Above- and below-ground resource acquisition strategies determine plant species responses to nitrogen enrichment

Dianye Zhang1,3, Yunfeng Peng1, Fei Li1,2, Guibiao Yang1,2, Jun Wang1,2, Jianchun Yu1,3, Guoying Zhou1 and Yuanhe Yang1,2,*

1State Key Laboratory of Vegetation and Environmental Change, Institute of Botany, Chinese Academy of Sciences, Beijing 100093, China, 2University of Chinese Academy of Sciences, Beijing 100049, China, and 3Key Laboratory of Tibetan Medicine Research, Northwest Institute of Plateau Biology, Chinese Academy of Sciences, Xining 810008, China

*For correspondence. E-mail yhyang@ibcas.ac.cn

Received: 16 September 2020 Returned for revision: 18 February 2021 Editorial decision: 19 February 2021 Accepted: 23 February 2021 Electronically published: 25 February 2021

INTRODUCTION

The amount of reactive nitrogen (N) input to terrestrial ecosystems has dramatically increased over time due to intensified human activities (e.g. agricultural fertilization) and continuous atmospheric N deposition (Galloway et al., 2008). Reactive N enrichment can directly affect ecosystem functions such as gross primary productivity and the carbon (C) cycle by altering plant physiology and soil biogeochemistry (Manning et al., 2006; Zhang et al., 2019). Indirectly, external N input-induced changes in community structure, such as altered community composition and decreased species diversity, can also mediate the trajectories of ecosystem functions under N enrichment (Manning et al., 2006). Considering that the indirect effects of reactive N inputs on community structure might even dominate ecosystem responses (Hooper et al., 2012), our knowledge of the dynamics of community structure and the associated mechanisms is crucial for accurately predicting the responses of ecosystem functions to N enrichment.

During the last few decades, many studies have been conducted to investigate the effects of external N inputs on community structure and composition as well as the underlying mechanisms (Hautier et al., 2009; Borer et al., 2014; Dickson et al., 2014; DeMalach et al., 2017; Tian et al., 2020). These studies highlighted that aggravated light competition was the primary driver of the contrasting responses of various species to N enrichment (Dickson et al., 2014; DeMalach et al., 2017; Xiao et al., 2021). Considering that light is a unidirectional (decay from the top of the canopy to the bottom) and size-asymmetrical (taller individuals receive more light per unit of size than shorter individuals) resource (Onoda et al., 2014), N inputs would favour tall species, as they can compete for light effectively (Tilman, 1987; Dickson et al., 2014; Gross and Mittelbach, 2017). Meanwhile, the light deficiency induced by the shadow of tall species would suppress the biomass and richness of short species (Dickson et al., 2014; DeMalach et al., 2017). However, none of these studies has quantified the amount of light acquired by various species, which leaves unexplored...
the fundamental linkages between the responses of species relative abundance to N enrichment and plant light acquisition capacity. Moreover, N-induced increases in leaf area and specific leaf area (Zhang et al., 2019) would also favour light acquisition by short species even when being shaded by their taller competitors (Hirose and Werger, 1995; Kohyama and Takada, 2009; Onoda et al., 2014). For this reason, the accurate quantification of the amount of light acquired by various species is essential for better exploring the mechanisms underlying the fates of various species under N enrichment.

Apart from increasing light acquisition, plants would cope with N-induced light competition by altering their acquisition of non-N nutrients [e.g. phosphorus (P) and some micronutrients] due to the strong regulation of light use efficiency by leaf nutrient status (Wright et al., 2004). However, few studies have focused on the different trajectories of leaf non-N nutrient concentrations among different species under N input. Taking leaf P concentration as an example, the prevailing perspective suggests that leaf P concentration would decline under external N input due to the imbalance between the N-induced increase in plant P demand and the elevated soil P supply as a result of the enhancement of root and soil phosphatase activity (Li et al., 2016; Deng et al., 2017). In addition to secreting phosphatase, plants could also acquire P by altering their P resorption, mycorrhizal colonization, root morphology, root vitality, root carboxylate exudation and utilization of different soil P fractions (Shen et al., 2011; Lambers et al., 2015; Yu et al., 2020). More importantly, these P acquisition strategies could allow some species (e.g. plants with cluster roots) to acquire P effectively even under P-poor conditions (Vance et al., 2003) and further induce different responses to N enrichment in the leaf P concentrations of various species. However, to date few studies have quantified the differences in these nutrient acquisition strategies among species and considered the combined effects of above-ground (light) and below-ground (nutrients) resource acquisition in regulating the responses of species relative abundance to N enrichment.

To fill this knowledge gap, we explored the effects of N input on species relative abundance, light acquisition and leaf nutrient concentrations of two species with different resource acquisition strategies [the taller, dominant species *Stipa purpurea*, which is colonized by arbuscular mycorrhizal fungi (AMF), and the shorter subordinate species *Carex stenophylloides*, which has cluster roots] and quantified the linkages between species relative abundance and above-/below-ground resource acquisition in a Tibetan alpine steppe. We also examined the relationship between species relative abundance and resource acquisition for two extra species (subordinate species *Poa poophagorum* and subordinate species *Potentilla multifida*) whose relative abundance declined under N enrichment. To further investigate the drivers of the contrasting resource acquisition trends of *S. purpurea* and *C. stenophylloides*, we measured a series of plant and soil parameters, including plant height, leaf and root morphology, mycorrhizal colonization, root vitality, root extracellular enzyme activity, root carboxylate exudation, leaf nutrient resorption efficiency and rhizosphere soil nutrient status. The aim of our study was to explore the mechanisms underlying the species-specific responses of plant relative abundance to N enrichment. We hypothesized that above-ground (light) and below-ground (nutrients) resource acquisition would co-determine the effects of N input on species relative abundance. Specifically, plants would invest additional N in shoot growth and further acquire more light. Species relative abundance would then increase with the improved light acquisition. Meanwhile, both *S. purpurea* (colonized by AMF) and *C. stenophylloides* (with cluster roots) might alter their nutrient acquisition traits (e.g. synthesizing and secreting more phosphatase and carboxylates) to take up more non-N nutrients and thus enhance leaf nutrient concentrations. The increased leaf nutrient concentrations would then promote species relative abundance, even under a strong light competition scenario.

**MATERIALS AND METHODS**

**Site description and experimental design**

The study was carried out in an alpine steppe (37°18′ N, 100°15′ E; 3290 m a.s.l.) located on the north-eastern Tibetan Plateau, China. The study site experiences an alpine continental climate in which cold and dry winters alternate with relatively warm and wet summers. The mean annual temperature is 0.08 °C, with average precipitation of ~390 mm, falling predominantly in May–September. The local vegetation is composed of the dominant species (relative abundance >20 %; Ye et al., 2018) *Stipa purpurea*, subordinate species (relative abundance between 1 % and 20 %; Sánchez-Castillo et al., 2008), including *Carex stenophylloides*, *Poa poophagorum*, *Leymus secalinus*, *Agropyron cristatum*, *Potentilla multifida* and *Heteropappus altaicus*, and several rare species (relative abundance <1 %; Mouillot et al., 2013), including *Dracocephalum heterophyllum*, *Leontopodium nanum* and *Potentilla bifurca*. The soil type is a Haplic Calcisol according to the FAO classification system, with 11.6 ± 3.6 mg kg⁻¹ of inorganic N (extracted by 1 m KCl) and 2.2 ± 0.3 mg kg⁻¹ of Olsen-P (labile inorganic P extracted by 0.5 m NaHCO₃) in the top 30 cm of soil (Peng et al., 2017). This alpine steppe was historically used as a winter pasture for sheep, and no additional management practices (such as fertilization or irrigation) were applied before our experiment.

We fenced the experimental field (0.5 ha) and established the N manipulation experiment in May 2013. The experiment was set in a randomized complete block design, with five blocks (isolated by buffer zones of 2 m) and eight N treatments (0, 1, 2, 4, 8, 16, 24 and 32 g N m⁻² year⁻¹). The N treatments ranged from N limitation to N saturation, which may occur at ~15 g N m⁻² year⁻¹ in grassland ecosystems around the world (Peng et al., 2020). Considering that the N saturation threshold appears to be ecosystem-dependent [e.g. the results from an alpine meadow on the Tibetan Plateau showed that below-ground net primary productivity first increased and then decreased at 2–4 g N m⁻² year⁻¹ (Wang et al., 2019), while the addition of 10 g N m⁻² year⁻¹ had a weak effect on most functional traits in Konza Prairie (La Pierre and Smith, 2015)], we established this N addition gradient (eight N levels up to 32 g N m⁻² year⁻¹) to ensure that non-linear responses would be included and that the saturation threshold would be detected. In each block, the eight N addition levels were randomly assigned to 6 × 6-m² plots (isolated by buffer zones of 1 m). The N fertilizer applied...
Species relative abundance, light acquisition and leaf nutrient concentration measurements

The relative abundances of S. purpurea, C. stenophylloides, P. poophagorum and P. multifida were determined in mid-August 2016. Specifically, we recorded the species richness and the number of individuals per species (i.e. the abundance of the targeted species) based on a permanent 1 x 1-m$^2$ quadrat. The species relative abundance was then calculated as the ratio of the number of individuals of a species to the number of individuals within the quadrat.

To quantify the light acquisition of S. purpurea, C. stenophylloides, P. poophagorum and P. multifida, we divided the canopy into several layers (equal to the number of species) according to the maximum vegetative height of the different species in each plot (Supplementary Data Fig. S1A). We then calculated the amount of photosynthetically active radiation (PAR; 400–700 nm waveband) acquired by each species using the following formula (Anten and Hirose, 1999; Kamiyama et al., 2010):

$$PAR_i = \sum_{j}^{n} PAR_{ij}$$

where $PAR_i$ is the amount of PAR acquired by species $i$ in the whole canopy ($\mu$mol photons m$^{-2}$ s$^{-1}$), $PAR_{ij}$ is the PAR acquired by species $i$ in layer $j$ ($\mu$mol photons m$^{-2}$ s$^{-1}$) and $n$ is the number of layers (i.e. the number of species). $PAR_{ij}$ was quantified using the following equation (Hirose and Werger, 1995; Anten and Hirose, 1999; Kamiyama et al., 2010):

$$PAR_{ij} = PAR_j \times \left( \frac{LA_{ij}}{\sum_{i}^{s} LA_{ij}} \right)$$

where $PAR_j$ is the PAR absorbed by layer $j$ ($\mu$mol photons m$^{-2}$ s$^{-1}$), $LA_{ij}$ is the leaf area of species $i$ in layer $j$ (cm$^2$) and $s$ is the number of species existing in layer $j$. To estimate $LA_{ij}$, all plants in three 0.25 x 0.25-m$^2$ quadrats were harvested and sorted by individual species. The harvested plant materials for each species were cut into several segments starting at the base. The species with a maximum height taller than 15 cm were cut into 5-cm segments and the other species were cut into 2-cm segments (Hirose and Werger, 1995; Anten and Hirose, 1999). The total leaf area of each segment was measured for each species, and empirical functions of the relationship between leaf area and height of the segments were established for the target species (Ramesh et al., 2007). Then, $LA_{ij}$ was estimated on the basis of the established empirical function and the height of layer $j$ for species $i$. $PAR_{ij}$ can be estimated according to the intercepted and reflected PAR in layer $j$ (Supplementary Data Fig. S1B; Tagesson et al., 2015):

$$PAR_j = PAR_{inc,j} - PAR_{inc,j+1} - PAR_{ref,j} + PAR_{ref,j+1}$$

where $PAR_{inc,j}$ is the incoming PAR to layer $j$ ($\mu$mol photons m$^{-2}$ s$^{-1}$) and $PAR_{ref,j}$ is the PAR reflected by layer $j$ ($\mu$mol photons m$^{-2}$ s$^{-1}$).

To obtain the incoming and reflected PAR for a given layer, we measured these values at eight heights (0, 5, 10, 20, 40, 60, 80 and 100 cm) in each plot using a canopy analyser [an analyser with a linear light ceptometer (50 cm length x 2 cm width x 2 cm height); Top-1000; Zhejiang Top Instrument, Hangzhou, Zhejiang, China]. The PAR measurements were taken between 11:00 a.m. and 14:00 p.m. on a sunny and cloudless day in mid-August, with ten repetitions in each plot. We then quantified the association between the observed PAR and canopy height according to the following light decay equation (Supplementary Data Fig. S1C; Anten and Hirose, 1999; DeMalach et al., 2017):

$$\log (PAR) = a \times \ln (H) + b$$

where $PAR$ is the incoming or reflected PAR ($\mu$mol photons m$^{-2}$ s$^{-1}$) and $H$ is the canopy height (cm). Finally, we calculated the incoming and reflected PAR for the specific layers based on the light decay equation and the height of the layer in each plot.

For the leaf nutrient concentrations of S. purpurea, C. stenophylloides, P. poophagorum and P. multifida, we measured the leaf P, potassium (K), calcium (Ca), sodium (Na), magnesium (Mg), aluminium (Al), copper (Cu), iron (Fe), manganese (Mn) and zinc (Zn) concentrations because N addition can aggravate plant non-N nutrient limitation (Li et al., 2016). Specifically, the non-N nutrient concentrations of the leaves were measured by inductively coupled plasma atomic emission spectroscopy (ICAP6300, Thermo Fisher Scientific, Waltham, MA, USA) after digesting the leaf samples with acid. In addition, the leaf N concentration was determined using an elemental analyser (Vario EL III, Elementar, Germany).

Plant resource acquisition trait measurements

For the plant light acquisition traits, we measured the plant height, leaf area and specific leaf area (a large plant size and specific leaf area can promote plant light acquisition; Pérez-Harguindeguy et al., 2013; Laurans and Vincent, 2016). Specifically, we measured 20 healthy and mature individuals to obtain the vegetative height for S. purpurea and C. stenophylloides in the field and then harvested the leaves of these individuals. The collected leaves were scanned and analysed by WinFOLIA software (Regent Instruments, Quebec City, Quebec Canada) to determine their leaf area, and oven-dried to obtain their leaf weight. The specific leaf area was calculated as the ratio of leaf area to leaf weight (Pérez-Harguindeguy et al., 2013). For details of the procedure for measuring light acquisition traits see Zhang et al. (2019).

For the plant nutrient acquisition traits, we measured mycorrhizal colonization, specific root length, root vitality (plants with higher mycorrhizal colonization, specific root length and root vitality can take up more soluble inorganic nutrients from the soil; Shen et al., 2011; Treseder, 2013), root phosphomonoesterase (PME) activity and root...
carboxylate exudation (plants can secrete PME to hydrolyse soluble organic nutrients and carboxylates to mobilize insoluble nutrient forms; thus, exuding more PME and carboxylates can increase the soil available nutrient supply; Marklein and Houlton, 2012; Lambers et al., 2015). Specifically, we excavated the roots from the top 0–10 cm of soil at five random locations within each plot. The collected root samples within each plot were mixed and classified as S. purpurea, C. stenophylloides or other species by their morphological characteristics. Then, the root samples of S. purpurea and C. stenophylloides were classified into different orders, with the most distal root tips being termed the first order (Xia et al., 2010). We chose to measure root P acquisition traits of the first three orders of roots since they are closely related to plant nutrient uptake (McCormack et al., 2015).

After root sampling, mycorrhizal colonization was measured using the trypan blue staining technique (Johnson et al., 2003). Briefly, fresh roots were cleared with KOH solution, bleached with alkaline H₂O₂, and soaked in an acid solution. The acidified roots were stained in acidic trypan blue solution, destained in acidic glycerol and then used to estimate mycorrhizal colonization. Specific root length refers to the ratio of root length to root weight (Pérez-Harguindeguy et al., 2013). The rinsed roots were scanned and analysed with WinRHIZO software (Regent Instruments, Quebec City, Quebec, Canada) to measure their root length. Then, the samples were oven-dried and weighed to determine their specific root length. Root vitality was measured using the triphenyltetrazolium chloride (TTC) reduction method (Comas et al., 2000). For the TTC reduction test, ~500 mg of rinsed roots was soaked in TTC buffer solution, vacuum-infiltrated, and subsequently incubated for 24 h in the dark. During the thermostatic incubation (25 °C), colourless TTC was reduced to red triphenyl-formazan. The amount of TTC reduction, which is proportional to root vitality, was then determined with a spectrophotometer (UV-2550, Shimadzu, Kyoto, Japan) at 490 nm.

Root carboxylate (citrate and malate) exudation was measured with a citric acid assay kit (K-CITR, Megazyme, Wicklow, Ireland) and a malic acid assay kit (K-LMALQR, Megazyme, Wicklow, Ireland) following the manufacturer’s protocols. The citric acid assay kit quantified citrate according to the oxidation of nicotinamide adenine dinucleotide (NADH), while the malic acid assay kit determined malate by the formation of NADH (Dinkelaker et al., 1989; Kabir et al., 2013). The changes in NADH were determined with a spectrophotometer (UV-2550, Shimadzu, Kyoto, Japan) at 340 nm. Root PME activity was determined according to the method (Marx et al., 2001). First, we soaked ~200 mg of rinsed roots in 50 mL of Tris-buffer solution and incubated them for 1 h at 25 °C. Subsequently, 200 μL of root-extracted solution was placed in a 96-well microplate and mixed with 50 μL of 4-methylumbellifere-related substrate, and the mixture was incubated for 2 h at 25 °C in the dark. After incubation, the fluorescence of each sample, which is proportional to the PME activity, was determined with an automatic microplate reader (DTX 880 Multimode Detector, Beckman Coulter, Fullerton, CA, USA) with 365-nm excitation and 450-nm emission wavelengths.

We also measured the plant nutrient resorption efficiency (Vergutz et al., 2012) and root:shoot ratio (Gedroc et al., 1996), which are closely associated with plant nutrient acquisition. Phosphorus resorption efficiency was calculated according to the following equation (Vergutz et al., 2012):

\[ \text{P resorption efficiency} = \left(1 - \frac{P_i}{P_5 \times \text{MLCF}}\right) \times 100 \]

where \( P_i \) is the P concentration of mature green leaf (mg g⁻¹; sampled in mid-August), \( P_5 \) is the P concentration of fresh leaf litter (mg g⁻¹; sampled in late October) and MLCF is the mass loss correction factor (the MLCFs of S. purpurea and C. stenophylloides are considered to be 0.713 and 0.640, respectively; Vergutz et al., 2012). The root:shoot ratios for S. purpurea and C. stenophylloides were calculated as the ratio of the root biomass in the top 30 cm of soil to the corresponding above-ground biomass.

We further determined the rhizosphere soil P fractions using the sequential P fractionation method (Hedley et al., 1982; Tiessen and Moir, 1993). Rhizosphere soil collection was performed at the same time as the root sampling; we collected the soil that was still adhered to the roots after gently shaking the roots (Marilley et al., 1998). The soil samples were air-dried and analysed by the sequential extraction method (resin extraction followed by 0.5 M NaHCO₃ extraction, 0.1 M NaOH extraction and 1 M HCl extraction) to extract labile resin-inorganic P (Pi), NaHCO₃-Pi and NaHCO₃–organic P (Po); moderately labile NaOH-Pi and NaOH-Po; and recalcitrant HCl-Pi (Hedley et al., 1982, 1983; Tiessen and Moir, 1993). The Pi in the extractant was measured using the ammonium molybdate method, and Po was calculated as the difference between the total P (determined by inductively coupled plasma atomic emission spectroscopy) and Pi. In addition, we measured the rhizosphere soil-available micronutrients (Cu, Fe, Mn, Zn) using inductively coupled plasma atomic emission spectroscopy after extracting soil samples according to the diethylenetriaminepentaacetic acid procedure (Lindsay and Norvell, 1978).

**Statistical analyses**

We analysed the data with the following three steps. First, one-way ANOVAs were used to explore the effects of N enrichment on species relative abundance, plant resource acquisition (plant PAR acquisition, leaf nutrient concentration), plant traits, and rhizosphere soil nutrient status. During these analyses, the N addition level was assigned as a fixed factor, and the block was assigned as a random factor (because the effects of the block were caused by heterogeneity; Dutilleul, 1993). Subsequently, post hoc analyses (Tukey’s honestly significant difference test; Hothorn et al., 2008) were conducted to examine the differences among treatments for those variables that exhibited significant changes along the N gradient.

Second, linear mixed-effects regression models were used to identify the drivers of species relative abundance under extra N input. Specifically, we examined the single-variable relationships of species relative abundance with resource acquisition based on linear mixed-effects regression models. During these analyses, resource acquisition was treated as a fixed factor and
the block was treated as a random factor. Subsequently, single-variable linear mixed-effects regression models were developed to examine the relationships of resource acquisition to plant resource acquisition traits and rhizosphere soil nutrient status. In these linear mixed-effects models, plant traits and rhizosphere soil nutrient status were treated as fixed factors and the block was treated as a random factor. All residuals of the models were tested for homoscedasticity and normality. The ANOVAs and linear mixed-effects models were performed using the lme4 package (Bates et al., 2015) in R 3.6.0 (R Core Team, 2019).

Third, structural equation models were used to explore the relative effects of the predictors on species relative abundance. In the initial structural equation model, the N addition level was considered as an exogenous variable; plant resource acquisition, plant traits and rhizosphere soil nutrient status were set as the endogenous variables; and species relative abundance was the final response variable (Supplementary Data Table S1).

Then, iterative model optimization was performed to improve the goodness-of-fit of the model. Fisher’s C statistic, Akaike’s information criterion corrected for small sample size (AICc) and the whole-model P value were used to evaluate the model performance (Lefcheck, 2016). Lower Fisher C and AICc values reflected a better-fitting model (Lefcheck, 2016). The structural equation models were conducted using the piecewiseSEM package in R 3.6.0 (R Core Team, 2019) with the block as the random effect. Notably, as some plant relative abundance and light acquisition parameters showed unimodal responses to N enrichment with a threshold of 8 g N m⁻² year⁻¹, we divided the corresponding data into two parts (≤8 g N m⁻² year⁻¹, low N; and >8 g N m⁻² year⁻¹, high N) and conducted the linear mixed-effects regression model and structural equation model under the low-N and high-N treatments, respectively.

RESULTS

Nitrogen-induced changes in species relative abundance, resource acquisition, plant traits and rhizosphere soil nutrient status

Our results showed that species relative abundance, PAR acquisition and leaf P and micronutrient concentrations showed different responses to N enrichment (Fig. 1; Supplementary Data Table S1). Specifically, along the N addition gradient, the relative abundance and PAR acquisition of the dominant species *S. purpurea* increased (all *P* < 0.05; Fig. 1A, B),

![Fig. 1](image_url). Changes in species relative abundance, light acquisition and leaf P concentration along the experimental N gradient. Values are means ± s.e. (*n* = 5), and the same letter denotes a non-significant difference among N treatments (*P* > 0.05). DW, dry weight; N0, N1, N2, N4, N8, N16, N24 and N32 represent N addition rates of 0, 1, 2, 4, 8, 16, 24 and 32 g N m⁻² year⁻¹, respectively.
while its leaf P concentration declined ($P < 0.05$; Fig. 1C). For *C. stenophylloides*, relative abundance almost doubled ($P < 0.05$; Fig. 1D) and leaf P and micronutrient (Cu, Mn and Zn) concentrations increased along the N addition gradient (all $P < 0.05$; Fig. 1F; Supplementary Data Table S1); PAR acquisition showed a unimodal response to N enrichment with a threshold of 8 g N m$^{-2}$ year$^{-1}$ (Fig. 1E). For *P. poophagorum*, relative abundance, PAR acquisition and leaf P concentration all declined along the experimental N addition gradient (all $P < 0.05$; Fig. 1G–I). For *P. multifida*, relative abundance and PAR acquisition exhibited hump-shaped responses to N enrichment (all $P < 0.05$; Fig. 1J, K), while leaf P concentration declined along the N addition gradient ($P < 0.05$; Fig. 1L).

In terms of light acquisition traits, the plant height, leaf area and specific leaf area of *S. purpurea* and *C. stenophylloides* all increased along the experimental N addition gradient (all $P < 0.05$; Fig. 2). For nutrient acquisition traits, the mycorrhizal colonization of *S. purpurea* decreased, while the root vitality, root phosphatase activity and root carboxylate exudation of *S. purpurea* and *C. stenophylloides* increased along the N addition gradient (all $P < 0.05$; Fig. 3A, C–E; the results are shown separately for citrate and malate in Supplementary Data Fig. S3). Moreover, the root:shoot ratio of *C. stenophylloides* declined under N enrichment ($P < 0.05$; Supplementary Data Fig. S4), while the specific root length and P resorption efficiency of *S. purpurea* and *C. stenophylloides* remained stable (Fig. 3B, F). In terms of the rhizosphere soil P fraction, N enrichment significantly reduced the resin-Pi, NaHCO$_3$-Pi and NaHCO$_3$-Po of *S. purpurea* rhizosphere soil (all $P < 0.05$; Table 1). The NaOH-Pi, NaOH-Po and HCl-Pi of *C. stenophylloides* rhizosphere soil decreased under N enrichment (all $P < 0.05$; Table 1). For rhizosphere soil micronutrient availability, N enrichment significantly decreased the rhizosphere soil Cu, Fe, Mn and Zn availability for both *S. purpurea* and *C. stenophylloides* (all $P < 0.05$; Supplementary Data Table S2).

**Linkages between species relative abundance and resource acquisition, plant traits and rhizosphere soil nutrient status**

Our results indicated that the relative abundances of *S. purpurea*, *P. poophagorum* and *P. multifida* exhibited positive correlations with plant PAR acquisition under both low and high N levels (all $P < 0.05$; Fig. 4A, E, G). Moreover, although the leaf P concentration was positively correlated with the relative abundance for *P. poophagorum* under both low and high N levels, and for *P. multifida* under high N levels, it exhibited negative correlations with the relative abundance for *S. purpurea* and *P. multifida* under low N levels (all $P < 0.05$; Fig. 4B, F, H). For *C. stenophylloides*, relative abundance was positively correlated with both plant PAR acquisition and leaf nutrient (P, Mn and Zn) concentrations under low N levels (all $P < 0.05$; Fig. 4C, D; Supplementary Data Fig. S5B, C). However, under high N levels, the relative abundance of *C. stenophylloides* exhibited positive correlations only with leaf nutrient (P, Mn and Zn) concentrations ($P < 0.05$; Fig. 4D; Supplementary Data Fig. S5B, C).

Our results also revealed that the PAR acquisition of *S. purpurea* and *C. stenophylloides* was closely associated with plant height and leaf area (all $P < 0.05$; Fig. 5A, B). The leaf P concentration of *S. purpurea* showed a positive relationship with mycorrhizal colonization under low N levels, and decreased with increasing root vigour, root phosphatase activity and root carboxylate exudation along the entire N gradient (all $P < 0.05$; Fig. 5A). However, leaf P concentration of *C. stenophylloides* exhibited positive associations with root vigour, root phosphatase activity and root carboxylate...
exudation, and negative correlations with the moderately labile (NaOH-Pi and NaOH-Po) and recalcitrant (HCl-Pi) fractions in rhizosphere soil (all $P < 0.05$; Fig. 5B). In addition, the leaf micronutrient concentration of Carex stenophylloides showed a positive relationship with root carboxylate exudation ($P < 0.05$; Supplementary Data Fig. S6).

Further analyses demonstrated that light was the dominant factor driving the response of the relative abundance of Stipa purpurea to N enrichment (Fig. 6A, B). Acquisition of PAR could explain 57 and 75 % of the variation in the relative abundance of Stipa purpurea at low N and high N levels, respectively. For Carex stenophylloides, PAR acquisition and leaf P concentration co-determined the response of relative abundance under low N levels (Fig. 6C). However, the increased leaf P concentration induced by increased carboxylate exudation was the most important driver of the relative abundance of Carex stenophylloides under high N levels (Fig. 6D). The combination of the leaf P concentration and carboxylate exudation explained 59 % of the variation in the relative abundance of Carex stenophylloides under high N levels.

DISCUSSION

Based on the quantification of above-ground (light) and below-ground (P and other non-N nutrients) resources acquired by specific species, this study provided the first evidence that light and nutrients co-drove the response of plant species to N enrichment (Fig. 7). Specifically, structural equation model and stepwise regression analyses (Supplementary Data Table S3) revealed that
Table 1. Responses of thyrhesphere soil P fractions (mg kg$^{-1}$) of S. purpurea and C. stenophylloides to increasing N addition

| Species          | Soil P Fraction | P-value | 0  | 2  | 4  | 8  | 16 | 24 | 32 |
|------------------|-----------------|---------|----|----|----|----|----|----|----|
| S. purpurea      | NaHCO$_3$-Pi    | 0.04    | 3.4 ± 0.1 a | 3.2 ± 0.3 ab | 3.3 ± 0.3 ab | 3.4 ± 0.4 ab | 2.5 ± 0.4 ab | 2 ± 0.3 ab | 2.2 ± 0.3 ab | 1.7 ± 0.2 b |
|                  | NaHCO$_3$-Po    | <0.01   | 2.7 ± 0.1 ab | 1.9 ± 0.3 ab | 3.1 ± 0.2 a  | 3 ± 0.3 ab   | 2 ± 0.1 b   | 2.1 ± 0.2 ab | 2.1 ± 0.2 ab | 2.2 ± 0.1 b |
|                  | NaOH-Pi         | 0.05    | 14.6 ± 0.5  | 14.5 ± 0.8  | 16.7 ± 0.8  | 12.8 ± 0.3  | 14.4 ± 0.5  | 12.9 ± 0.8  | 13.8 ± 0.7  | 13.8 ± 0.5  |
|                  | NaOH-Po         | 0.05    | 11.8 ± 0.5  | 9.6 ± 0.6   | 11 ± 0.7    | 10.3 ± 0.8  | 11.6 ± 0.9  | 10.8 ± 0.7  | 10.4 ± 0.4  | 10.9 ± 0.3  |
|                  | HCL-Pi          | 0.3     | 65.7 ± 1.2  | 65.4 ± 1.0  | 65.2 ± 0.6  | 63.2 ± 1.6  | 62.8 ± 0.8  | 61 ± 1.1    | 62.7 ± 3.7  | 63.4 ± 2.1  |
|                  | Resin-Pi        | 0.08    | 5.0 ± 0.3   | 5.3 ± 0.3   | 4.6 ± 0.5   | 4.3 ± 0.3   | 4.5 ± 0.5   | 4.2 ± 0.6   | 4.3 ± 0.5   | 4.2 ± 0.2   |
| C. stenophylloides | NaHCO$_3$-Po   | 0.88    | 2.6 ± 0.2   | 2.4 ± 0.1   | 2.6 ± 0.1   | 2.5 ± 0.2   | 2.6 ± 0.1   | 2.5 ± 0.2   | 2.4 ± 0.1   | 2.4 ± 0.2   |
|                  | NaOH-Pi         | <0.01   | 14.5 ± 1.2  | 14 ± 1.2 ab | 12.7 ± 0.6  | 13.3 ± 1.8 ab | 11.8 ± 1.1 ab | 12.4 ± 0.8 ab | 10.9 ± 0.7 b | 11.8 ± 0.8 ab |
|                  | NaOH-Po         | 0.04    | 11.3 ± 0.6 ab | 10.5 ± 0.7 ab | 11.5 ± 0.4 a | 10.2 ± 0.6 ab | 10.3 ± 1.1 ab | 8.9 ± 0.9 b   | 9.7 ± 1.1 ab | 8.7 ± 0.7 b |
|                  | HCL-Pi          | <0.01   | 67.3 ± 1.2 a | 63 ± 1.7 ab | 61.5 ± 1.9 abc | 57.7 ± 2.2 bcd | 54.5 ± 1.2 bcd | 54.8 ± 1.1 bcd | 53.8 ± 2 cd | 52.5 ± 1.1 d |

Values in a specific row followed by the same letter denote a non-significant difference among N treatments (P > 0.05).

Light acquired by each species dominated the different fates of S. purpurea, P. poophagorum and P. multifida under N enrichment. This finding advances our understanding of the crucial role of light in shaping the complex responses (increasing, decreasing and unimodal) of plant species to N enrichment beyond the traditional perspective about the inhibition of short species by monotonously increasing community light interception (Borer et al., 2014) or light asymmetry (Supplementary Data Fig. S7; DeMalach et al., 2017). For S. purpurea, its stronger light acquisition promoted its relative abundance, while the weaker light acquisition of P. poophagorum induced a decrease in its relative abundance. It has been reported that light is a unidirectional and asymmetrical resource input from the top of the canopy to the bottom (Onoda et al., 2014) and that N enrichment would increase light asymmetry (DeMalach et al., 2017). In this study, S. purpurea acquired light more efficiently than other species because S. purpurea included tall individuals and showed increased leaf area along the N addition gradient. Meanwhile, the light acquisition of P. poophagorum may be suppressed by increased shading and light asymmetry (Borer et al., 2014; DeMalach et al., 2017); thus, its relative abundance decreased. However, for P. multifida, whose light acquisition showed a unimodal response to N enrichment, its relative abundance increased initially but decreased after a threshold of 8 g N m$^{-2}$ year$^{-1}$. The initial increase in light acquisition at low N levels may have occurred because N enrichment also promoted the growth of shorter species and increase their plant height and leaf area (Zhang et al., 2019), which was beneficial for light acquisition. Consequently, the relative abundance of P. multifida increased, especially under low N levels when the shading effect of taller species and the light asymmetry were not strong. Nevertheless, the inhibitory effect of the intensified shading exceeded the facilitation of increased plant size at high N levels, leading to decreased light acquisition and relative abundance for P. multifida. Overall, the quantification of light by different species, as performed in this study, offers the possibility of revealing the mechanisms underlying the species-specific responses to various N addition levels.

Our results also demonstrated that the increased leaf nutrient (especially P) concentrations of C. stenophylloides offset the negative effect of decreased light acquisition under high N levels, and promoted its relative abundance even in the shade of taller competitors (Fig. 6C, D). This finding agrees with our hypothesis that above- and below-ground resource acquisition strategies co-determine the response of community structure in the case of N input and does not support the previously suggested view that light itself determines species persistence under N enrichment (Hautier et al., 2009; Borer et al., 2014; DeMalach et al., 2017). Although light competition is important in regulating community structure under N enrichment, plants would cope with light competition not only by modifying their light acquisition capacity but also by altering their light use efficiency (Anten, 2005; Onoda et al., 2014). In this study, light acquisition by S. purpurea increased along the N addition gradient, which stimulated its relative abundance. However, for C. stenophylloides the light acquired initially increased but subsequently decreased at levels higher than 8 g N m$^{-2}$ year$^{-1}$. Nevertheless, N enrichment improved the light use efficiency of C. stenophylloides (Supplementary Data Fig. S8A), and thus induced its high relative abundance at high N levels. The increased light use efficiency of C. stenophylloides...
Fig. 4. Relationships of species relative abundance with light acquisition and leaf P concentration. DW, dry weight; N0, N1, N2, N4, N8, N16, N24 and N32 represent N addition rates of 0, 1, 2, 4, 8, 16, 24 and 32 g N m⁻² year⁻¹, respectively.
Fig. 5. Linkages of light acquisition and leaf P concentration with plant traits and rhizosphere soil P fraction for *S. purpurea* (A) and *C. stenophylloides* (B). The plant and soil parameters shown in the figure all exhibit significant effects on plant resource acquisition (all $P < 0.05$). The data points represent standardized regression coefficients obtained from single-variable regression analysis and the lines represent 95% credible intervals. N0, N1, N2, N4, N8, N16, N24 and N32 represent N addition rates of 0, 1, 2, 4, 8, 16, 24 and 32 g N m$^{-2}$ year$^{-1}$, respectively.
could be driven by its higher leaf P concentrations under N enrichment, due to the close associations between light use efficiency and plant nutrient levels (Supplementary Data Fig. S8B; Wright et al., 2004).

We further explored the mechanisms underlying the increased leaf P concentrations for *C. stenophylloides* under N enrichment. The prevailing perspective suggests that N enrichment should elevate soil P availability but that the increased soil P supply always fails to meet the increased plant P demand and thus induces a decrease in leaf P concentrations (Deng et al., 2017). In contrast, we observed that *C. stenophylloides* showed efficient P acquisition under N enrichment through its cluster roots, which further elevated its leaf P concentration. Under the external N input, *C. stenophylloides* increased its investment in below-ground nutrient acquisition, and its cluster roots became more efficient in acquiring P. Specifically, N enrichment stimulated the exudation of carbon-rich carboxylates (Fig. 3E), which can release soluble Pi and Po from insoluble P forms that are strongly sorbed onto soil particles by ligand exchange (mainly NaOH-Pi and Po, and HCl-Pi; Vance et al., 2003; Lambers et al., 2015; Table 1). The soluble Pi released during this process can be directly absorbed by cluster roots, while the soluble Po is first converted into soluble Pi by phosphatase and then taken up by plants (Lambers et al., 2015). Meanwhile, the root vitality and root PME activity of *C. stenophylloides* also increased significantly after additional N input. Consequently, the leaf P concentration of *C. stenophylloides* increased along the N addition gradient. Interestingly, though the root vitality, root PME activity and root carboxylate exudation of *S. purpurea* also increased under N enrichment, its leaf P concentration declined. This might be due to the fact that *S. purpurea* invested more photosynthate in shoot growth, reduced its below-ground C inputs, and induced a decline in AMF root colonization along the N gradient (the plant supplied all the C that the fungi required; Smith et al., 2011). Although AMF could only utilize the labile P forms, it usually takes up soluble Pi more efficiently than roots (by exploring more soil volume; Smith et al., 2011; Raven et al., 2018). As a result, the decreased AMF root coloni- zation might restrict P absorption of *S. purpurea*, and contrib- uted to the decline of its leaf P concentration.

In addition to P, the changed nutrient acquisition traits could also help in the acquisition of other micronutrients. Consistent with this deduction, our results revealed a positive association between leaf micronutrient concentrations of *C. stenophylloides* and its root carboxylate exudation (Supplementary Data Fig. S6). Nevertheless, apart from alleviating nutrient limitations, the accumulation of micronutrients in leaves can have a negative effect on *C. stenophylloides* due to the inhibition of the plant photosynthetic rate and growth by metal toxicity (Tian et al., 2016). These contrasting effects of leaf micronutrients may have led to the weak linkages between the relative abundance and leaf micronutrient concentration of *C. stenophylloides* along the N addition gradient.

In summary, this study revealed that above- and below-ground resource acquisition strategies co-determine the responses of plant species to N enrichment (Fig. 7). The relative abundance of the taller *S. purpurea* increased with its increased light acquisition. Meanwhile, the increased non-N nutrient (especially P) concentrations in leaves promoted the relative abundance of cluster-rooted *C. stenophylloides* even in the shade of taller competitors. These results demonstrated that the increased leaf P in the shorter species stimulated their light use efficiency and facilitated their relative abundance along the N addition gradient. These findings conflict with previous findings that the aggravation of light limitation induced by N input drives shorter species to extinction (Borer et al., 2014; DeMalach et al., 2017). Given that...
decreases in species diversity and changes in community composition always constrain the responses of ecosystem productivity to N enrichment (Hooper et al., 2012), the persistence of short species may contribute to the consistent positive effects of N inputs on ecosystem productivity. Therefore, considering the differences in species resource acquisition strategies can help to reveal the mechanisms underlying the dynamics of community structure and ecosystem functions under an N enrichment scenario.

SUPPLEMENTARY DATA

Supplementary data are available online at https://academic.oup.com/aob and consist of the following. Figure S1: schematic diagram showing the quantification of photosynthetically active radiation acquired by a specific species within a given plot. Figure S2: the initial structural equation model. Figure S3: response of root citrate and malate exudation to N enrichment. Figure S4: response of root:shoot ratio of *C. stenophylloides* to N enrichment. Figure S5: relationships between species relative abundance and leaf micronutrient concentrations for *C. stenophylloides*. Figure S6: linkages of leaf micronutrient concentrations with plant traits and micronutrient availability for *C. stenophylloides*. Figure S7: changes in community-level light asymmetry along the N addition gradient. Figure S8: response of light use efficiency of *C. stenophylloides* to N enrichment and its linkage with leaf P concentration. Table S1: effects of N addition on plant leaf nutrient concentrations. Table S2: responses of rhizosphere soil micronutrients availability to N enrichment. Table S3: optimal models for the relative abundances of *Poa poophagorum* and *Potentilla multifida*.

FUNDING

This work was supported by the National Natural Science Foundation of China (31825006, 31988102 and 91837312),
LITERATURE CITED

Anten NPR. 2005. Optimal photosynthetic characteristics of individual plants in vegetation stands and implications for species coexistence. Annals of Botany 95: 495–506.

Anten NPR, Hirose T. 1999. Interspecific differences in above-ground growth patterns result in spatial and temporal partitioning of light among species in a tall-grass meadow. Journal of Ecology 87: 583–597.

Bates D, Maechler M, Bolker B, Walker S. 2015. Fitting linear mixed-effects models using lme4. Journal of Statistical Software 67: 1–48.

Borer ET, Seabloom EW, Gruner DS, Anten NPR. 2005. Herbivores and nutrients control grassland plant diversity via light limitation. Nature 508: 517–520.

Comas LH, Eissenstat DM, Lakso AN. 2000. Assessing root death and root system dynamics in a study of grape canopy pruning. New Phytologist 147: 171–178.

DeMichele N, Zandy E, Kadmon R. 2017. Light asymmetry explains the effect of nutrient enrichment on grassland diversity. Ecology Letters 20: 60–69.

Deng Q, Hui DF, Dennis S, Reddy KC. 2017. Responses of terrestrial ecosystem phosphorus cycling to nitrogen addition: a meta-analysis. Global Ecology and Biogeography 26: 713–728.

Dickson TL, Mittelbach GG, Reynolds HL, Gross KL. 2014. Height and clumpiness traits determine plant community responses to fertilization. Ecology 95: 2443–2452.

Dinkelaker B, Römheld V, Marschner H. 2001. Transformations of the study of enzyme diversity in soils. Soil Biology and Biochemistry 33: 1633–1649.

McCormack ML, Dickie IA, Eissenstat DM, et al. 2015. Redefining fine roots improves understanding of below-ground contributions to terrestrial biosphere processes. New Phytologist 207: 505–518.

Mouillot D, Bellwood DR, Baraloto C, Mouillot D, Bellwood DR, Baraloto C, et al. 2013. Rare species support vulnerable functions in high-diversity ecosystems. PLoS Biology 11: e1001569.

Mubunyi AR, Houlton BZ. 2012. Nitrogen inputs accelerate phosphorus cycling rates across a wide variety of terrestrial ecosystems. New Phytologist 193: 696–704.

Marx MC, Wood M, Jarvis SC. 2001. A microplate fluorimetric assay for the study of enzyme diversity in soils. Soil Biology and Biochemistry 33: 1633–1649.

McKee AR, Houlton BZ, Baraloto C, et al. 2013. Rare species support vulnerable functions in high-diversity ecosystems. PLoS Biology 11: e1001569.

Onoda Y, Saihiga JB, Akutsu K, Aiba SI, Yahara T, Anten NP. 2014. Trade-off between light interception efficiency and light use efficiency: implications for species coexistence in one-sided light competition. Journal of Ecology 102: 167–175.

Peng YF, Chen HYH, Yang YH. 2020. Global pattern and drivers of nitrogen saturation threshold of grassland productivity. Functional Ecology 34: 1979–1990.

Pérez-Harguindeguy N, Diaz S, Garnier E, et al. 2013. New handbook for standardised measurement of plant functional traits worldwide. Australian Journal of Botany 61: 167–234.

R Core Team. 2019. R: a language and environment for statistical computing. Vienna: R Foundation for Statistical Computing.

Ramesh K, Naleeni R, Virendra S. 2007. Leaf area distribution pattern and non-destructive estimation methods of leaf area for Stevia rebaudiana (Bert.) Bertoni. Asian Journal of Plant Sciences 6: 1037–1043.

Raven JA, Lambers H, Smith SE, Westoby M. 2018. Costs of acquiring phosphorus by vascular land plants: patterns and implications for plant coexistence. New Phytologist 217: 1420–1427.

Sánchez-Castillo PM, Linares-Cuesta JE, Fernández-Moreno D. 2008. Changes in epilithic diatom assemblages in a Mediterranean high mountain lake (Laguna de la Caldera, Sierra Nevada, Spain) after a period of drought. Journal of Limnology 67: 49–55.

Shen JB, Yuan LX, Zhang JL, et al. 2011. Phosphorus dynamics: from soil to plant. Plant Physiology 156: 1007–1009.

Smith SE, Jakobsen I, Grønlund M, Smith FA. 2011. Roles of arbuscular mycorrhizas in plant phosphorus nutrition: interactions between pathways of phosphorus uptake in arbuscular mycorrhizal roots have important implications for understanding and manipulating plant phosphorus acquisition. Plant Physiology 156: 1050–1057.

Tamm T, Fensholt R, Guir G, et al. 2015. Ecosystem properties of semiarid savanna grassland in West Africa and its relationship with environmental variability. Global Change Biology 21: 250–264.
Tian QY, Liu NN, Bai WM, et al. 2016. A novel soil manganese mechanism drives plant species loss with increased nitrogen deposition in a temperate steppe. *Ecology* 97: 65–74.

Tian QY, Yang LY, Ma PF, et al. 2020. Below-ground-mediated and phase-dependent processes drive nitrogen-evoked community changes in grasslands. *Journal of Ecology* 108: 1874–1887.

Tiessen H, Moir JO. 1993. Characterization of available P by sequential extraction. In: Carter MR, Gregorich EG, eds. *Soil sampling and methods of analysis*. Ann Arbor: Lewis Publishers, 75–86.

Tilman D. 1987. Secondary succession and the pattern of plant dominance along experimental nitrogen gradients. *Ecological Monographs* 57: 189–214.

Treseder KK. 2013. The extent of mycorrhizal colonization of roots and its influence on plant growth and phosphorus content. *Plant and Soil* 371: 1–13.

Vance CP, Uhde-Stone C, Allan DL. 2003. Phosphorus acquisition and use: critical adaptations by plants for securing a nonrenewable resource. *New Phytologist* 157: 423–447.

Vergutz L, Manzoni S, Pregitzer KS, Jackson RB. 2012. Global resorption efficiencies and concentrations of carbon and nutrients in leaves of terrestrial plants. *Ecological Monographs* 82: 205–220.

Wang JS, Song B, Ma FF, et al. 2019. Nitrogen addition reduces soil respiration but increases the relative contribution of heterotrophic component in an alpine meadow. *Functional Ecology* 33: 2239–2253.

Wright IJ, Reich PB, Westoby M, et al. 2004. The worldwide leaf economics spectrum. *Nature* 428: 821–827.

Xia MX, Guo DL, Pregitzer KS. 2010. Ephemeral root modules in Fraxinus mandshurica. *New Phytologist* 188: 1065–1074.

Xiao Y, Liu X, Zhang L, Song ZP, Zhou SR. 2021. The allometry of plant height explains species loss under nitrogen addition. *Ecology Letters* 24: 553–562.

Ye BB, Chu ZS, Wu AP, et al. 2018. Optimum water depth ranges of dominant submerged macrophytes in a natural freshwater lake. *PLoS ONE* 13: e0193176.

Yu RP, Li XX, Xiao ZH, Lambers H, Li L. 2020. Phosphorus facilitation and covariation of root traits in steppe species. *New Phytologist* 226: 1285–1298.

Zhang DY, Peng YF, Li F, et al. 2019. Trait identity and functional diversity co-drive response of ecosystem productivity to nitrogen enrichment. *Journal of Ecology* 107: 2402–2414.