	Elevation (m a.s.l.)	No. of stems per clump	Mean height (m)	Mean diameter at breast height (cm)
1	3589	2.3 ± 1.2	1.8 ± 0.6	5.5 ± 1.0
2	3441	7.5 ± 2.9	2.8 ± 0.5	7.4 ± 1.0
3	3327	7.4 ± 2.9	2.8 ± 0.9	5.1 ± 1.7
4	3159	6.1 ± 4.5	2.8 ± 0.9	5.6 ± 2.0
5	2978	3.5 ± 1.6	3.1 ± 0.7	7.1 ± 2.8
6	2843	7.0 ± 1.8	3.7 ± 0.3	8.2 ± 2.0

samples were immediately stored in a cool box in the field, killed in a microwave oven in the evening (40 s at 600 W), and dried to a constant weight at 65 °C in Wolong Forest Ecosystem Research Station located at 2800 m a.s.l., and kept dry until laboratory treatments.

On 18–19 July 2014, mineral soils (0–10 cm depth) were also taken, after removing soil organic matter, from 4 to 6 locations in each plot and then mixed homogeneously to get a composite soil sample for each plot (n = 6 plots). Soils were taken from the 0–10 cm layer only because soil depth was <20 cm on that slope and the majority of fine root biomass in forest stands occurs in the top 0–20 cm soil layer (Jackson et al., 1996; Meinen et al., 2009). For each soil sample, a subsample was restored in 4 °C to test the quick-acting N, and the other subsample was dried for the measurement of P and other elements.

2.3. Non-structural carbohydrate (NSC) analysis

Dried plant material was shattered through a ball mill instrument. The powdered material (0.1 g) was put into a 15 mL centrifuge tube, where 5 mL of 80% ethanol was added. The mixture was incubated at 80 °C in a water bath shaker for 30 min, and then centrifuged at 4000 rpm for 5 min. The pellets were extracted two more times with 80% ethanol. Supernatants were retained for soluble sugar determinations, and the ethanol-insoluble pellet was used for starch extraction. Concentrations of NSC for a sample was defined as soluble sugar concentration plus starch concentration.

Soluble sugars in the supernatants were determined using the anthrone method (Seifter et al., 1950). An aliquot of the extract was hydrolysed in 5 mL of 0.4% anthrone solution (4 g anthrone in 1000 mL 95% H_2SO_4) in a boiling water bath for 10 min. After cooling, the sugar concentration was determined spectrophotometrically (ultraviolet-visible spectrophotometer 752S; Cany Precision Instruments Co., Ltd., Shanghai, China) at 620 nm. Glucose was used as a standard. The sugar concentration was calculated on a dry mass basis (% d.m.) and also on a leaf area basis (g/m², for leaves only).

Ethanol in the ethanol-insoluble pellet was removed by evaporation. Starch in the residue was released in 2 mL distilled water for 15 min in a boiling water bath. After cooling to room temperature, 2 mL of 9.2 mol L^{-1} HClO₄ was added. Starch was hydrolysed for 15 min. Distilled water (4 mL) was added to the samples. The samples were then centrifuged at 4000 rpm for 10 min. The pellets were extracted one more time with 2 mL of 4.6 mol L^{-1} HClO₄. Supernatants were retained, combined and filled to 25 mL, to measure the glucose concentration spectrophotometrically (ultraviolet-visible spectrophotometer 752S) at 620 nm using anthrone reagent, and the starch concentration was then calculated by multiplying glucose concentration by the conversion factor of 0.9 (Osaki et al., 1991). The starch concentration was described on a dry mass basis (% d.m.) and also on a leaf area basis (g/m², for leaves only).

2.4. Nitrogen and phosphorus analysis

Concentrations of total N of tissues were analyzed using a C/N analyzer (Vario Micro cube, Germany). Concentrations of total P were determined following ammonium molybdate method after nitric acid and perchloric acid (2:1) digestion (Sparks et al., 1996).

The availability of soil N (nitrate and ammonium) was determined colorimetrically from 1 M KCL soil extracts from fresh soil samples using an Auto Continuous Flow Analyzer (Bran & Luebbe, Norderstedt, Germany). The availability of soil P was determined by the molybde-num blue colorimetric method (Murphy and Riley, 1962) after extraction by 0.5 M NaHCO₃ (Olsen, 1954).

2.5. Statistical analysis

Data were tested for normality and homogeneity of variance, and transformed to logarithm when normality and homogeneity of variance

Table 2

The effects of elevation, sampling season (summer and winter), tissue type, and their interactions on concentrations of non-structural carbohydrates (NSC), total nitrogen (N), total phosphorus (P), ratio of NSC to N (NSC:N), and ratio of N to P (N:P) in *Quercus aquifoliodes* plants, tested with three-way ANOVAs.

Effects	df	NSC	Ν	Р	NSC:N	N:P
Season (S) Tissue (T) Elevation (E) $S \times T$ $T \times E$ $S \times E$ $S \times T \times E$	1, 240 3, 240 5, 240 3, 240 15, 240 5, 240 15, 240	3.09 ^{ns} 34.42*** 5.21*** 75.28*** 4.25*** 10.77*** 4.39***	74.88*** 1160.77*** 26.33*** 11.33*** 2.93*** 3.73*** 1.52 ^{ns}	109.32*** 104.49*** 20.46*** 31.24*** 10.32*** 7.19*** 2.71***	0.01 ^{ns} 114.87*** 8.401*** 11.05*** 3.01*** 4.42*** 1.31 ^{ns}	18.37*** 90.05*** 9.045*** 47.69** 2.89*** 3.92*** 2.72***

F values are given. Note, degree of freedom (df) and significance levels (^{ns} P > 0.05, *P < 0.05, *P < 0.01, and ***P < 0.001).

could not be met. First, we tested the effects of sampling season (summer, winter), tissue type (4 types), elevation (6 levels), and their interactions on tissue NSC, N, P, NSC:N, and N:P, and found that tissue type interacted with elevation to affect the parameters studied (Table 2). We, therefore, analyzed the effects of sampling date and elevation on each parameter for each tissue type separately (Table 3). To explore the dynamic pattern along the elevational gradient, we tested the relationship between physiological indexes (NSC, N, P, NSC:N, and N:P) and altitude with regression analysis (Figs. 1–6), as well as the correlation analysis between soil and each tissue (Table 4). All the statistical analyses were done using SPSS (v. 20.0, SPSS Inc., Chicago, IL, USA).

3. Results

3.1. Climatic factors and soil N and P along the elevational gradient

The climatic pattern along the elevational gradient showed almost the same trend for the two years, and annual and growing season patterns differed only in the numeric value (Fig. 1). Air temperature (Fig. 1A, B) decreased linearly with elevation, but soil temperature did not (Fig. 1C, D). Temperature lapse rate was 0.46 °C per 100 m. Growing season air humidity (Fig. 1H) increased linearly with elevation, but annual air humidity did not (Fig. 1G). The precipitation was significantly less at lower elevations than at the highest elevation (Fig. 1E, F). Available soil N increased linearly with increasing elevation, but available soil P decreased linearly with elevation (Fig. 1I, J).

3.2. Plant tissue N and P along the elevational gradient

Plant tissue N and P concentration varied significantly with sampling season, tissue type, and elevation (Table 2). Total N and P level in different plant tissues changed significantly with sampling season, except for N in shoots (Table 3). In summer, tissue N concentration did not show any clear elevational trends (Fig. 2A, C, E, G). In winter, N concentration

in leaves and fine roots decreased linearly with increasing elevation (Fig. 2B, H). In summer, P level in all plant tissues decreased linearly with increasing elevation (Fig. 3A, C, E, G). In winter, P levels in leaves and fine roots decreased linearly with increasing elevation (Fig. 3B, H).

3.3. Plant tissue NSC along the elevational gradient

There was no season effect on tissue NSC concentrations (Table 2), but NSC levels varied significantly with both sampling season and tissue type (Tables 2 and 3). In summer, the NSC concentrations in leaves increased significantly linearly with increasing elevation (Fig. 4A), but NSC levels in other tissues did not show any elevational trends (Fig. 4C, E, G). In winter, the NSC concentrations in fine roots significantly decreased with increasing elevation (Fig. 4H).

3.4. Plant tissue NSC:N and N:P along the elevational gradient

Plant NSC:N and N:P ratio were significantly affected by interactions between sampling season and tissue type, as well as interactions between elevation and tissue type (Table 2). Except for fine roots (Fig. 5G), tissue NSC:N increased linearly with elevation in summer (Fig. 5A, C, E). In winter, except for a linear increase in NSC:N in leaves (Fig. 5B), tissue NSC:N did not have clear trends with elevation (Fig. 5D, F, H). Except for shoots (Fig. 6C), tissue N:P increased linearly with elevation in summer (Fig. 6A, E, G). In winter, except for a linear increase in N:P in roots (Fig. 5H), tissue N:P did not have clear trends with elevation (Fig. 6B, D, F).

4. Discussion

4.1. Elevational effects on NSC and nutrients in different tissues

In summer (growing season), NSC in any tissues of *Q. aquifolioides* did not decrease with elevation (Fig. 4), which is consistent with previous studies on treeline species (Hoch and Körner, 2005; Shi et al., 2006; Li et al., 2008a, 2008b; Zhu et al., 2012a, 2012b; Yu et al., 2014; Fajardo and Piper, 2014; Dang et al., 2015). This result suggests that the carbon supply was sufficient for *Q. aquifolioides* during growing season in the study site. However, low temperature and relative shorter growing season at high elevation reduced the metabolic processes needed for wood production and tissue maintenance, which led to a linear increase of NSC levels in leaves in summer (Richardson and Berlyn, 2002; Piper et al., 2006). A delay in bud break can also be related to the increase in NSC levels in leaves with increasing elevation (Gruber et al., 2011).

More interestingly, we revealed a linear decrease in NSC concentrations in fine roots with increasing elevation in winter, which is consistent with some previous studies conducted in winter (Li et al., 2008a, 2008b; Genet et al., 2010; Zhu et al., 2012a; Dang et al., 2015). The linear decreasing pattern in sink tissues (especially roots) at upper altitude at

Table 3

The effects of elevation, sampling season (summer and winter), and their interaction on concentrations of non-structural carbohydrates (NSC), total nitrogen (N), total phosphorus (P), ratio of NSC to N (NSC:N), and ratio of N to P (N:P) in *Quercus aquifoliodes* plants, tested with two-way ANOVAs.

Tissue	Effect	df	NSC	Ν	Р	NSC:N	N:P
Leaves	Season (S)	1,60	311.81***	63.04***	24.35***	313.79***	52.65***
	Elevation (E)	5,60	5.28***	11.37***	3.93**	7.53**	1.05 ^{ns}
	$S \times E$	5,60	3.39**	1.30 ^{ns}	0.92 ^{ns}	1.72 ^{ns}	1.16 ^{ns}
Shoots	Season (S)	1,60	0.18 ^{ns}	1.08 ^{ns}	36.27***	0.09 ^{ns}	27.70***
	Elevation (E)	5,60	2.34**	6.62***	1.83 ^{ns}	4.89***	7.17***
	S×E	5,60	1.27 ^{ns}	2.21 ^{ns}	3.04*	1.55 ^{ns}	4.95***
Stems	Season (S)	1,60	52.15***	9.96**	52.11***	7.72**	47.66***
	Elevation (E)	5,60	2.32 ^{ns}	3.46**	5.61***	3.32*	6.60***
	$S \times E$	5,60	1.02 ^{ns}	0.98 ^{ns}	8.32***	0.53 ^{ns}	4.72***
Fine roots	Season (S)	1,60	69.90***	52.59***	39.26***	0.05 ^{ns}	32.17***
	Elevation (E)	5,60	5.64***	4.79***	14.67***	4.99***	14.47***
	S × E	5,60	8.74***	0.55 ^{ns}	3.24**	3.77**	4.79***

F values are given. Note, degree of freedom (df) and significance levels ($^{ns} P > 0.05$, $^*P < 0.05$, $^{**}P < 0.01$, and $^{***}P < 0.01$).



Fig. 1. Annual average and growing season meteorological conditions in relation to elevation in 2012 and 2013 (A–H) and soil available nutrient (I and J) conditions in growing season in 2014. P₁ means value of 2012; P₂ means value of 2013.

the end of the growing season may influence winter survival and early spring re-growth, playing an important role in the persistence and development of *Q. aquifoliodes* at its upper distributional elevations. The

root NSC concentrations decreased significantly with increasing elevation in winter, this phenomenon, indeed, did not imply fully depletion of mobile carbohydrates in plants growing at their elevational limit



Fig. 2. Relationships between elevation and concentration of total nitrogen in different plant tissues (A–H) in two sampling seasons.

(Fig. 4H). Moreover, increasing root biomass with increasing elevation (Leuschner et al., 2007; Graefe et al., 2008; Moser et al., 2011; Zhu et al., 2012a, 2012b) may compensate for the decreasing NSC concentration when considering the root NSC pool size.

We found that soil available N increased significantly with increasing elevation (Fig. 11), which may be a result of decreasing mineralization (Kitayama et al., 1998; Soethe et al., 2008) and increasing organic matter (Dieleman et al., 2013; Du et al., 2014) and thus increasing organic N content in soils with altitude. In general, N cycling rates (e.g., nitrification and denitrification) tend to be more active at lower elevations than higher elevations in subtropical and tropical mountain forests (Vitousek, 1994; Pendry and Proctor, 1996; Leuschner et al., 2007), which may lead to decreasing N availability with increasing elevation (Sjögersten and Wookey, 2005), and thus, alpine ecosystems are often thought to be N limited (Vitousek and Howarth, 1991). In our study, high frequency of thunderstorm and snowfall may provide sufficient available N in soils at high elevations (Fig. 11). However, plant tissue N concentrations (Fig. 2A–H) did not positively respond to increases in soil N availability with increasing elevation (Fig. 11). At higher elevations, lower temperature and shorter growing season may restrict plants' N uptake and use (Vitousek, 1994; Liptzin et al., 2013; Sundqvist et al., 2014), leading to a decoupling between soil N availability (Fig. 11) and plant tissue N (Fig. 2A–H) concentrations (Table 4). Some studies described an increase in N accumulation and conservation with increasing elevation as adaptive responses to low temperature in alpine ecosystems, to enhance or maintain the metabolic capacity of plants in cold environment (Richardson et al., 2001; Shi et al., 2006). Our result showed a stable pattern of N in different tissues along the elevational gradient in summer, suggesting that N resource does not limit *Q. aquifolioides* growth at high elevations.



Fig. 3. Relationships between elevation and concentration of total phosphorus in different plant tissues (A-H) in two sampling seasons.

We found a decreasing tendency for both plant tissue P (Fig. 3) and available soil P (Fig. 1J) with increasing elevation. Consequently, there was a strong and positive correlation between plant tissue P and soil available P concentration (Table 4), suggesting that P is more limited with increasing elevation. Some studies have also provided evidence for a change in the relative importance of P versus N limitation with increasing elevation (Van de Weg et al., 2009). Low P availability in soils may directly restrict nutrient absorption and utilization, and indirectly affect P-associated physiological processes in plants, which further lead to a general decrease in total P levels in different tissues at high elevations (Fig. 3). However, Chen et al. (2013) measured leaf N and P concentrations of 386 woody species in 14 forest sites across eastern China, and found that both leaf N and P concentrations were negatively correlated with mean annual temperature but positively correlated with soil N and P contents. Other studies showed that leaf N and P concentrations first increased and then decreased with increasing elevation (Van de Weg et al., 2009; Fisher et al., 2013). Previous studies demonstrated that changes in leaf N and P concentrations reflected environmental conditions more than plant intrinsic characteristics, such as genotype (Ågren and Weih, 2012) and taxonomy (Zhang et al., 2012).

Plant N and P contents often influence each other during the growing season, especially in leaves (Kang et al., 2011; Yuan and Chen, 2012). Our results showed an increasing pattern of N:P in leaves in summer, which is consistent with previous studies (Yuan and Chen, 2012; Fisher et al., 2013). Several studies found that tissue N:P ratio increased with increasing elevation (Kang et al., 2011; Yuan and Chen, 2012; Chen et al., 2013; Zhao et al., 2016). However, based on data gained from 386 woody species across eastern China, Chen et al. (2013) reported that growing season leaf N:P ratio was positively correlated with mean annual temperature, i.e. N:P ratio decreases with increasing elevation.



Fig. 4. Relationships between elevation and concentration of non-structural carbohydrates in different tissues (A-H) in two sampling seasons.

The N:P ratio was found to be much higher in leaves than in woods (Fig. 6), Yuan and Chen (2012) combined data from 211 published studies and showed that leaves and roots had similar N:P ratios but the former had significantly higher level of N and P concentration than the latter. Decreasing elevational trend of soil P (Fig. 1J) in combination with increasing elevational trend of soil N (Fig. 1I) and plant tissue N:P (Fig. 6) suggest a P limitation at high elevations.

Increasing P limitation can be coupled with higher N deposition and redistribution of N accumulated in snow via melt-water with increasing elevation (Weintraub, 2011), if leaching is minor. This occurs in soils

dominated by low temperatures, which generally show low rates of organic matter decomposition, though N mineralization rates need to account for N demand by plants (Ruess et al., 1996). While a decline in nutrient mineralization and N availability has been associated with an increase in elevation in some studies (Sveinbjörnsson et al., 1995), others have revealed contrasting results (Griffiths et al., 2009). Increasing elevation may have contrasting effects on nutrient cycling processes, and it is not always associated with a decline in N mineralization and N availability. It is possible that higher N and lower P mineralization at higher elevation reflects decreasing microbial N demand and P



Fig. 5. Relationships between elevation and ratio of non-structural carbohydrates to concentration of total nitrogen in tissues (A–H) in two sampling seasons.

immobilization with increasing elevation (Rinnan et al., 2007; Nadelhoffer et al., 1991).

4.2. Seasonal variation of NSC and nutrients in different tissues

Seasonal variation of NSC indicated a season-dependent carbon balance between carbon acquisition (photosynthesis) and carbon investment (growth and respiration), which is consistent with previous studies (Shibata and Nishida, 1993; Palacio et al., 2008; Zhu et al., 2012a, 2012b). In summer, tissue NSC concentration did not decrease with increasing elevation (Fig. 4A, C, E, G), whereas in winter, root NSC concentration significantly linearly decreased with elevation (Fig. 4H). NSC may transfer among different tissues depending on the relative activity between tissues or between sources and sinks (Finn and Brun, 1982), or at the expenses of growth during the growing season to guarantee the survival of plants in winter, because high tissue NSC concentrations help to avoid intra- and intercellular ice formation and thus freezing damage. Moreover, tissue NSC levels may also be either passively (source-to-sink flow) or actively (gene expression) controlled (Wiley and Helliker, 2012). For example, when irradiance or source activity is insufficient, biomass of stems and roots decreases, NSC can be transferred from roots to leaves (Lee et al., 2007). Variations of environmental conditions at higher elevations may, therefore, cause resource remobilization among different tissues, which is also a strategy of plants to face the harsh habitat (Gaucher et al., 2005; Kilpeläinen et al., 2005) such as low temperature.

Available soil N increased with increasing elevation (Fig. 1I), indicating that soil N resources were sufficient to support *Q. aquifolioides* growth along the elevational gradient, even at the upper limit. A worldwide comparison revealed that trees at their upper limits did not have any disadvantages in N supply compared to plants at lower elevations (Körner, 1989). Unlike N in soil and plant tissues, both soil available P (Fig. 1J) and plant tissue P (both in summer and winter) decreased with increasing elevation (Fig. 3A–H), which may, as mentioned



Fig. 6. Relationships between elevation and ratio of total nitrogen to total phosphorus in tissues (A-H) in two sampling seasons.

 Table 4

 Correlations of concentrations of NSC, total N and total P of each tissue with soil available N and P in summer.

	NSC	Ν	Р	NSC	Ν	Р
Soil N Soil P	Leaves 0.62 ^{**} - 0.13 ^{ns}	-0.09 ^{ns} 0.29 ^{ns}	-0.38^{*} 0.34 *	Shoots 0.21^{ns} -0.05^{ns}	0.13 ^{ns} 0.11 ^{ns}	-0.32^{ns} 0.14^{ns}
Soil N Soil P	Stems 0.29 ^{ns} 0.17 ^{ns}	-0.06 ^{ns} 0.16 ^{ns}	-0.48^{**} 0.45^{**}	Roots - 0.07 ^{ns} - 0.08 ^{ns}	-0.11^{ns} -0.05^{ns}	-0.60 ^{**} 0.53 ^{**}

The given are F values and significance levels (^{ns} P > 0.05, *P < 0.05, **P < 0.01, and ***P < 0.001).

above, imply a P-limitation for plant growth at high elevations. Foliar concentration of P, in organic (i.e., nuclear acid) and inorganic (i.e., orthophosphate) forms (Chapin and Kedrowski, 1983), often decreases as altitude increases (Sundkvist et al., 2013). The inorganic P forms comprise a considerable proportion (Chapin and Kedrowski, 1983; Sterner and Elser, 2002), mainly existing as storage materials from accumulation, and reflecting the surrounding habitat (Sterner and Elser, 2002; Oyarzabal and Oesterheld, 2009). In summer and winter, total P concentrations in different tissues showed a decrease along the elevational gradient, correlating positively with available soil P. This pattern suggests a P limited habitat along the elevational gradient in the study area, especially in winter. Declining P availability, but highly variable or increasing availability of soil N, associated with increasing elevation, suggests that increasing temperature with global warming may lead to a decrease in the relative importance of P versus N limitation for *Q. aquifolioides*.

NSC:N in shoots and fine roots was similar between winter and summer, suggesting that the pattern of carbon and N balance was relatively stable (Figs. 2 and 4). Carbon and N levels in plants generally reflect the difference between uptake (photosynthesis; source activity) and demand (metabolism, growth and export; sink activity) (Körner, 2003). Increasing NSC:N with altitude indicated that carbon source did not limit tree growth and development along this elevational gradient. Other factors can be more limiting for plant growth and distribution in this study area. A linear increase in the NSC:N with altitude may indicate relative increases in N limitation with altitude (He et al., 2006). In the current study, however, variation in NSC:N was caused more by increasing NSC levels rather than by decreasing N concentrations. Nevertheless, nutritional constraints at high elevation may still hamper the conversion of carbohydrates into N- and/or P-based compounds (e.g., amino acids, proteins), as well as the transport of these compounds from leaves to other plant organs (Körner, 2003).

The ratio of N to P can be regulated by soil nutrient availability, tree growth rates and plant needs (Tessier and Raynal, 2003; Elser et al., 2003; Reich and Oleksyn, 2004; Hogan et al., 2010) and, as such, is regarded as an important index to explain nutrient limitation pattern (Wardle et al., 2004; He et al., 2006). A review of 40 fertilization studies revealed that an N:P > 16 indicated P limitation, while an N:P < 14 is indicative of N limitation. At N:P between 14 and 16, either N or P can be limiting or plant growth is co-limited by N and P together (Koerselman and Meuleman, 1996). In summer, N:P in leaves increased significantly with elevation, and much more than in other tissues. N:P was higher than 16 in leaves but lower than 14 in other tissues in summer. In both summer and winter, we found a tendency for increasing N:P with altitude in all tissues. We suggest that P limitation plays a more important role than N limitation on Q. aquifolioides growth at higher elevations. Zhao et al. (2014) found that tree leaf N:P ratios increased, while leaf N and P concentrations decreased with elevation (500-2300 m a.s.l.) in northeastern China. Conversely, N:P ratios decreased significantly as elevation increased, especially at the transition from krummholz to the alpine tundra in a Himalayan treeeline ecotone (Müller et al., 2017). He et al. (2016) found that plant and soil nutrient properties did not change linearly with elevation from 50 to 950 m a.s.l. in subtropical China.

5. Conclusions

Our results fully supported our 1st and 2nd hypotheses but are only partly in line with our 3rd hypothesis (see Introduction). Plant NSC levels depend on tissue type, elevation, season, and tissue P, but it was not correlated with tissue N. This study showed that the availability of soil N significantly increased with increasing elevation, probably because of the slowing organic matter cycling under low temperature and N deposition from snow at high elevations. Conversely, the availability of soil P decreased progressively with increasing elevation, which implies increasing P limitation with increasing elevation. Soil nutrient availability influenced by climate (temperature and precipitation), soil (leaching and weathering) and biotic factors (litter quality and microbial activity), affects tissue nutrient and carbohydrate concentrations. However, regardless of the relative changes in N and P limitation, and carbohydrate availability, tissue stoichiometry (NSC:N, N:P) revealed that transfer of NSC among different organs may occur in current conditions and that a decrease in the relative importance of P versus N limitation can be expected with a warming climate. In summer, carbon supply is sufficient for Q. aquifolioides growth and development at its upper limit, but P may be or may become a limiting factor for plant growth. Stoichiometric ratios and relationships suggest that the upper limit of this species likely depends on the nutrient source (namely P), on one hand, and on the root winter NSC storage, on the other hand.

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References

- Ågren, G.I., Weih, M., 2012. Plant stoichiometry at different scales: element concentration patterns reflect environment more than genotype. New Phytol. 194, 944–952.
- Atkin, O.K., Tjoelker, M.G., 2003. Thermal acclimation and the dynamic response of plant respiration to temperature. Trends Plant Sci. 8, 343–351.
- Atkin, O., Bruhn, D., Hurry, V., Tjoelker, M., 2005. The hot and the cold: unraveling the variable response of plant respiration. Funct. Plant Biol. 32, 87–105.
- Attiwill, P.M., Adams, M.A., 1993. Nutrient cycling in forests. New Phytol. 124, 561–582. Bansal, S., Germino, M.J., 2008. Carbon balance of conifer seedlings at timberline: relative changes in uptake, storage, and utilization. Oecologia 158, 217–227.
- Bowman, E.H., Hurry, D., 1993. Strategy through the option lens: an integrated view of resource investments and the incremental-choice process. Acad. Manag. Rev. 8, 760–782.
- Chapin III, F.S., Kedrowski, R.A., 1983. Seasonal changes in nitrogen and phosphorus fractions and autumn retranslocation in evergreen and deciduous taiga trees. Ecology 64, 376–391.
- Chapin III, F.S., Schulze, E.D., Mooney, H.A., 1990. The ecology and economics of storage in plants. Annu. Rev. Ecol. Syst. 21, 423–447.
- Chen, Y., Han, W., Tang, L., Tang, Z., Fang, J., 2013. Leaf nitrogen and phosphorus concentrations of woody plants differ in responses to climate, soil and plant growth form. Ecography 36, 178–184.
- Cordell, S., Goldstein, G., Mueller-Dombois, D., Webb, D., Vitousek, P.M., 1998. Physiological and morphological variation in *Metrosideros polymorpha*, a dominant Hawaiian tree species, along an altitudinal gradient: the role of phenotypic plasticity. Oecologia 113, 188–196.
- Dang, H.S., Zhang, R., Zhang, Q.F., Xu, Y.M., 2015. Temporal variations of mobile carbohydrates in *Abies fargesii* at the upper tree limits. Plant Biol. 17, 106–113.
- Davis, J., Schober, A., Bahn, M., Sveinbjörnsson, B., 1991. Soil carbon and nitrogen turnover at and below the elevational treeline in northern Fennoscandia. Arct. Alp. Res. 23, 279–286.
- Dieleman, W.I.J., Venter, M., Ramachandra, A., Krockenberger, A.K., Bird, M.I., 2013. Soil carbon stocks vary predictably with altitude in tropical forests: implications for soil carbon storage. Geoderma 204-205, 59–67.
- Du, B., Kang, H., Pumpanen, J., Zhu, P., Yin, S., Zou, Q., Wang, Z., Kong, F., Liu, C., 2014. Soil organic carbon stock and chemical composition along an altitude gradient in the Lushan Mountain, subtropical China. Ecol. Res. 29, 433–439.
- Elser, J.J., Acharya, K., Kyle, M., Cotner, J., Makino, W., Markow, T., Watts, T., Hobbie, S., Fagan, W., Schade, J., Hood, J., Sterner, R.W., 2003. Growth rate-stoichiometry couplings in diverse biota. Ecol. Lett. 6, 936–943.
- Fajardo, A., Piper, F.I., 2014. An experimental approach to explain the southern Andes elevational treeline. Am. J. Bot. 101, 788–795.
- Finn, G.A., Brun, W.A., 1982. Effect of atmospheric CO₂ enrichment on growth, nonstructural carbohydrate content, and root nodule activity in soybean. Plant Physiol. 69, 327–331.
- Fisher, J.B., Malhi, Y., Torres, I.C., Metcalfe, D.B., Van de Weg, M.J., Meir, P., Silva-Espejo, J.E., Huasco, W.H., 2013. Nutrient limitation in rainforests and cloud forests along a 3,000m elevation gradient in the Peruvian Andes. Oecologia 172, 889–902.
- Gaucher, C., Gougeon, S., Mauffette, Y., Messier, C., 2005. Seasonal variation in biomass and carbohydrate partitioning of understory sugar maple (*Acer saccharum*) and yellow birch (*Betula alleghaniensis*) seedlings. Tree Physiol. 25, 93–100.
- Genet, M., Li, M., Luo, T., Fourcaud, T., Clément-Vidal, A., Stokes, A., 2010. Linking carbon supply to root cell-wall chemistry and mechanics at high altitudes in *Abies georgei*. Ann. Bot. 107, 311–320.
- Graefe, S., Hertel, D., Leuschner, C., 2008. Fine root dynamics along a 2,000-m elevation transect in South Ecuadorian mountain rainforests. Plant Soil 313, 155–166.
- Graham, E.A., Mulkey, S.S., Kitajima, K., Phillips, N.G., Wright, S.J., 2003. Cloud cover limits net CO₂ uptake and growth of a rainforest tree during tropical rainy seasons. Proc. Natl. Acad. Sci. U. S. A. 100, 572–576.
- Griffiths, R.P., Madritch, M.D., Swanson, A.K., 2009. The effect of topography on forest soil characteristics in the Oregon Cascade Mountains (USA): implications for the effects of climate change on soil properties. For. Ecol. Manag. 257, 1–7.
- Gruber, A., Pirkebner, D., Oberhuber, W., Wieser, G., 2011. Spatial and seasonal variations in mobile carbohydrates in *Pinus cembra* in the timberline ecotone of the Central Austrian Alps. Eur. J. For. Res. 130, 173–179.
- Grytnes, J.A., 2003. Species-richness patterns of vascular plants along seven altitudinal transects in Norway. Ecography 26, 291–300.
- Handa, I.T., Körner, C., Hättenschwiler, S., 2005. A test of the treeline carbon limitation hypothesis by in situ CO2 enrichment and defoliation. Ecology 86, 1288–1300.

He, J.S., Fang, J., Wang, Z., Guo, D., Flynn, D.F., Geng, Z., 2006. Stoichiometry and large-scale patterns of leaf carbon and nitrogen in the grassland biomes of China. Oecologia 149, 115–122.

He, X., Hou, E., Liu, Y., Wen, D., 2016. Altitudinal patterns and controls of plant and soil nutrient concentrations and stoichiometry in subtropical China. Sci. Rep. 6, 24261.

Hoch, G., Körner, C., 2003. The carbon charging of pines at the climatic treeline: a global comparison. Oecologia 135, 10–21.

Hoch, G., Körner, C., 2005. Growth, demography and carbon relations of Polylepis trees at the world's highest treeline. Funct. Ecol. 19, 941–951.

Hogan, E.J., Minnullina, G., Sheppard, L.J., Leith, I.D., Crittenden, P.D., 2010. Response of phosphomonoesterase activity in the lichen *Cladonia portentosa* to nitrogen and phosphorus enrichment in a field manipulation experiment. New Phytol. 186, 926–933.

Hughes, L., 2000. Biological consequences of global warming: is the signal already apparent? Trends Ecol. Evol. 15, 56–61.

Jackson, R.B., Canadell, J., Ehleringer, J.R., Mooney, H.A., Sala, O.E., Schulze, E.D., 1996. A global analysis of root distributions for terrestrial biomes. Oecologia 108, 389–411.

Kang, H., Zhuang, H., Wu, L., Liu, Q., Shen, G., Berg, B., Liu, C., 2011. Variation in leaf nitrogen and phosphorus stoichiometry in *Picea abies* across Europe: an analysis based on local observations. For. Ecol. Manag. 261, 195–202.

Kessler, M., 2001. Patterns of diversity and range size of selected plant groups along an elevational transect in the Bolivian Andes. Biodivers. Conserv. 10, 1897–1921.

Kilpeläinen, A., Peltola, H., Ryyppö, A., Kellomaki, S., 2005. Scots pine responses to elevated temperature and carbon dioxide concentration: growth and wood properties. Tree Physiol. 5, 75–83.

Kitayama, K., Aiba, S.I., Majalap-Lee, N., Ohsawa, M., 1998. Soil nitrogen mineralization rates of rainforests in a matrix of elevation and geological substrates on Mount Kinabalu, Borneo. Ecol. Res. 13, 301–312.

Koerselman, W., Meuleman, A.F.M., 1996. The vegetation N:P ratio: a new tool to detect the nature of nutrient limitation. J. Appl. Ecol. 33, 1441–1450.

Körner, C., 1989. The nutritional status of plants from high altitudes. Oecologia 81, 379–391.

Körner, C., 2003. Alpine Plant Life: Functional Plant Ecology of High Mountain Ecosystems. Springer Science & Business Media, Berlin.

Körner, C., 2007. The use of 'altitude' in ecological research. Trends Ecol. Evol. 22, 569–574.

Körner, C., 2012. Alpine Treelines: Functional Ecology of the Global High Elevation Tree Limits. Springer Science & Business Media, Berlin.

Lee, K.S., Park, S.R., Kim, Y.K., 2007. Effects of irradiance, temperature, and nutrients on growth dynamics of seagrasses: a review. J. Exp. Mar. Biol. Ecol. 350, 144–175.

Leuschner, C., Moser, G., Bertsch, C., Roderstein, M., Hertel, D., 2007. Large altitudinal increase in tree root/shoot ratio in tropical mountain forests of Ecuador. Basic Appl. Ecol. 8, 219–230.

Li, M.H., Yang, J., 2004. Effects of microsite on growth of *Pinus cembra* in the subalpine zone of the Austrian alps. Ann. For. Sci. 61, 319–325.

Li, M.H., Hoch, G., Körner, C., 2001. Spatial variability of mobile carbohydrates within *Pinus cembra* trees at the alpine treeline. Phyton 41, 203–213.

Li, M.H., Hoch, G., Körner, C., 2002. Source/sink removal affects mobile carbohydrates in *Pinus cembra* at the Swiss treeline. Trees 16, 331–337.

- Li, M.H., Yang, J., Kräuchi, N., 2003. Growth responses of *Picea abies* and *Larix decidua* to elevation in subalpine areas of Tyrol, Austria. Can. J. For. Res. 33, 653–662.
- Li, M.H., Xiao, W.F., Wang, S.G., Cheng, G.W., Cherubini, P., Cai, X.H., Zhu, W.Z., 2008a. Mobile carbohydrates in Himalayan treeline trees I. Evidence for carbon gain limitation but not for growth limitation. Tree Physiol. 28, 1287–1296.

Li, M.H., Xiao, W.F., Shi, P., Wang, S.G., Zhong, Y.D., Liu, X.L., Shi, Z.M., 2008b. Nitrogen and carbon source–sink relationships in trees at the Himalayan treelines compared with lower elevations. Plant Cell Environ. 31, 1377–1387.

Liptzin, D., Sanford, R.L., Seastedt, T.R., 2013. Spatial patterns of total and available N and P at alpine treeline. Plant Soil 365, 127–140.

Mayor, J.R., Sanders, N.J., Classen, A.T., Bardgett, R.D., Clement, J., Fajardo, A., Lavorel, S., Sundqvist, M.K., Bahn, M., Chisholm, C., Cieraad, E., Gedalof, Z., Grigulis, K., Kudo, G., Oberski, D.L., Wardle, D.A., 2017. Elevation alters ecosystem properties across temperate treelines globally. Nature 542, 91–95.

Meinen, C., Hertel, D., Leuschner, C., 2009. Biomass and morphology of fine roots in temperate broad-leaved forests differing in tree species diversity: is there evidence of below-ground overyielding? Oecologia 161, 99–111.

Mooney, H.A., 1972. The carbon balance of plants. Annu. Rev. Ecol. Syst. 3, 315–346.

Moser, G., Leuschner, C., Hertel, D., Graefe, S., Soethe, N., Iost, S., 2011. Elevation effects on the carbon budget of tropical mountain forests (S Ecuador): the role of the belowground compartment. Glob. Chang. Biol. 17, 2211–2226.

Müller, M., Oelmann, Y., Schickhoff, U., Böhner, J., Scholten, T., 2017. Himalayan treeline soil and foliar C:N:P stoichiometry indicate nutrient shortage with elevation. Geoderma 291, 21–32.

Murphy, J., Riley, J.P., 1962. A modified single solution method for the determination of phosphate in natural waters. Anal. Chim. Acta 27, 31–36.

Nadelhoffer, K.J., Giblin, A.E., Shaver, G.R., Laundre, J.A., 1991. Effects of temperature and substrate quality on element mineralization in six arctic soils. Ecology 72, 242–253. Olsen, S.R., 1954. Estimation of Available Phosphorus in Soils by Extraction with Sodium

Bicarbonate. United States Department of Agriculture, Washington. Osaki, M., Shinano, T., Tadano, T., 1991. Redistribution of carbon and nitrogen compounds

from the shoot to the harvesting organs during maturation in field crops. Soil Sci. Plant Nutrition 37, 117–128.

Oyarzabal, M., Oesterheld, M., 2009. Phosphorus reserves increase grass regrowth after defoliation. Oecologia 159, 717–724.

Palacio, S., Hester, A.J., Maestro, M., Millard, P., 2008. Browsed Betula pubescens trees are not carbon-limited. Funct. Ecol. 22, 808–815. Pauli, H., Gottfried, M., Dullinger, S., Abdaladze, O., Akhalkatsi, M., Benito, Alonso J.L., Coldea, G., Dick, J., Erschbamer, B., Fernandez, Calzado R., Ghosn, D., Holten, J.I., Kanka, R., Kazakis, G., Kollar, J., Larsson, P., Moiseev, P., Moiseev, D., Molau, U., Mesa, Molero, Nagy, L., Pelino, G., Puscas, M., Rossi, G., Stanisci, A., Syverhuset, A.O., Theurilla, J.P., Tomaselli, M., Unterluggauer, P., Villar, L., Vittoz, P., Grabherr, G., 2012. Recent plant diversity changes on Europe's mountain summits. Science 226, 353–355.

Pendry, C.A., Proctor, J., 1996. The causes of altitudinal zonation of rain forests on Bukit Belalong, Brunei. J. Ecol. 1996, 407–418.

Peterson, D.L., 1998. Climate, Limiting Factors and Environmental Change in High-Altitude Forests of Western North America, the Impacts of Climate Variability on Forests. Springer, Berlin, Heidelberg, pp. 191–208.

Piper, F.I., Cavieres, L.A., Reyes-Diaz, M., Corcuera, L.J., 2006. Carbon sink limitation and frost tolerance control performance of the tree *Kageneckia angustifolia* D. Don (Rosaceae) at the treeline in central Chile. Plant Ecol. 185, 29–39.

Reich, P.B., Oleksyn, J., 2004. Global patterns of plant leaf N and P in relation to temperature and latitude. Proc. Natl. Acad. Sci. U. S. A. 101, 11001–11006.

Reich, P.B., Tjoelker, M.G., Pregitzer, K.S., Wright, I.J., Oleksyn, J., Machado, J.L., 2008. Scaling of respiration to nitrogen in leaves, stems and roots of higher land plants. Ecol. Lett. 11, 793–801.

Richardson, A.D., Berlyn, G.P., 2002. Spectral reflectance and photosynthetic properties of *Betula papyrifera* (Betulaceae) leaves along an elevational gradient on Mt. Mansfield, Vermont, USA. Am. J. Bot. 89, 88–94.

Richardson, A.D., Berlyn, G.P., Gregoire, T.G., 2001. Spectral reflectance of Picea rubens (Pinaceae) and Abies balsamea (Pinaceae) needles along an elevational gradient, Mt. Moosilauke, New Hampshire, USA. Am. J. Bot. 88, 667–676.

Richardson, A.D., Lee, X., Friedland, A.J., 2004. Microclimatology of treeline spruce-fir forests in mountains of the northeastern United States. Agric. For. Meteorol. 125, 53–66.

Rinnan, R., Michelsen, A., Bååth, E., Jonasson, S., 2007. Mineralization and carbon turnover in subarctic heath soil as affected by warming and additional litter. Soil Biol. Biochem. 39, 3014–3023.

Ruess, R.W., Cleve, K.V., Yarie, J., Viereck, L.A., 1996. Contributions of fine root production and turnover to the carbon and nitrogen cycling in taiga forests of the Alaskan interior. Can. J. For. Res. 26, 1326–1336.

Ryan, M.G., 1991. Effects of climate change on plant respiration. Ecol. Appl. 1, 157–167. Sala, A., Woodruff, D.R., Meinzer, F.C., 2012. Carbon dynamics in trees: feast or famine?

Tree Physiol. 32, 764–775. Seifter, S., Dayton, S., Novic, B., Muntwyler, E., 1950. The estimation of glycogen with the

anthrone reagent. Arch. Biochem. 25, 191–200. Sexton, J.P., McIntyre, P.J., Angert, A.L., 2009. Evolution and ecology of species range limits.

Annu, Rev. Ecol. Evol. Syst. 40, 415–436. Shaver, G.R., Chapin, F.S., 1995. Long-term responses to factorial, NPK fertilizer treatment

by Alaskan wet and moist tundra sedge species. Ecography 18, 259–275. Shi, P., Körner, C., Hoch, G., 2006. End of season carbon supply status of woody species

near the treeline in western China. Basic Appl. Ecol. 7, 370–377. Shi, P., Körner, C., Hoch, G., 2008. A test of the growth-limitation theory for alpine tree line

Snl, P., Korner, C., Hoch, C., 2008. A test of the growth-limitation theory for alpine tree line formation in evergreen and deciduous taxa of the eastern Himalayas. Funct. Ecol. 22, 213–220.

Shibata, O., Nishida, T., 1993. Seasonal changes in sugar and starch content of the alpine snowbed plants, *Primula cuneifolia* ssp. *hakusanensis* and *Fauria crista-galli*, in Japan. Arct. Alp. Res. 25, 207–210.

Sjögersten, S., Wookey, P.A., 2005. The role of soil organic matter quality and physical environment for nitrogen mineralization at the forest-tundra ecotone in Fennoscandia. Arct. Antarct. Alp. Res. 37, 118–126.

Soethe, N., Lehmann, J., Engels, C., 2008. Nutrient availability at different altitudes in a tropical montane forest in Ecuador. J. Trop. Ecol. 24, 397–406.

Sparks, D.L., Fendorf, S.E., Page, A.L., Helmke, P.A., Loeppert, R.H., Soltanpour, P.N., Tabatabai, M.A., Sumner, M.E., 1996. Kinetic Methods and Measurements. Soil Science Society of America, American Society of Agronomy, pp. 1275–1307.

Sterner, R.W., Elser, J.J., 2002. Ecological Stoichiometry: The Biology of Elements from Molecules to the Biosphere. Princeton University Press, Princeton.

Sundqvist, M.K., Sanders, N.J., Wardle, D.A., 2013. Community and ecosystem responses to elevational gradients: processes, mechanisms, and insights for global change. Annu. Rev. Ecol. Evol. Syst. 44, 261–280.

Sundqvist, M.K., Liu, Z., Giesler, R., Wardle, D.A., 2014. Plant and microbial responses to nitrogen and phosphorus addition across an elevational gradient in subarctic tundra. Ecology 95, 1819–1835.

Sveinbjörnsson, B., David, J., Abadie, W., Butler, A., 1995. Soil carbon and nitrogen mineralization at different elevation in the Chugach Mountains of south-central Alaska, U.S.a. Arct. Alp. Res. 27, 29–37.

Sveinbjörnsson, B., Smith, M., Traustason, T., Ruess, R.W., Sullivan, P.F., 2010. Variation in carbohydrate source-sink relations of forest and treeline white spruce in southern, interior and northern Alaska. Oecologia 163, 833–843.

Tessier, J.T., Raynal, D.J., 2003. Use of nitrogen to phosphorus ratios in plant tissue as an indicator of nutrient limitation and nitrogen saturation. J. Appl. Ecol. 40, 523–534.

Van de Weg, M.J., Meir, P., Grace, J., Atkin, O.K., 2009. Altitudinal variation in leaf mass per unit area, leaf tissue density and foliar nitrogen and phosphorus content along an Amazon-Andes gradient in Peru. Plant Ecolog. Divers. 2, 243–254.

Vetaas, O.R., Grytnes, J.A., 2002. Distribution of vascular plant species richness and endemic richness along the Himalayan elevation gradient in Nepal. Glob. Ecol. Biogeogr. 11, 291–301.

Vitousek, P.M., 1994. Beyond global warming: ecology and global change. Ecology 75, 1861–1876.

Vitousek, P.M., Howarth, R.W., 1991. Nitrogen limitation on land and in the sea: how can it occur? Biogeochemistry 13, 87–115.

- Wardle, D.A., Bardgett, R.D., Klironomos, J.N., Setälä, H., Van Der Putten, W.H., Wall, D.H., 2004. Ecological linkages between aboveground and belowground biota. Science 304, 1629–1633.
- Weintraub, M.N., 2011. Biological Phosphorus Cycling in Arctic and Alpine Soils. Phosphorus in Action. Springer, Berlin, Heidelberg, pp. 295–316.
 White, M.A., Running, S.W., Thornton, P.E., 1999. The impact of growing-season length
- White, M.A., Running, S.W., Thornton, P.E., 1999. The impact of growing-season length variability on carbon assimilation and evapotranspiration over 88 years in the eastern US deciduous forest. Int. J. Biometeorol. 42, 139–145.
- Wiley, E., Helliker, B., 2012. A re-evaluation of carbon storage in trees lends greater support for carbon limitation to growth. New Phytol. 195, 285–289.
- Yu, D., Wang, Q., Liu, J., Zhou, W., Qi, L., Wang, X., Dai, L., 2014. Formation mechanisms of the alpine Erman's birch (*Betula ermanii*) treeline on Changbai Mountain in Northeast China. Trees 28, 935–947.
- Yuan, Z.Y., Chen, H.Y., 2012. A global analysis of fine root production as affected by soil nitrogen and phosphorus. Proc. R. Soc. Lond. B Biol. Sci. 279, 3796–3802.
 Zhang, S.B., Zhang, J.L., Slik, J.W., Cao, K.F., 2012. Leaf element concentrations of terrestrial
- Zhang, S.B., Zhang, J.L., Slik, J.W., Cao, K.F., 2012. Leaf element concentrations of terrestrial plants across China are influenced by taxonomy and the environment. Glob. Ecol. Biogeogr. 21, 809–818.

- Zhao, N., He, N., Wang, Q., Zhang, X., Wang, R., Xu, Z., Yu, G., 2014. The altitudinal patterns of leaf C:N:P stoichiometry are regulated by plant growth form, climate and soil on Changbai Mountain, China. PLoS One 9, e95196.
- Zhao, N., Yu, G., He, N., Wang, R., Xu, Z., Yu, G., 2016. Invariant allometric scaling of nitrogen and phosphorus in leaves, stems, and fine roots of woody plants along an altitudinal gradient. J. Plant Res. 129, 647–657.
- Zhou, Z., 1992. The fossil history of Quercus. Acta Bot. Yunnanica 15, 21-33.
- Zhu, W.Z., Cao, M., Wang, S.G., Xiao, W.F., Li, M.H., 2012a. Seasonal dynamics of mobile carbon supply in *Quercus aquifolioides* at the upper elevational limit. PLoS One 7, e34213.
- Zhu, W.Z., Xiang, J.S., Wang, S.G., Li, M.H., 2012b. Resprouting ability and mobile carbohydrate reserves in an oak shrubland decline with increasing elevation on the eastern edge of the Qinghai–Tibet Plateau. For. Ecol. Manag. 278, 118–126.
- Zotz, G., 2004. The resorption of phosphorus is greater than that of nitrogen in senescing leaves of vascular epiphytes from lowland Panama. J. Trop. Ecol. 20, 693–696.