

Vegetation Patterns and Competitive Dynamics along Elevation Gradients: Interactions between Environmental Factors and Vegetation in the Central Himalayas

Prashant Paudel¹, Stefan Olin², Mark Tjoelker¹, Mikael Pontarp³, Daniel Metcalfe⁴, and Benjamin Smith^{1,2}

¹ Hawkesbury Institute for the Environment, Western Sydney University, Penrith, NSW, Australia

² Department of Physical Geography and Ecosystem Science, Lund University, Lund, Sweden

³ Department of Biology, Lund University, Lund, Sweden

⁴ Department of Ecology and Environment Science, Umeå University

Corresponding Author: Benjamin Smith (ben.smith@nateko.lu.se)

1. Abstract

Elevation gradients are generally characterized by a steady reduction in temperature with altitude, ~~potentially leading to strong variation in zonation of vegetation structure with resulting in differences in growth conditions that often produce clear patterns of vegetation zonation along the gradient.~~ A south-north transect ~~of across~~ the central Himalayas spans from a tropical ~~climate in the south~~ to alpine ~~conditions climates in the north~~, offering an opportunity to investigate the relative roles of abiotic stress and competitive interactions in shaping plant community assembly. We hypothesise ~~a shift from that~~ vegetation composition and productivity ~~shift from being characterised by competition-driven being characterised by~~ realised niches, ~~defined by competitive interactions,~~ at lower elevations, to ~~stress-driven~~ physiological niches, ~~shaped by stress (freezing temperature)~~ at higher elevations. To investigate how these niche transitions influence community ~~dynamics assembly~~ and ecosystem processes, we used a dynamic vegetation model with regional plant functional types (PFTs) parameterised with trait data, including allometric relationships. The model ~~effectively~~ captured spatial ~~and temporal variability patterns~~ in vegetation structure and productivity along the gradient, ~~with simulated patterns closely matching observed vegetation zonation across the transect. The establishment and performance of the PFTs' establishment and performance were dependent~~ depended on their climatic niche and the local ~~competitive interaction abundance of competing PFTs,~~ with persistence shaped by specific ~~functional traits and adaptive strategies.~~ At low elevations, where competitive interactions

dominate, tropical shade-intolerant raingreen and tropical shade-tolerant evergreen PFTs dominated ~~carbon above-ground biomass~~ production and vegetation cover. In contrast, shorter stature, evergreen ~~phenology~~, and cold-tolerant ~~types~~ PFTs were favoured at high elevations, reflecting reduced ~~interspecific~~ competition and ~~physiological adaptation to low~~ increasing temperature ~~limitation~~ stress. Along the ~~elevation gradient~~, PFT functional diversity declined with elevation, while ~~compositional~~ but evenness in composition increased, with evidence of a mid-elevation diversity peak after accounting for stochasticity. Conversely ~~Despite~~, low-elevation communities supported higher functional diversity at low elevations, yet vegetation structure and function ~~characterised~~ reflected by in leaf-area index (LAI), foliar projective cover (FPC) (e.g., LAI, FPC, and ~~carbon above-ground biomass~~) were dominated by a few competitively superior PFTs. ~~We conclude~~ Overall, these results indicate that ~~our model predictions are consistent with that~~ vegetation dynamics along the ~~temperature~~ elevation gradient ~~being~~ are governed by a trade-off between competitive ability and stress tolerance. ~~This trade-off drives as reflected in~~ leads to shifts in structure, composition, and productivity along the gradient, reflecting a transition from realised to ~~physiological niche dominance mediated by environmental conditions and trait responses, which are shaped by environmental conditions, functional traits, and adaptive strategies of the vegetation communities along the gradient.~~

Keywords: Elevation gradient, temperature, competition, community composition, Himalayas, traits, dynamic vegetation model.

3.2. Introduction

Elevation gradients are characterized by rapid changes in climatic conditions, particularly a decline in temperature with altitude (Zhu et al., 2022). With rising elevation, temperature changes predictably in accordance with the 'lapse rate', creating differences in growth conditions, which may be further ~~exaggerated~~ accentuated by small-scale ~~variations in precipitation, topography, aspect, exposure, geology, soil properties~~ types, and biogeochemical processes governing nutrient cycling and resource availability. ~~These environmental gradients variations in growth conditions act as environmental filters (stress factors) (Asner et al., 2014; De Frenne et al., 2013; Zhu et al.,~~

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

2022), which influence that structure ecological processes and species interactions, resulting in that give rise to emergent eco-evolutionary patterns such as niche differentiation, functional trait distribution, plant strategies, and competitive exclusion (Asner et al., 2014; Muñoz Mazón et al., 2020). Distinct community composition and structure may result, with clear vegetation zonation shaped by the interplay of environmental filtering and competition interaction along the gradient.

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

Temperature gradients are among the most influential environmental axes shaping vegetation patterns globally. The Himalayas provide a striking example of this, with distinct zonation observed from the tropical lowlands in the south to the alpine conditions in the north. The strong climatic contrast, i.e. cooler and wetter conditions at higher elevations, and hot, with seasonally dry conditions at low elevations, influences vegetation fitness and community compositions, and distribution of functional traits, play a significant role in shaping the composition and survival of plant species (Måren et al., 2015). Vegetation at higher elevations often exhibits thicker leaves and frost tolerance adaptations such as reduced leaf area, increased epidermal thickness, and increased antifreeze proteins (Satyakam et al., 2022). In contrast, lower elevation species have larger leaves with higher specific leaf area (SLA), larger and wider branching patterns, and fast growth rates to take advantage of abundant resources and warm temperatures (Shah et al., 2019; Sigdel et al., 2022). Additionally, In addition, biotic interactions, particularly competition for light, space, and soil resources also drives vegetation function, diversity (species richness and evenness), and species composition along the elevation gradient (Maharjan et al., 2021; Thakur & Chawla, 2019). However, the relative strength of biotic interactions and abiotic constraints varies along environmental gradients and remains difficult to quantify in complex mountain systems.

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

, creating a biotic filter that results in different vegetation zones exhibiting different community composition, and plant diversity (species richness and evenness). The combined effects of these abiotic and biotic filters on vegetation across elevations are reflected in differences shifts in functional traits, growth strategies of vegetation, and community composition at different positions along the gradient (Midolo et al., 2019; Silva et al., 2023). As a result, Vegetation at higher elevations often exhibits thicker leaves and frost tolerance adaptations such as reduced leaf area, increased epidermal thickness, and increased antifreeze proteins (Satyakam et al., 2022). In contrast, lower elevation

species have larger leaves with higher specific leaf area (SLA), larger and wider branching patterns, and fast growth rates to take advantage of abundant resources and warm temperatures (Shah et al., 2019; Sigdel et al., 2022).

Research on community assembly and vegetation zonation in the Himalayas has largely been descriptive, based on empirical field observations (Acharya et al., 2011; Dhakal, 2018; Shah et al., 2019; Sigdel et al., 2022; Thakur and Chawla, 2019; Thorne et al., 2022) and existing research descriptive interpretation of elevational patterns (Ayer et al., 2025; Drollinger et al., 2017; Dyola et al., 2022; Liang et al., 2020; Maharjan et al., 2021; Maletha et al., 2022; Sigdel et al., 2020). Studies have in the Himalayas have largely remained descriptive, focusing on species distribution, occurrence, morphological traits distributions and their relationships with existing environment. However, No currently available studies a synthetic framework that explains community assembly and ecosystem dynamics across the Himalayan bioclimatic range, where traits and plant strategies drive considering both abiotic and biotic influences on species distribution patterns, remains lacking, limiting our understanding of how competition and abiotic stress interact along the elevation gradient, particularly in driving the transition from realised to physiological niches and how species traits mediate these shifts.

(Ayer et al., 2025; Thakur and Chawla, 2019) Despite these clear vegetation patterns, the relative contributions of trait-mediated competitive interactions and adaptation to stress on vegetation structure, composition, and productivity remain poorly understood along the Himalayas' elevation gradient.

The complex interaction between growth conditions and vegetation functional traits, and their role in structural dynamics and competitive interactions along this gradient, can be effectively captured by incorporating characteristic plant traits into dynamic vegetation models (DVMs) (De Paula et al., 2021; Sitch et al., 2003; Smith et al., 2014). By encoding the traits and response mechanisms of real-world vegetation for different plant functional types (PFTs) in these models and demonstrating agreement between modelled and observed vegetation structural and functional characteristics along the gradient, we can may disentangle the interaction between environmental conditions and vegetation environmental and community dynamics drivers of ecosystem productivity. DVMs represent vegetation as groups of functionally similar species (represented as PFTs),

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

defined by shared ~~which share similar traits (~~morphological, ~~and~~ ecophysiological ~~straits,~~
and life history ~~characteristics)~~strategies and climatic niches (bioclimatic limits for
establishment or survival). The trait-based parameterisation enables representation of
~~Distinct traits and life history strategies, encoded as PFT parameters, influence their~~
~~performance and interactions in model simulations, reflecting~~ functional trade-offs in the
~~carbon allocation and adaptations of species~~ecological strategies ~~that affect their~~
~~performance when growing under different environmental conditions~~ (Díaz et al., 2016;
Pierce et al., 2013)~~), providing a mechanistic link between plant-level adaptations and~~
regional vegetation patterns.

Formatted: Font color: Red

~~We tested how plant strategies reflected in their traits, competition, and interactions with~~
~~environmental factors collectively shape vegetation structure, composition, and~~
~~productivity along the Himalayas elevation gradient.~~ We employed an individual-based
dynamic vegetation model, LPJ-GUESS (Smith et al., 2001, 2014), to simulate structural,
compositional, and functional variability along a south-north transect of the central
Himalayas spanning from a tropical ~~seasonal climate in the lowlands~~ to alpine conditions.
~~Using~~ ~~Our approach leverages~~ empirical data on vegetation traits and life history
strategies ~~of regional PFTs, we evaluated how abiotic stress and competitive interactions~~
~~jointly shape plant community assembly, vegetation structure, and productivity, along the~~
~~same gradient. The simulated PFTs were defined based on empirical data on vegetation~~
~~traits and life history strategies for major species and taxa found along the Himalayan~~
~~transect. Through simulations with this model, incorporating representations of key~~
~~mechanisms and trait data from the Himalayan flora,~~Here, ~~Specifically,~~ we teste the
hypothesis that community assembly shifts from a competition-driven regime at lower
elevations (realised niche dominance) to a stress-driven regime at higher elevations
(physiological niche dominance). By comparing simulated outcomes with field
observations, we assess how well these niche transitions explain spatial patterns in
vegetation structure, composition and productivity across one of the world's most
complex mountain ecosystems.~~By comparing simulated outcomes with field~~
~~observations, we evaluate how well the mechanisms embedded in the model explain~~
~~spatial patterns in structure, composition, and function variability. We also examine~~
~~species richness and evenness as emergent properties of functional composition,~~
~~reflecting how changing growth conditions influence diversity. Overall, this study~~

Formatted: Font color: Red

~~illustrates the complex interplay between growth conditions, traits, and competitive dynamics, thereby enhancing our understanding of the ecological processes that shape plant communities along the elevation gradient of the central Himalayas. We further evaluate species richness and evenness as emergent properties of functional composition, providing insight into how changing growth conditions influence biodiversity patterns along the Himalayan elevation gradient.~~

3. Methods

3.1 Study site

The study focuses on a species-rich elevation gradient of the central Himalayas (Figure 1). Along the gradient, temperature decreases by approximately 6.5 °C per vertical kilometer, reflecting the standard lapse rate, where the average annual temperature in the tropical zone is 28 °C and around 10 °C in the alpine region (MoFSC, 2016; Poudel et al., 2020). Nearly 80% of the total annual rainfall occurs during the monsoon (June to September) (Maharjan et al., 2021), where average annual rainfall ranges from 165 mm (northern end of the gradient, i.e., Trans-Himalayan region) to 5244 mm (1550-2000 m altitude) in the lower and middle part of the gradient (Luitel et al., 2020; Poudel et al., 2020). ~~Generally, precipitation reaches a maximum at mid-elevations (approximately 1000-2000 m) and declines rapidly at higher~~~~Precipitation peaks at around 1000 m altitude asl and then rapidly decreases with an increase in elevation~~ (Luitel et al., 2020; Maharjan et al., 2021; Poudel et al., 2020). Vegetation follows the temperature patterns ranging from tropical (24 °C) to temperate forests and to colder sub-alpine vegetation (6.9 °C) (Shrestha et al., 2015), where growth condition varies significantly due to variations in temperature and precipitation patterns. Within a horizontal span of 100 km along the gradient, 160 different tree species were recorded in a plot-level survey conducted in forest areas, where vegetation composition and dominance changes with elevation (DFRS, 2015; Pokhrel and Sherpa, 2020).

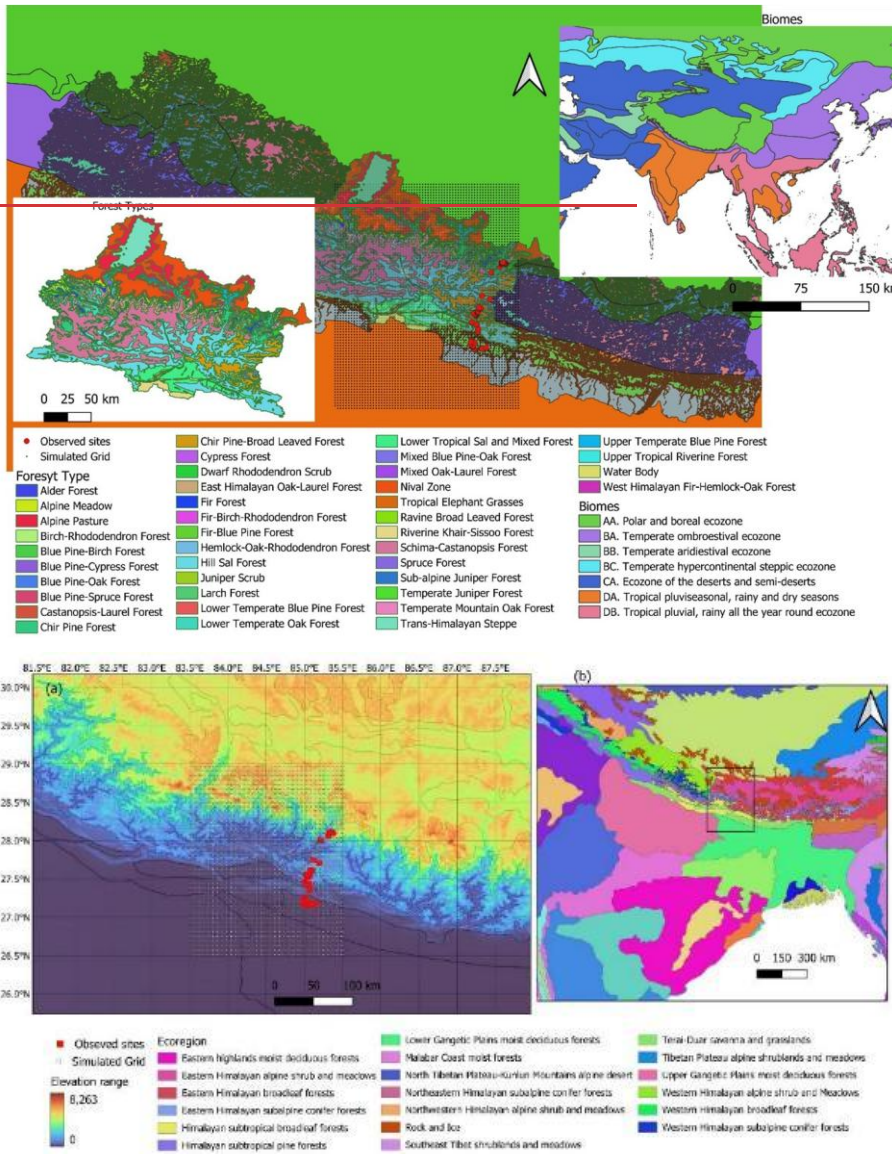


Figure 1: Map of the study area with major global Biomes ecoregion (data source: Olson et al., 2001), elevation range Forest types (data source: Earth Resources Observation and Science (EROS) Center, 2017 (Land Resource Mapping Project, 1986) (<https://rds.icimod.org/>)), trait data observed sites (Maharjan et al., 2021), and

203 simulated grids along the elevation gradient of the central Himalayas [\[Supplementary](#)
204 [Figure 1 shows the spatial distribution of forest types along the gradient\]](#).
205

206 3.2 Vegetation model description and customization

207 The Lund-Potsdam-Jena General Ecosystem Simulator (LPJ-GUESS) is a process-based
208 dynamic vegetation model used ~~to for simulating~~[simulate and predicting](#) ecosystem
209 responses to environmental changes at ~~the~~ regional or global level based on local,
210 neighbourhood-scale interactions among simulated plants (Smith et al., 2001, 2014). It
211 represents generalized ecophysiological processes such as photosynthesis, autotrophic
212 (plant) and heterotrophic (soil) respiration, carbon, water, and nitrogen cycling. The
213 model adopts gap dynamics theory (Bugmann et al., 1996; Scherstjanoi et al., 2014) to
214 simulate tree population dynamics ~~emerging from the balance of~~[through](#) plant
215 establishment, growth, and mortality (De Paula et al., 2021; Sitch et al., 2003; Smith et al.,
216 2001). The model is applied across a continuous geographic grid. Vegetation in each grid
217 cell is represented as a mixture of PFTs whose ~~potential~~ distribution ~~in climate space~~ is
218 governed by bioclimatic envelopes for ~~the~~ establishment and survival. Within its
219 bioclimatic envelope, ~~the presence and abundance of a~~ PFT [abundance is further](#)
220 [influenced by](#) ~~are additionally subject to vegetation dynamics determined by the~~
221 interactions between co-occurring individuals ~~of other PFTs and the cascading that~~
222 ~~affects on~~ carbon assimilation and allocation, reproduction, and survival ~~of individuals~~
223 ~~co-occurring~~ within ~~replicate~~ local patches, nominally 0.1 ha in size. The overall
224 vegetation of a grid cell is aggregated across multiple patches (here 15), representing
225 random samples of the wider landscape ~~of the~~[within each](#) grid cell. PFT-specific
226 parameters and growth strategies determine performance under different climates, CO₂
227 concentrations, and stages of vegetation development.
228

229 The model accounts for structural responses to competitive and environmental
230 conditions through adaptive allometric relations (DBH-Height, DBH-Crown Area, DBH-
231 Crown Volume) and a dynamic bole height scheme for each cohort. This allows allometric
232 scaling and carbon allocation to respond dynamically to ~~canopy the competition~~
233 ~~{crowding }~~ conditions defined by the availability of photosynthetically active radiation
234 ~~(Paudel et al., 2026)(Paudel et al., Unpublished)~~. [The model simulates leaf area index](#)

Formatted: Font color: Red

Formatted: Font color: Red

(LAI) from leaf carbon pool using PFT-specific ~~specific leaf areas~~ SLA (prescribed as a PFTs parameter), emerging from carbon allocation, phenology and turnover processes.

To represent the diversity of vegetation composition along the Himalayan elevation gradient, we modified the following features to customize the model for application to our study area.

- The default model has 12 PFTs, defined to represent the dominant biomes of the world. For this study, we defined a new set of regional tree PFTs tailored to local conditions using a multivariate hybrid (hierarchical) clustering approach (Ssee Section 3.3 and Supplementary Materials for the details of the clustering approach and derived PFTs). These PFTs represented major ~~hypothesised vegetation strategies and adaptation mechanisms employed by vegetation to~~ coping with competition (biotic stress) and harsh climatic conditions (abiotic stress) ~~along the elevation across the~~ gradient. Tree PFT parameters and their derivation are further discussed in Section 3.3 below. Both C₃ and C₄ grass PFTs with default parameter values (defined for the global level) were retained for simulations (Peng et al., 2024), representing distinct photosynthetic pathways and associated physiological differences that influence their relative abundance along the elevation gradient.
-
- Bioclimatic limits (mean minimum and maximum temperature for the 20 years of coldest month for establishment and survival; mean minimum warmest month temperature for establishment) control each PFT's establishment and survival in a given grid cell. The model defines these limits for global biomes ranging from tropical to boreal ecosystems. For this study, four climatic limits are defined: tropical, subtropical, temperate, and alpine. Climatic ranges ~~for our study~~ were adopted from Jackson (1994) and modified based on the ranges recorded by Maharjan et al. (2021).

3.3 Data Sources for model input and parameterisation

The CRU-JRA (v2.4.5d) global gridded climate dataset was downscaled to a 3 km spatial resolution using bicubic interpolation (Latombe et al., 2018) and used as climate-forcing data for our model simulations. Using the method, we This method increases spatial

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: English (United States)

Formatted: Indent: Left: 0.5", No bullets or

Formatted: Font color: Red

Formatted: Font color: Red

268 resolution but does not explicitly account for topographic effects such as lapse-rate-
269 driven elevation and associated terrain condition. The CRU-JRA is a gridded daily dataset
270 with 0.5° × 0.5° spatial resolution from 2001 to 2022 (Araghi and Martinez, 2024). We
271 used monthly mean air temperature, precipitation, wind speed, incoming solar radiation,
272 specific humidity, number of wet days, and minimum and maximum temperature as
273 inputs. The number of wet days are defined as a days with non-zero precipitation (>
274 0 mm) and monthly counts are obtained using daily precipitation data for model
275 simulation. All variables except for precipitation were interpolated to daily values; for
276 precipitation, the monthly sum was divided equally across the number of wet days per
277 month. Soil properties and the atmospheric nitrogen deposition rates (Lamarque et al.,
278 2013) were configured using 0.5° × 0.5° spatial resolution. Annual atmospheric CO₂
279 concentration data from NOAA (1901- 2022) are used as input data (Friedlingstein et al.,
280 2023). Elevation values of each simulated grid, whereas extracted from the GTOPO30
281 dataset provided by the U.S. Geological Survey (Earth Resources Observation and Science
282 (EROS) Center, 2017), with a horizontal resolution of 30 arc-seconds (approx. 1 km),
283
284 Trait values of the 31 most abundant tree species - based on the frequency of observation
285 and total carbon contribution identified by Maharjan et al. (2021) were compiled from
286 Maharjan et al. (2021), Jackson (1994), and Thakur and Phulara (2014). Tree allometry
287 data (DBH, total tree height, crown radius, crown height) were compiled from the Tallo
288 database (a global tree allometry and crown architecture database) (Jucker et al., 2022)
289 and BAAAD (a biomass and allometry database for woody plants) (Falster et al., 2015).
290 Elevation data across the simulated grid whereas extracted from Earth Resources
291 Observation and Science (EROS) Center (2000) and whereas used for plotting simulated
292 outputs. (Supplementary Figure 2 shows the patterns of environmental variables along
293 the gradient are).
294
295 A divisive hierarchical clustering approach was used to group tree species into distinct
296 PFTs based on similarities in traits and life-history strategies. Both phenological and
297 morphological characteristics traits were considered. Species were first stratified into
298 four temperature-defined groups (tropical, subtropical, temperate, and alpine) based on
299 their dominant climatic distribution (Jackson, 1994; Thakur and Phulara, 2014), ensuring
300 that broad-scale bioclimatic filtering were accounted for prior to clustering. Within each

Formatted: Font color: Red
Formatted: Font color: Red
Formatted: Font color: Red
Formatted: Font color: Red
Formatted: Font color: Red
Formatted: Font color: Red
Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red
Formatted: Font color: Red
Formatted: Font color: Red

Formatted: Font color: Red
Formatted: Font color: Red

Formatted: Font color: Red
Formatted: Font color: Red

301 temperature stratum, hierarchical clustering using Ward's method (Murtagh and
302 Legendre, 2014). was applied to standardized trait data, including specific leaf area (SLA),
303 wood density, leaf phenology, and leaf longevity, to group species into functionally similar
304 clusters. The clustering further incorporated ecological strategy variables, including life-
305 history strategy (fast vs. slow growing; early vs. late successional), shade tolerance
306 (tolerant, intermediate, intolerant), and drought resistance (tolerant vs. sensitive),
307 alongside structural traits such as maximum tree height. Species were progressively
308 grouped based on multivariate similarity in these traits, resulting in 13 distinct functional
309 clusters, including three PFTs within the conifer group (see Supplementary for further
310 details of the clustering approach and derived PFTs). ~~We clustered tree species into 13~~
311 ~~functional groups based on their dominant temperature range (tropical, sub-tropical,~~
312 ~~temperate or alpine), leaf phenology (evergreen, raingreen, summergreen, broadleaved,~~
313 ~~and conifers), life history strategies (growing fast and slow, late and early successional~~
314 ~~species), shade requirement (shade tolerant, intermediate tolerant and intolerant);~~
315 ~~drought resistance class (drought tolerance, drought sensitive), as well as traits such as~~
316 ~~wood density and total tree height (Supplementary Table 1).~~ The resultant PFTs are
317 designed to represent the functional diversity required to simulate structure,
318 composition, and productivity along the elevation gradient.

319
320 The following parameters were updated for each tree PFT: leaf phenology, drought
321 tolerance, wood density, SLA, shade tolerance, leaf longevity, and leaf turnover rate (Table
322 1). The values of these parameters, compiled from the sources mentioned above, were
323 averaged across species included in the PFTs emerging from the clustering procedure
324 described above. In the model, shade tolerance is linked to life history strategies, which
325 influence growth, reproduction, and survival of PFTs across different light environments.
326 Here, parameters and their values (Supplementary Table 2), as defined by Hickler et al.
327 (2004), were adapted in model simulation to represent these dynamics. Similarly,
328 quantile regression was used to estimate allometric scaling parameters (DBH relative to
329 height, DBH relative to crown area, and DBH relative to crown volume) under three
330 different stand crowding conditions (5%, 50% and 95%), allowing us to evaluate
331 structural response to competition and their influence on ecosystem dynamics.

Table 1: Tree PFTs and parameter values used for simulation, including bioclimatic limits, Shade tolerance parameters and their values, and allometric relations (see Supplementary Table 1 & 2 for details).

PFTs	Parameters					
	Leaf phenology	Leaf longevity (years)	Shade tolerance	Wood density (kgC m ⁻³)	SLA (m ² kgC ⁻¹)	Leaf Turnover (fraction/year)
Alpine broadleaved evergreen (ABE)	evergreen	2	intolerant	273	13	0.5
Alpine needleleaved (ANE)	evergreen	3	tolerant	280	9	0.33
Temperate summergreen (TeBSG)	summergreen	0.5	intolerant	265	18	1
Temperate broadleaved shade tolerant evergreen (TeBEt)	evergreen	2	Tolerant	281	15	0.5
Temperate broadleaved shade intolerant evergreen (TeIBE)	evergreen	2	intolerant	282	15	0.5
Temperate needleleaved (TeNE)	evergreen	3	intolerant	235	18	0.33
Sub-tropical needleleaved (STrNE)	evergreen	3	intolerant	210	10	0.33
Sub-tropical rain green (STrRG)	raingreen	0.5	intolerant	200	25	1
Sub-tropical shade-tolerant evergreen (STrIBE)	evergreen	1	intermediate	265	20.85	1
Sub-tropical broadleaved evergreen (STrBE)	evergreen	2	tolerant	145	17	0.5
Tropical broadleaved shade-tolerant (TrBE)	evergreen	2	tolerant	290	22	0.5
Tropical Summergreen (TrBSG)	summergreen	0.5	intolerant	247	21	1
Tropical broadleaved raingreen (TrBRG)	raingreen	1	intolerant	245	18	1

Data source: (Jackson, 1994; Maharjan et al., 2021; Thakur and Phulara, 2014).

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

3.4 Simulation protocol and model validation

Using the aforementioned forcing data and PFTs parameterized with traits, the model was run with 15 patches in each grid cell of 1000 m², simulating the period from 1901 to

2022. A spin-up of 500 years, recycling the first 30 years of the observed climate data set, was performed to achieve an initial steady state for vegetation structure. We ran LPJ-GUESS in cohort mode (Smith et al., 2001, 2014), using the BLAZE fire model to account for the impacts of weather-related fire disturbances on vegetation structure (Rabin et al., 2017) and applied a generic return interval of 100 years for patch-destroying disturbances, following Pugh et al. (2019).

We implemented a neighbour removal experiment in the model to assess the effects of competitive neighbour individuals and PFTs on the performance of the selected PFTs (Monteux et al., 2024). Individuals of all other woody PFTs except the PFT of interest were removed from the simulation after model year 1950. C₃ grass or C₄ grass forming the understory of the woody stand was retained. However, to allow the ecosystem to re-equilibrate and account for the effects caused by removing neighbours, we allowed target PFTs to grow for another 50 years (i.e., model year 2000) until the ecosystem recovered and productivity stabilized according to the prevailing environmental conditions.

Gross primary productivity (GPP) data from We used Bi & Zhou (2022), (0.05° × 0.05° spatial resolution)'s, gross primary productivity (GPP), produced using the leaf light use efficiency model from 2010 to 2020 (DRYAD), were compared with simulated GPP from 2010 to 2020 along the elevation gradient, focusing on the evaluation of variation in GPP across elevations despite differences in spatial resolution. Plot level above-ground carbon biomass data by Khanal & Boer (2023), estimated from forested area in the national forest inventory, was compared with simulated patch-level carbon above-ground biomass from the model to validate the broader patterns in above-ground biomass distribution across the ecosystem regardless of land use conditions. Similarly, the bole height (height up to the first branch) measured and elevation range recorded by Maharjan et al. (2021) along the studied gradient were compared with simulated values to evaluate the model's ability to capture structural and compositional variability along the elevation gradient. Although these observed values predominantly represent dominant species along the gradient, they were treated as independent observational datasets for evaluating model performance in representing structural variability. Additionally, to further validate the structural component, MODIS derived LAI of 1 km resolution from 2000 to 2022 (Myneni et al., 2015), was compared with simulated monthly LAI, with model evaluation focusing

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

on elevation-dependent patterns given differences in spatial resolution between the observational and simulated datasets.

3.5 Competition index and rank abundance curve (RAC) of FPC for evenness

An index of competition was calculated for each PFT to quantify the effects of neighbour removal on the performance of the target PFT. The target PFT's performance was evaluated using simulated carbon mass production from 2000 to 2020 with and without competition. The competition index (CI) was calculated using the index matrices approach of Avolio et al. (2019) and Brooker & Kikvidze (2008) with modifications, rather than using empirically derived coefficients from species-removal experiments. We modified the equation by Avolio et al. (2019) and Brooker & Kikvidze (2008) to more intuitively represent each PFT's optimum competitive capacity relative to potentially co-occurring woody PFTs. This approach allows the CI to represent the relative competitive capacity of each PFT under different competition conditions. A CI value of 1 indicates no effect from competitors, while a value close to 0 indicates a significant impact from the competitor's presence.

$$CI_i = [1 - \frac{C_{mass-Ni} - C_{mass+Ni}}{\text{Max}(C_{mass-Ni}, C_{mass+Ni})}] \dots\dots\dots (\text{Equ i})$$

where CI is the competition index in year i, C_{mass-N} and C_{mass+N} are the carbon masses of the target PFTs in the presence (+N) and absence (-N) of a competitor in year i.

The rank abundance curve (RAC) of relative foliar projective cover (FPC) of each tree PFT present in a grid cell was used to quantify species-PFTs evenness, defined as similarity in local abundances among PFTs along the elevation gradient. PFTs level evenness and richness were calculated using FPC, we are used as indicators of dominant functional strategies and competitive hierarchy shaping ecosystem structure and carbon dynamics across the elevation gradient. Following Avolio et al. (2019), Smith & Wilson (1996) and Whittaker (1965), the natural logarithm of relative FPC was plotted against the inverse of PFT rank, and a regression between rank and log FPC was used to calculate the slope of the curve in each grid point using simulated FPC from 2000 to 2022. The slope of the RAC was used as a robust (independent of species richness) measure of evenness by Smith & Wilson (1996). A steeper slope would indicate greater dominance by one or a few PFTs

Formatted: Font color: Red

Formatted: Font color: Red

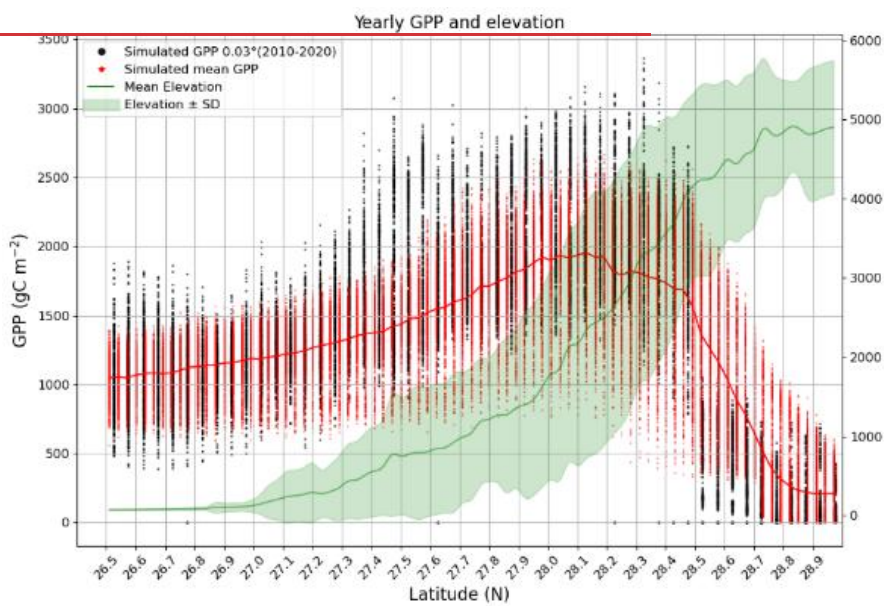
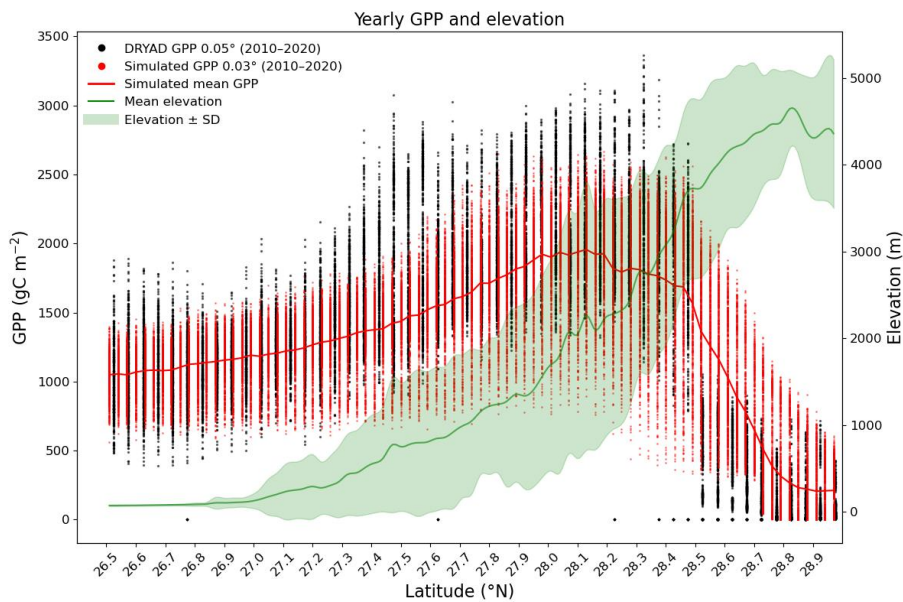
Formatted: Font color: Red

relative to other co-occurring PFTs, while a flatter slope indicates greater evenness. Evenness was plotted against elevation to examine the patterns of change in evenness in relation to changes in growth conditions. Similarly, the number of PFTs in each latitude band was used as an indicator of functional richness, representing the diversity of ecological strategies. We hypothesise that ~~with an increase in elevation,~~ evenness increases with elevation due to reduced competition, and the abundance of PFTs decreases as under environmental stress ~~limits species adaptations in~~ higher elevations.

4. Results

4.1 Spatial variability in productivity and structure along the gradient

Simulated annual GPP showed strong agreement with GPP estimated by Bi and Zhou et al. (2022) using the leaf light use efficiency model, both datasets displaying a gradual increase in productivity up to approximately 2500 meters. ~~In both datasets, at~~ the northern end of the gradient, annual GPP decreased sharply as temperature declined, with some grids having very low productivity (GPP value close to zero) (Figure 2). ~~Monthly GPP. These patterns shows a progressive align with~~ shortening of the productive seasons and a decrease in monthly productivity and with decreasing temperature ~~increasing elevation (Supplementary Figures 3 and 4 for monthly GPP) in alpine zones, characterized by fewer months of productivity reflecting a shift from productive lower elevations to physiological constraints (Supplementary Figures 2 & 3 for monthly GPP)~~ at higher elevations.

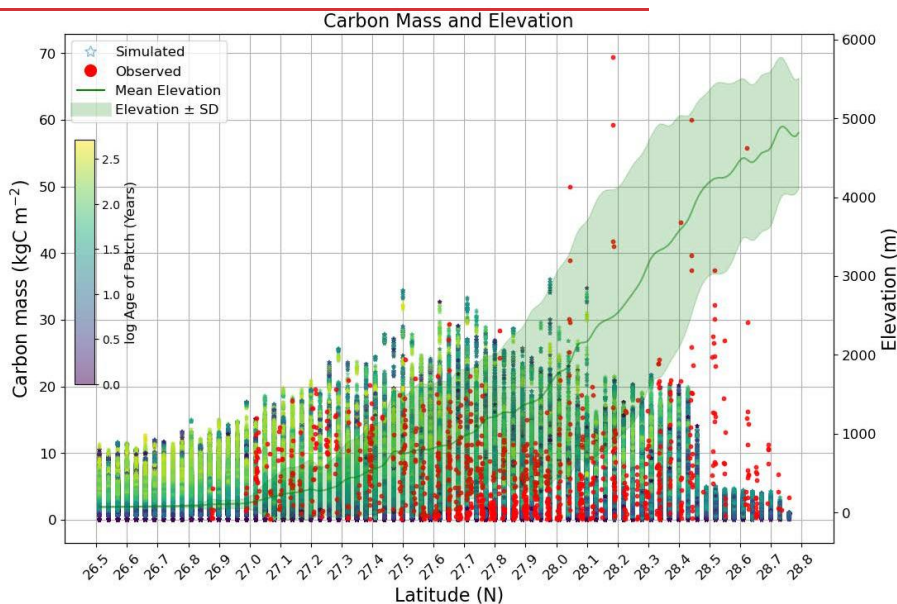


Formatted: Line spacing: 1.5 lines

Figure 2: Simulated annual GPP ($0.03^\circ \times 0.03^\circ$, red point) and annual GPP from the DRYAD database ($0.05^\circ \times 0.05^\circ$, black point) from 2010 to 2020 across grids along the elevation

gradient of the Himalayas along with mean elevation (green line) and elevation range
(greed shaded area) for each latitude.

Simulated above-ground biomass at the patch level closely matched observed plot-level
above-ground biomass in the measured at forest areas across of the gradient, indicating
that the model effectively captures spatial variability in carbon mass distribution. (Figure
3). However, in the northern regions of the gradient, where *Rhododendron* PFTs
dominate, the model underestimated the observed biomass (Figure 3). In contrast,
whereas in the mid-latitude range (27.4 – 28.0 °N), the model it slightly overestimated
above-ground carbon biomass, coinciding in the mid-latitude range (27.4 – 28.0 °N),
corresponding with to areas with of higher PFT abundance. Above-ground biomass
showed a peak at mid-elevations followed by a decline toward higher elevations. This
mid-elevation peak in biomass accumulation, followed by a decline at higher elevation,
suggests increasing physiological constraints under alpine conditions. Notably, the patch-
level above-ground biomass did not show any clear patterns with the simulated age of the
patch. In some simulated grids, young patches exhibited higher above-ground carbon
biomass productivity compared to older ones, suggesting that trait-based responses and
environmental conditions have a stronger influence on productivity than patch age.



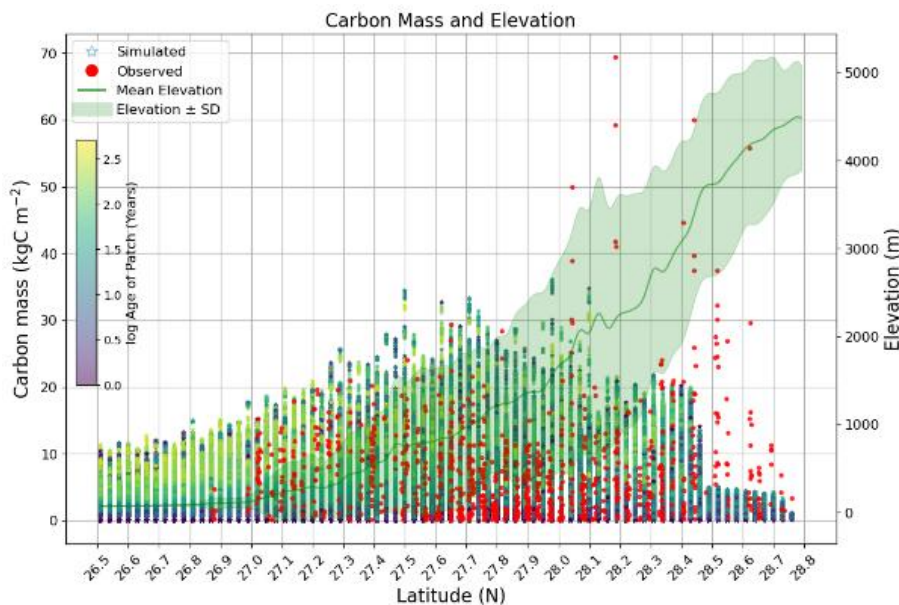
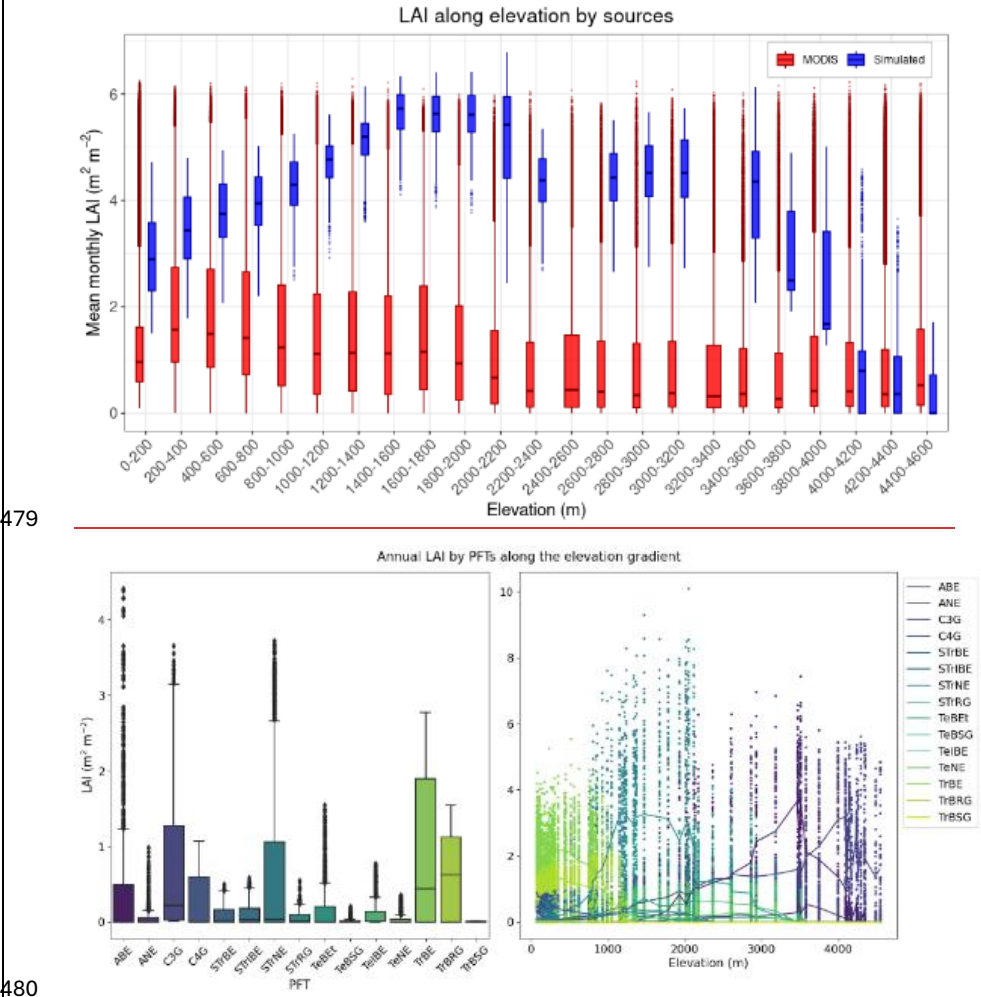


Figure 3: Simulated above-ground carbon-biomass per patch (2010-2015) for tree PFTs with the age of the patch (in years) and observed above-ground carbon-biomass per plot in forest areas along the elevation gradient of the Himalayas.

Similar to productivity and biomass, vegetation structural attributes also varied along the elevation gradient. Both simulated and MODIS LAI shows similar patterns in monthly LAI with substantial overlap between the two datasets (Figure 4- top panel). MODIS LAI shows greater variability in monthly LAI values compared to simulated LAI. Simulated LAI of tree PFTs increased from approximately $5 \text{ m}^2 \text{ m}^{-2}$ at lower elevations to about $8 \text{ m}^2 \text{ m}^{-2}$ at mid-elevations, followed by a gradual decline toward higher elevations (Figure 4 - bottom panel). Elevated LAI values at higher elevations were maintained primarily by evergreen PFTs, whose LAI remained relatively constant throughout the year and by C_3 grasses. Simulated leaf area index (LAI) for tree PFTs increases gradually along the gradient from around $5 \text{ m}^2 \text{ m}^{-2}$ to around $8 \text{ m}^2 \text{ m}^{-2}$, primarily due to the presence of evergreen PFTs at higher elevations, whose LAI remains constant throughout the year. Six PFTs contributed the most to LAI, each with varying zones (elevational range) of dominance. In the southern part of the gradient, two PFTs - tropical broadleaved raingreen and tropical broadleaved evergreen - exhibit higher PFT-specific LAI. Sub-

Formatted: Subscript

474 tropical needle-leaved PFTs form maximum LAI in mid-latitude elevation (~3.2 m² m⁻²),
475 followed by alpine evergreen broadleaved (~ 4 m² m⁻²) in higher elevation. C₄ and C₃
476 grasses contribute significantly to LAI along the gradient, with C₃ grasses being dominant
477 in cold regions, with an LAI of 3.2 m² m⁻² (Figure 4-bottom panel).
478



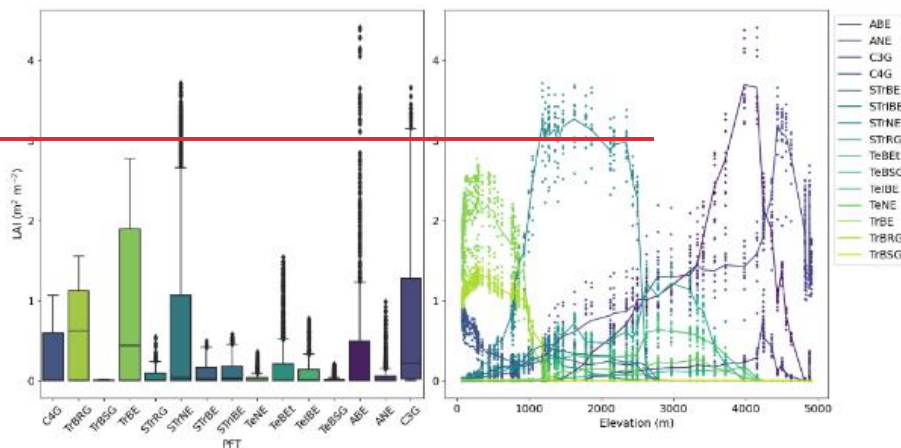


Figure 4: LAI across the elevational gradient. Top panel: comparison of monthly simulated LAI (3 km resolution) and MODIS LAI (1 km resolution) plotted against elevation, with points representing mean monthly LAI values at individual grid. Bottom panel: boxplot of simulated annual LAI by PFTs and line plot of total annual LAI (line represents mean LAI for each PFT and dots representing LAI at simulated grids across latitude) across the elevation gradient by PFTs.

The comparison of simulated (2010-2015) and measured bole heights from Maharjan et al. (2021) shows that both follow the same patterns, although the model exhibits more pronounced variability across most PFTs (Figure 5). It shows that bole height varies with PFTs, with tropical broad-leaved rainforest (TrBRG) having a large bole height, followed by temperate shade-intolerant evergreen (TeIBE) (Figure 5). The overall mean bole height remains similar up to the temperate zone (approximately up to 3500 m), after which it decreases and declines sharply at higher elevation, particularly in the upslope alpine zone, where bole heights remain consistently smaller (Figure 5). This overall pattern of taller bole height at lower elevation likely reflects intensive competition for light, where individuals grow taller to avoid shading. In contrast, the decline in bole height in higher elevations is consistent with both physiological constraints and adaptive strategies that favour short bole height under alpine conditions.

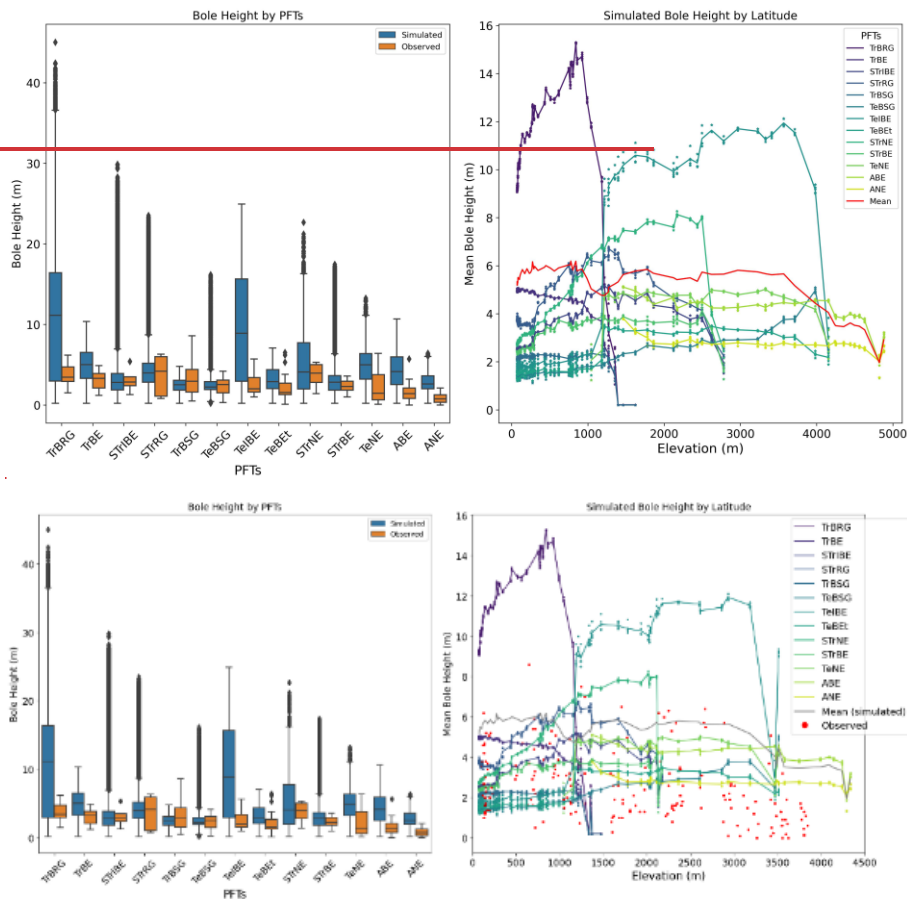


Figure 5: Boxplot of simulated and observed bole height [from Maharjan et. al. \(2021\)](#) (median with ranges and outliers) and bole height for PFTs (dot denoting the bole height of each cohort of PFTs simulated in the grid), [\(grey line\) and observed bole height \(red dots\) across PFTs along the elevation gradient, along with the mean bole height \(red line\) across PFTs along the elevation gradient.](#)

4.3 PFT performance and competition index along the elevation gradient

~~The result indicates shows that the T~~total ecosystem level carbon mass production varies along the gradient and depends on both—the abundance of PFTs' (Figure 6;

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

Supplementary Figure 5). and their location within the elevation range. It shows that
Total carbon mass production at the ecosystem level is highest in mid-lower elevations
(around 1000 m), where multiple PFTs contributed climatic conditions supported a
balanced contribution to ecosystem productivity from including broadleaved evergreen,
deciduous trees, and conifers (Supplementary Figure 54 for total carbon mass production
along the elevation). At lower elevations, tropical broadleaved raingreen PFTs had the
maximum contribution in carbon mass production, whereas temperate shade-intolerant
evergreen PFTs had the maximum carbon mass production in temperate regions. In
colder regions at higher elevations, alpine broadleaved evergreen PFTs (dominated by
Rhododendron species) contribute most to total carbon mass production (Figure 6). In
colder areas, where only two tree PFTs were adaptive within their prescribed bioclimatic
limits, alpine broadleaved evergreens (*Rhododendron* species) have the maximum
contribution to carbon mass production. These findings highlight that PFTs' composition
and their contribution to carbon mass production are driven by their climatic niche
(elevational range) and adaptive capacity to existing climatic and competition conditions
(Figure 6).

Formatted: Font color: Red

Formatted: Font: Italic

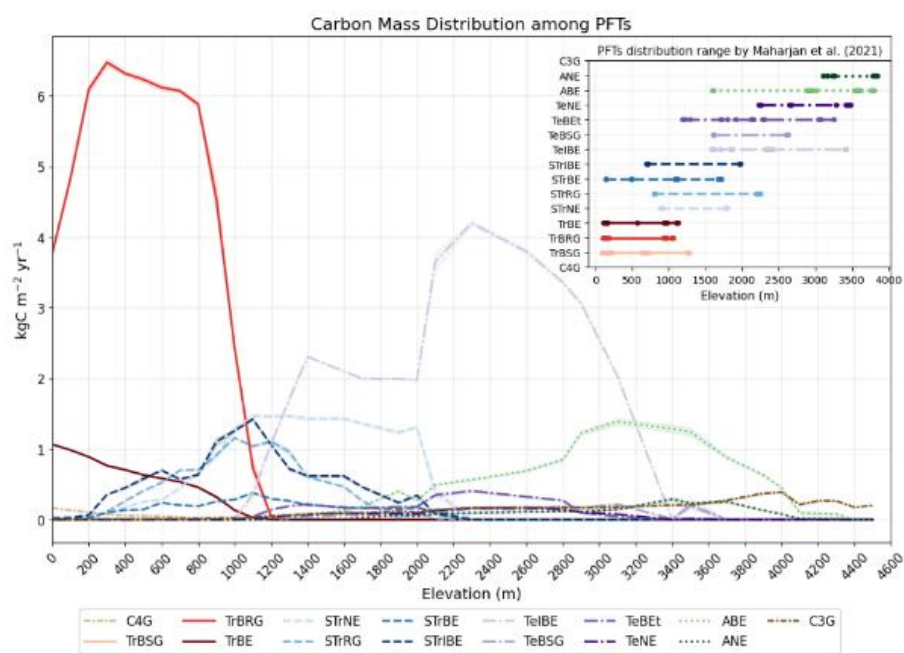
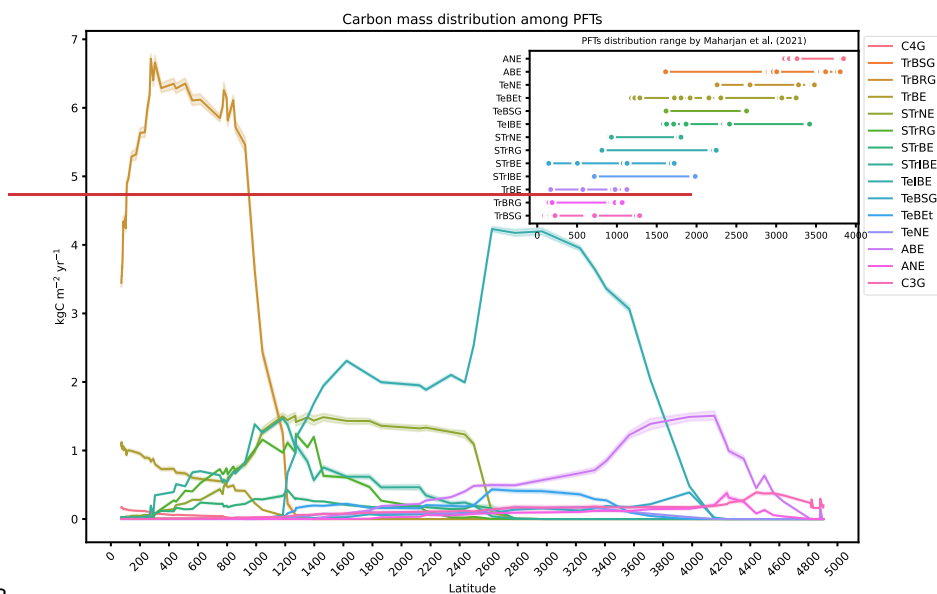
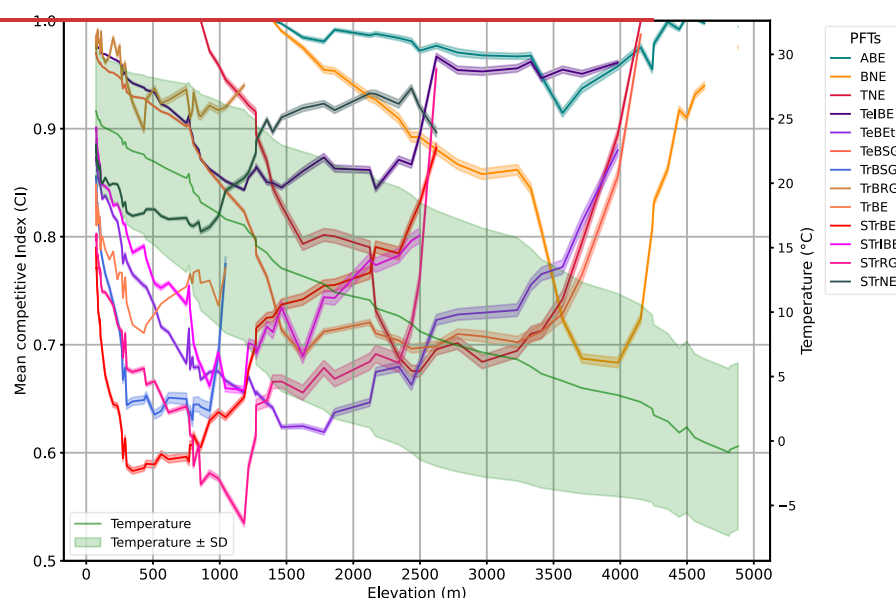


Figure 6: Carbon mass production by ~~each~~ PFT along the ~~latitude~~~~elevation gradient~~ (the inset figure shows the PFTs' distribution ranges recorded by Maharjan et al., (2021) in the ~~elevation~~ gradient).

The mean CI plotted against ~~latitude~~~~elevation~~ shows that removing competitors has ~~set~~ different levels of impact on ~~PFT performance in~~ carbon mass production ~~among PFTs~~ (Figure 7). ~~PFT responses to the absence of competition differ within and outside their dominant climatic ranges.~~~~PFTs' responses without competition differ within and outside their dominant climatic range. In their~~~~Within their~~ dominant growth regions, TrBRG and ABE were least impacted by the presence of neighbours, with CI values over 0.9 (Figure 7). ~~At~~~~the~~ the lower elevation range, the presence of temperate PFTs was more random, and their performance was not enhanced by neighbour removal. For example, the CI values of TelBE, TeBSG, and TeBEt were higher (~0.9), suggesting similar performance with and without competitors (Figure 7). In general, subtropical PFTs ~~highly~~ benefited from competitor removal in the model (Figure 7). There were no clear patterns and associations of PFTs' CI with average annual temperature, as PFTs' performance depends on climatic niche and ~~the~~~~neighbours'~~ presence ~~of neighbours~~ (Figure 7).



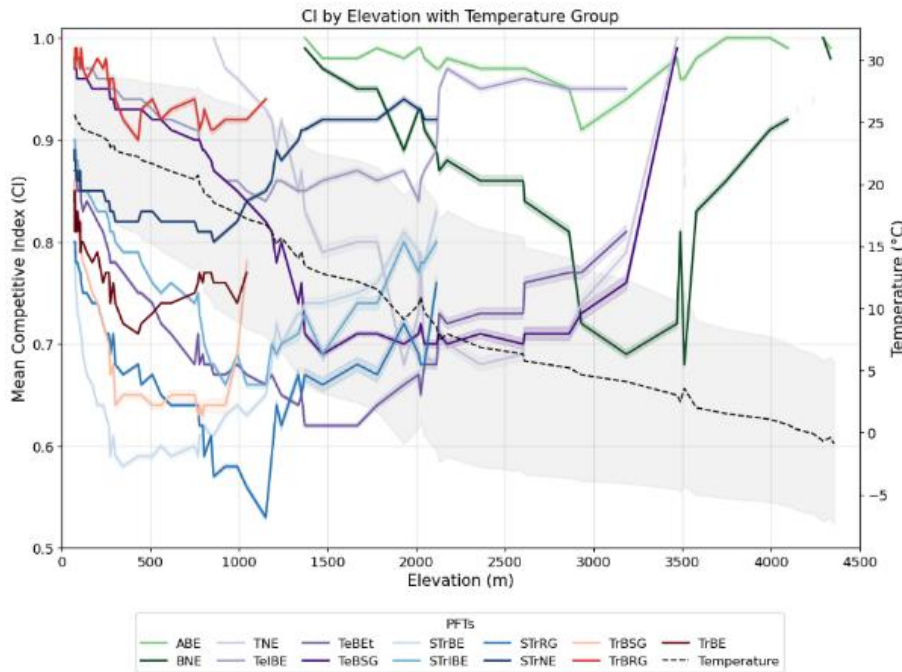


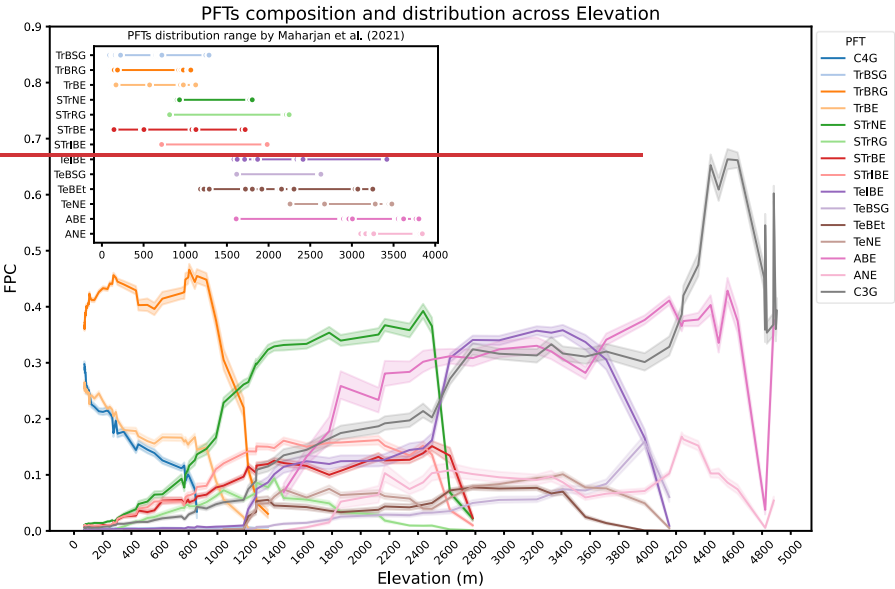
Figure 7: Mean competitive index of PFTs and mean annual temperature (green line) with standard deviation of temperature in the last 30 years (green shaded area) along the elevation gradient.

4.4 Community composition and evenness along the elevation gradient

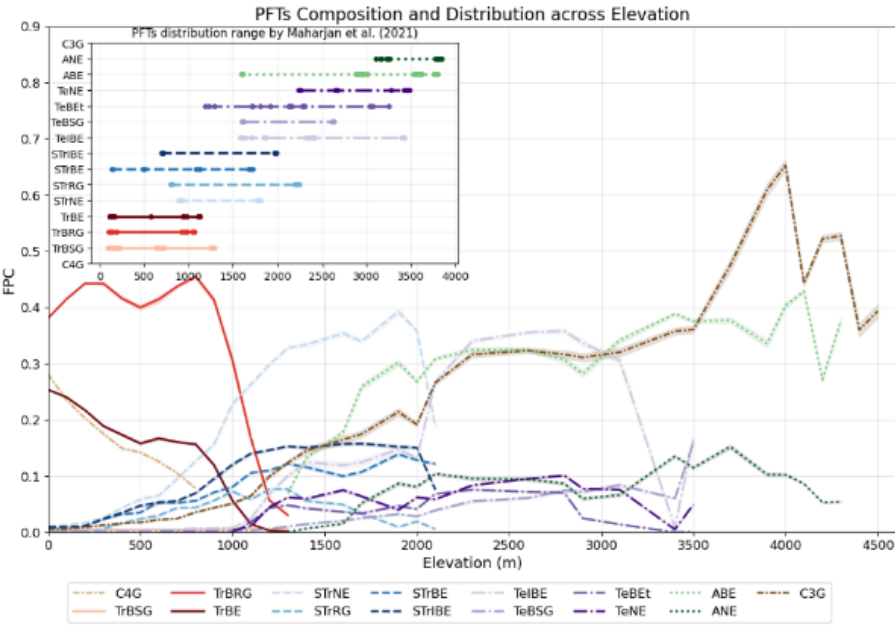
The simulated results show that PFT composition and dominance vary along the change with elevational gradient and associated temperature, where with the total number of PFTs decreases with an increase in elevation. At lower elevations, three PFTs - tropical broadleaved rainforest, tropical broadleaved evergreen, and C₄ grass - dominate despite the presence of over ten PFTs in the area. At the higher end of the gradient, alpine needle-leaved and alpine evergreen broadleaved form tree crown cover, where while C₃ grass dominance increases with elevation, reaching up to 70% in FPC. Sub-tropical needleleaved and temperate shade-intolerant evergreen PFTs dominate the mid-elevation range, showing variability in PFT composition and distribution along the gradient. In the mid-elevation gradient, an area dominated by sub-tropical and temperate

571 PFTs, PFT composition changes more frequently than at the gradient's other ends of the
572 gradient (Figure 8).

573



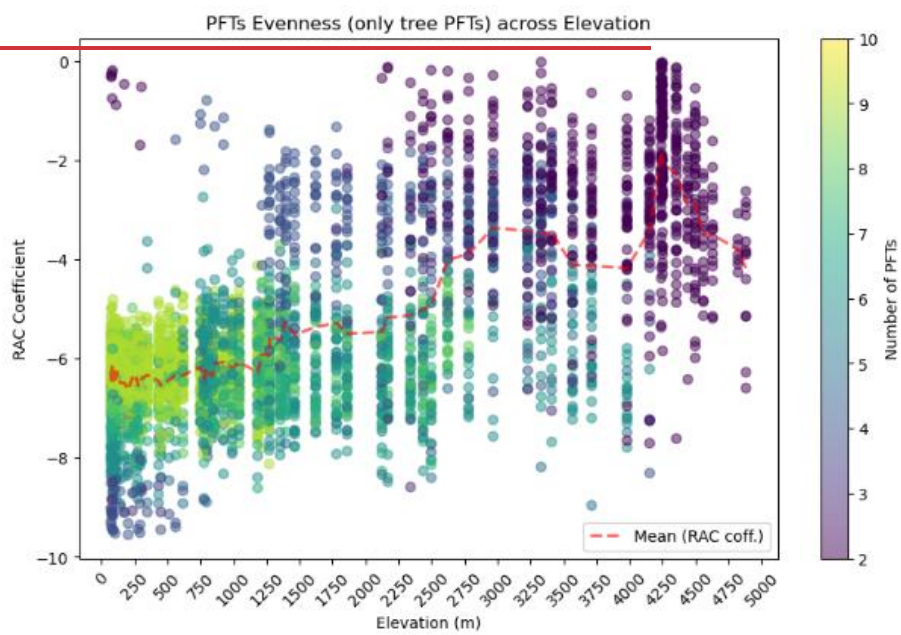
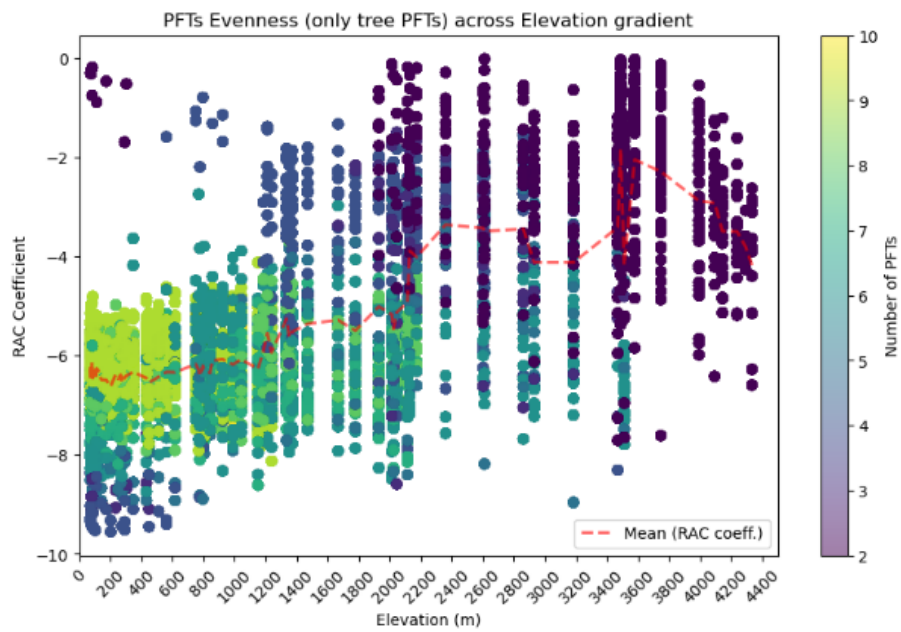
574



575

Figure 8: PFTs distribution and composition by fractional projective cover along the elevation gradient (inset figure shows the PFTs distribution ranges recorded by Maharjan et al., 2021 in the elevation gradient).

The RAC coefficient value, calculated for each simulated grid using the FPC value of trees, shows that PFT's evenness increases rapidly along the elevation gradient starting from mid-elevation (after 2000 m) (Figure 9). As ~~latitude-elevation~~ increases, PFT abundance decreases ~~significantly~~, from a maximum of 10 PFTs in lower elevations to just ~~two~~2 in higher elevations (Figure 9). In the high PFT abundance area, the competition was higher, ~~reflected-characterized~~ by the competitive dominance of a few PFTs (Supplementary figure ~~65~~5). For example, tropical broadleaved raingreen, tropical broadleaved evergreen in lower elevation, sub-tropical conifers, and subtropical intermediate shade tolerant broadleaves in the mid-elevation range (Supplementary figure 5). These patterns suggest a symmetrical competition for light and nutrients at lower elevation despite richness in species composition. In contrast, regions with low PFT abundance exhibited higher evenness (coefficient value close to 0), indicating reduced competition and a stronger role of environmental filtering. Here, plant adaptation to temperature and soil nutrient limitation becomes the primary driver of vegetation structure and composition. A more detailed breakdown of PFT-specific FPC distributions and dominance patterns along the elevation gradient is provided in Supplementary Figure ~~65~~5.



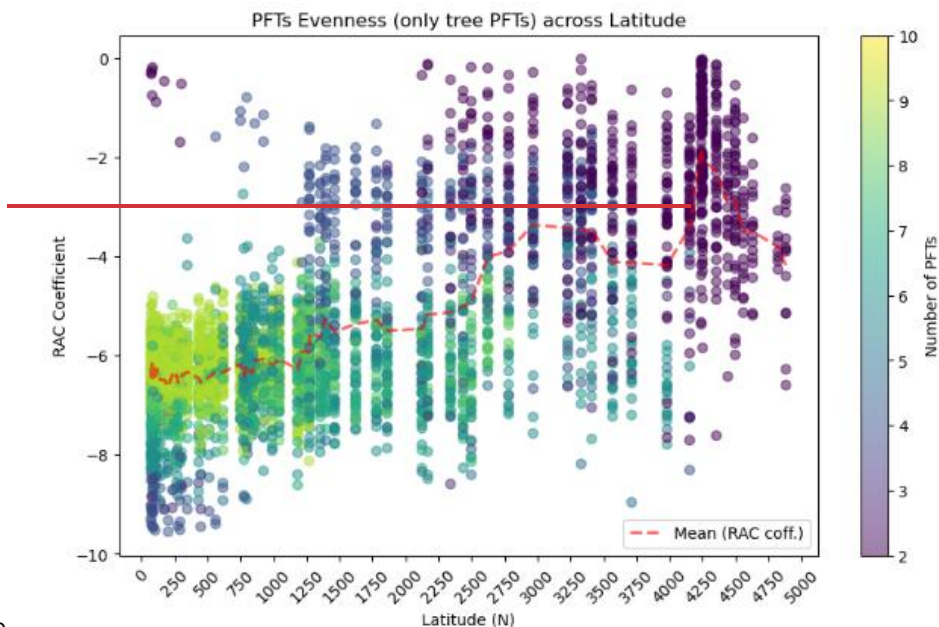


Figure 9: PFTs evenness (RAC coefficient) across latitude with the number of PFTs (present) in each simulated grid.

Deciduousness and shade tolerance emerges as the two most dominant plant adaptation mechanisms of vegetation which vary systematically along the elevation gradient. The proportion of shade-tolerant species distribution decreases with elevation. Intermediate shade tolerance (*Schima wallichii*) has maximum FPC in the mid-elevation ranges (1200 - 2500 m), suggesting that these elevation ranges offer a balance in light and temperature conditions, favouring flexible growth strategies. The deciduous broadleaved species form dominant crown coverage in lower elevations, but their contribution to FPC decreases with increased elevation, where cold temperature limit their performance. In contrast, Evergreen conifer's contribution to FPC increases with an increase in elevation up to 4500 m, corresponding to zones dominated by i.e., up to where Pinus coniferous species, mainly *Abies* and *Juniperus* species, mostly grow (Figure 10). At the cold end of the gradient, the alpine broadleaved evergreen, especially *Rhododendron spp.*, contributes the maximum in FPC. The dominance of evergreen trees at higher elevations reflects an adaptation strategy to cold climates, where year round leaf retention allows for rapid photosynthetic response to short favourable growing

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font: Not Italic, Font color: Red

Formatted: Font: Not Italic, Font color: Red

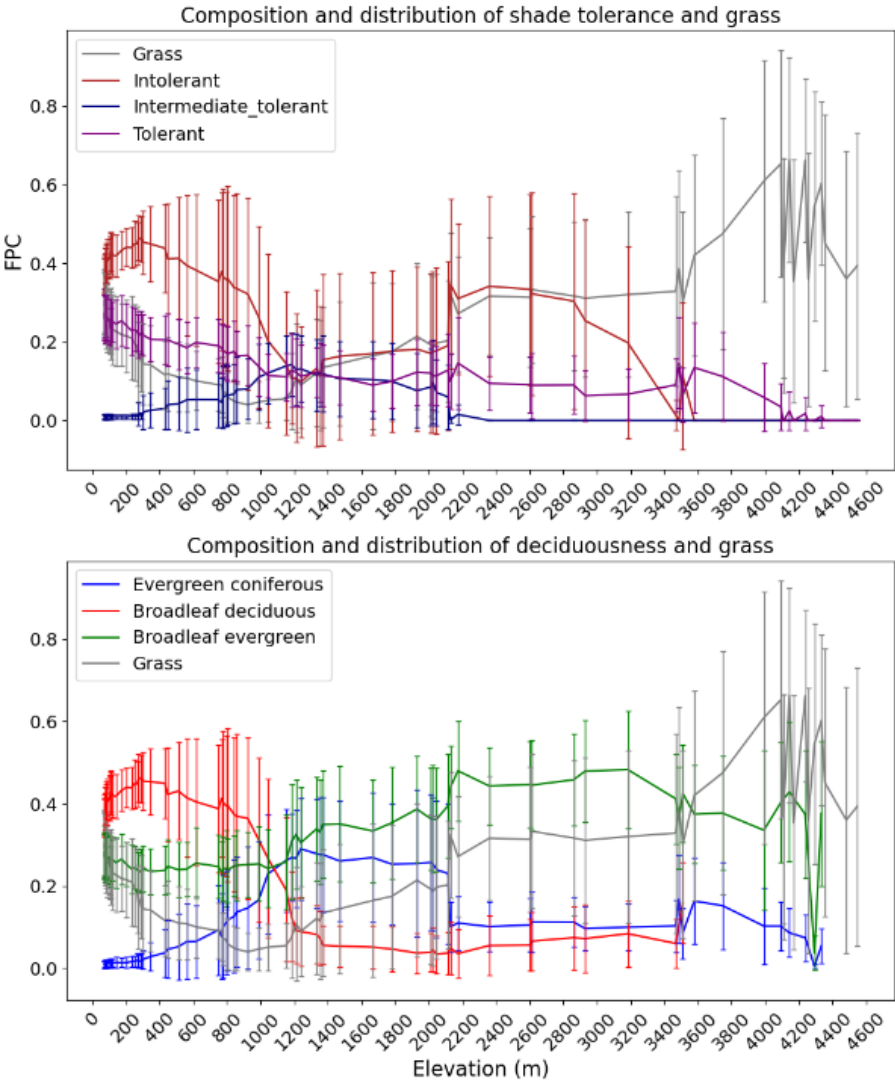
Formatted: Font color: Red

Formatted: Font: Not Italic, Font color: Red

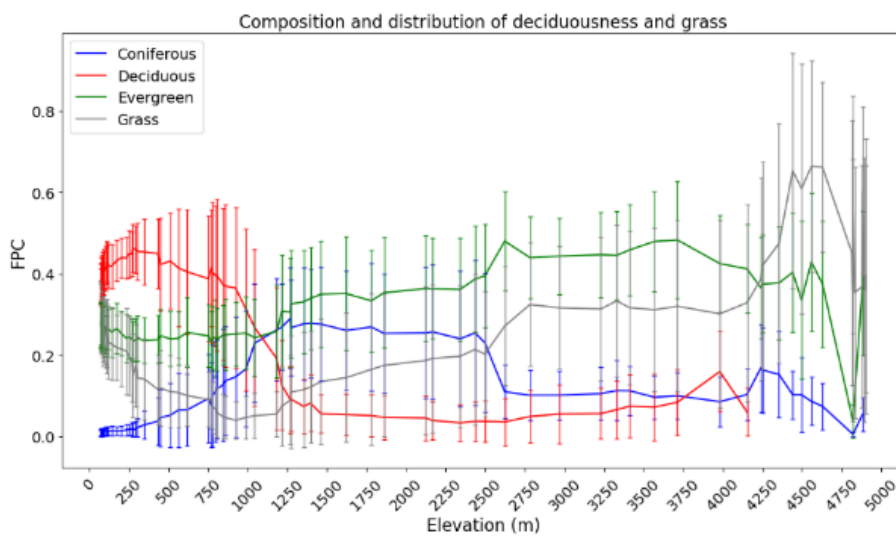
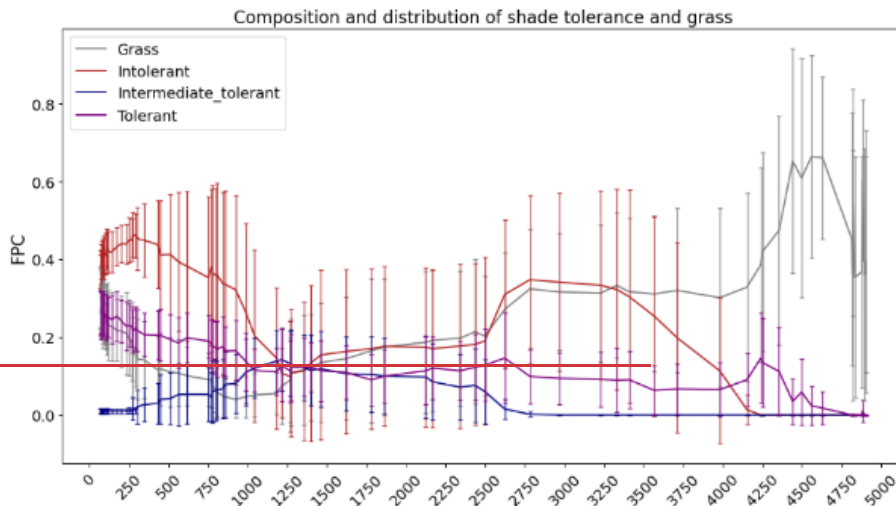
Formatted: Font color: Red

Formatted: Font color: Red

618 periods. Notably, around 1000 m elevation (Figure 10), both shade tolerance and
619 deciduousness exhibit high evenness in PFT distribution, indicating a transitional zone
620 where multiple plant adaptation strategies coexist due to overlapping ecological niches
621 shaped by complex interaction between plant traits and environmental conditions.



622



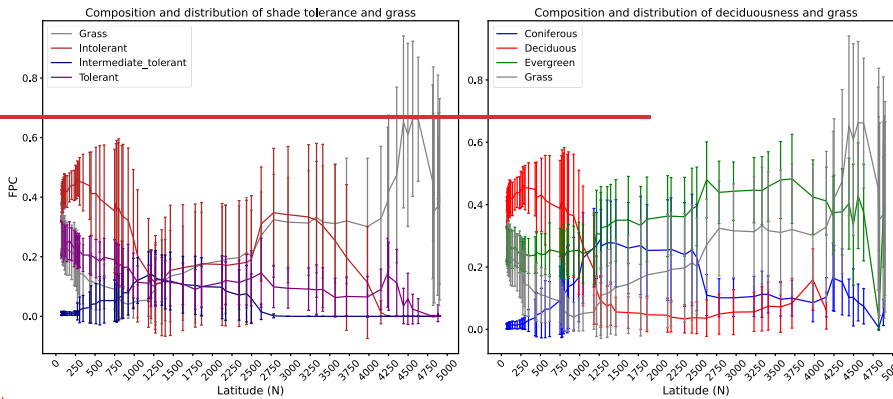


Figure 10: Distribution and composition of PFTs adaptation strategies (shade tolerance and deciduousness) and grasses (line represents mean and bar represents standard deviation) along the elevation gradient.

7.5. Discussion

We evaluated the complex interaction between growth conditions, PFT abundance, productivity, and competitive interactions by simulating these ~~assumed~~ factors and their interdependent dynamics along the Himalayan elevation gradient. Our results reveal that between both abiotic and biotic filters competition generate distinct ecosystem states through elevation-dependent niche differences. As expected, our model predicted pronounced shifts in ecosystem structure, composition, and productivity along the ~~Himalayan elevational gradient~~, reflecting a transition from competition-regulated community assembly at lower elevations to stress-filtered assembly at higher elevations. driven by shifting interactions between climatic conditions, especially temperature and nutrient availability, and plant adaptation strategies. At lower elevations, warmer temperatures and relatively higher nutrient availability allowed many PFTs to coexist within their physiological tolerance limits, resulting in intense asymmetric competition for light. Under these conditions, realised niches dominate community structure, with a few competitively superior PFTs suppressing others despite functional richness. Competition between co-occurring PFTs in crowded stands influenced the emergent composition and productivity, particularly at lower elevations, where warmer temperature and higher resource availability allowed species to occupy their realised

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

648 ~~niches.~~ In contrast, colder and nutrient-limited conditions at higher elevations
649 constrained community composition and structure to a few cold-tolerant species with
650 short bole heights, operating within their physiological niches, resulting in more even but
651 less diverse stands. These patterns support our hypothesis that vegetation structures,
652 composition, and productivity along the elevation gradient are structured by a shift from
653 realised niches, defined by competitive interactions at lower elevations, to physiological
654 niches, shaped by stress (freezing temperatures) at higher elevations and align with the
655 predictions of the stress gradient hypothesis.

Formatted: Font color: Red

659 The model reliably reproduced key ecological patterns along the elevation gradient, with
660 strong agreement between simulated outcomes and observations or independent
661 reconstructed variables, supporting the plausibility of the results and the underlying
662 mechanisms. Simulated GPP decreased with elevation, in agreement with the estimates
663 from the leaf light use efficiency model by Bi ~~and~~ Zhou (2022), reflecting the strong
664 control of temperature and growing-season length on carbon uptake in montane
665 ecosystems reflecting the dependency of GPP on temperature and growing-season length.
666 This decline aligned with the distribution of cold-tolerant PFTs such as *Rhododendron*
667 (ABE), Alpine and cConiferous species (ANE), and C₃ grasses found ~~at~~ higher elevations,
668 consistent with the National Forest Inventory, which identifies that *Rhododendron* and
669 *Abies* are the two most dominant species in higher elevations of the Himalayas (DFRS,
670 2015).

Formatted: Not Highlight

Formatted: Not Highlight

Formatted: Subscript, Not Highlight

Formatted: Not Highlight

672 ~~Although the model does not represent detailed soil chemistry, the presence of these PFTs~~
673 ~~in the simulation aligns with their known adaptation mechanisms to high pH and low~~
674 ~~nutrients, characterized by a wide distribution range. Conifers and *Rhododendron* species~~
675 ~~are native to the Himalayan range and exhibit a wide physiological niche space that allows~~
676 ~~adaptation in high pH and low nutrient soil from temperate to alpine (sub-alpine) habitats~~
677 ~~(Thakur et al., 2024).~~ Simulated above-ground biomass and bole height patterns closely
678 matched plot-based forest inventory measurements from Khanal and Boer (2024) and
679 bole height from Maharjan et al. (2021), indicating that the model captures key structural
680 and functional forest responses to elevation. The simulated results showed that the

Formatted: Font color: Red

Formatted: Font color: Red, Not Highlight

Formatted: Not Highlight

Above-ground carbon biomass gradually increased, before a significant decline with elevation (temperature) due to climatic stress, which is consistent with the patterns reported by Thakur et al. (2024). Observed biomass at forest area at higher elevations exhibits marked spatial variability. The higher estimation of observed values at the higher elevation may could arise from local topographic and microclimatic heterogeneity that favours the growth of fir (*Abies* spp.), brown oak (*Quercus* spp.), and *Rhododendron* spp. variability in topographic factors that create favourable microclimatic conditions for the growth of *Rhododendron* and *Abies* forests, particularly in deep gorges and on southern aspects. The Himalayas region is characterized by a unique environmental condition where temperature and soil water availability vary on a short spatial scale (Khanal et al., 2025; Thakur and Chawla, 2019; Tito et al., 2020), enabling certain PFTs to extend their realised niche beyond broader elevational trends. These species attain high biomass through contrasting strategies: brown oak and *Rhododendron* attain high biomass primarily through high wood density and structural investment, while fir attains high biomass through large stem volume and tall stature despite relatively lower wood density (Kumar et al., 2024), and these species have physiological tolerance to cold and nutrient limitation combined with persistent canopy cover. Although the model does not represent detailed soil chemistry, the presence of these *Rhododendron* and *Abies* PFTs in the simulation aligns with their known adaptation mechanisms to high pH and low nutrients, characterized by a wide distribution range (Thakur et al., 2024), which are implicitly represented by trait parameters within the PFT framework. Conifers and *Rhododendron* species are native to the Himalayan range and exhibit a wide physiological niche space that allows adaptation in high pH and low nutrient soil from temperate to alpine (sub-alpine) habitats (Thakur et al., 2024), resulting in structural diversity and competitive dynamics, allowing certain PFTs to extend their realised niche beyond broader patterns along the gradient.

Vegetation structure along the elevation gradient was significantly impacted by the environmental conditions. In lower elevations, where favourable climatic growth promotes conditions promote crowded stand conditions, taller bole heights, facilitating competitive advantages through increased light interception and asymmetric competition are more favourable and soil nutrients are abundant, PFTs tend to exhibit

Formatted: Font color: Red, Not Highlight

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font: Italic, Font color: Red

Formatted: Font color: Red

Formatted: Font: Italic, Font color: Red

Formatted: Font color: Red

Formatted: Font: Italic, Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red, Not Highlight

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red, Not Highlight

Formatted: Font color: Red

Formatted: Font: Italic, Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Not Highlight

Formatted: Not Highlight

Formatted: Font: Italic, Not Highlight

Formatted: Not Highlight

Formatted: Not Highlight

Formatted: Highlight

~~higher bole heights as a strategy to maximize light interception, avoid asymmetric competition for light, and reduce shading for competitors.~~ Additionally, taller bole height of trees in these conditions may result from a trade-off, where some resources are allocated away from other parts, such as stem and branch development (Tsunoda et al., 2025). Pokhrel and Sherpa (2020) also found ~~that significant associations between~~ tree height, DBH, and above-ground biomass ~~are significantly associated with~~ and elevation in ~~along the elevation gradient of the central Himalayas.~~ With the increase in altitude, tree growth declines, and light competition ~~also declines~~ weakens, resulting in forests with ~~where trees tend to have similar~~ similar basal area and total tree height (Coomes and Allen, 2007). In contrast, alpine environments favour structurally compact morphologies with reduced height growth but sustained leaf area through evergreen strategies. The relatively high LAI at higher elevations despite declining productivity ~~LAI increased with an increase in elevation.~~ In higher elevations, an increase in LAI was associated with the an increase in abundance of evergreen vegetation (Figure 5 & 10). ~~This pattern, especially~~ *Rhododendron* and *Abies* species. *Rhododendron* presence in higher elevations is linked with reflects survival strategies, including heat dissipation to avoid damage by excessive radiation in warm seasons ~~periods, as well as~~ and physiological mechanisms adjustments such as increasing intercellular fluid concentration and using reactive oxygen to withstand chilling temperatures to tolerate low-temperature stress (Li et al., 2022). ~~However~~ Although, these physiological mechanisms / strategies are ~~explicitly~~ not ~~explicitly represented~~ parameterized in our the model, they emerge implicitly through trait-based PFTs representation. Additionally, traits such as reduced height, smaller individual leaf area, lower SLA, and trade-offs in vessel diameter and density could also ~~facilitate the wide distribution of Rhododendron species (Pandey et al., 2021).~~

The simulated vegetation community exhibited a distinct shift in compositional patterns across the elevation gradient ~~that reflects changing competitive and abiotic conditions.~~ At lower elevations, favourable climatic conditions support high PFT richness and functional diversity; along with strong asymmetric competition, resulting in dominance by a few competitively superior PFTs operating within their realised niches. In lower elevations, ~~the coexistence of a large number of PFTs resulted in higher functional diversity, likely due to favourable growth conditions, as the presence of large PFTs within their physiological niches led to higher competitive dominance by a few PFTs.~~ Among 10 more

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

than ~~tend~~ different PFTs present, *Shorea robusta* (PFT-TrBRG) emerged as the dominant climax species ~~at the lower elevations~~, showing consistency with national forest inventory data, which shows that *Shorea robusta* and its associated species, such as *Terminalia alata*, *Mollunthous philippines*, and *Lagerstroemia parviflora*, are the most common and productive in terms of biomass in the southern ~~part of the~~ central Himalayas (DFRS, 2015). ~~The~~ its competitive dominance and higher productivity these species were associated with linked to drought and fire resistance (Gautam and Devoe, 2006). conferring competitive advantages under seasonal climatic variability. In contrast, higher elevations supported only a few PFTs (Supplementary Figure 5), with community composition shaped by abiotic stress such as low temperature and short growing seasons (Figure 7). Under these conditions, competitive exclusion weakens, and no single PFT dominates, resulting in more even but less diverse communities. Ahmad et al. (2025) also stated that functional diversity is higher in lower elevation, even though species richness and phylogenetic diversity are higher in mid-elevation across the Himalayas. These elevational shifts in dominance and diversity is pattern is broadly consistent with the stress-gradient hypothesis, which ~~posits~~ explains that abiotic stress dominates in harsher conditions, and competition is more influential in a benign environment (Bertness and Callaway, 1994). Additionally, the simulated PFTs composition is consistent with ~~m~~Mid-domain effects (Colwell and Lees, 2000; Smith and Wilson, 1996) with a large number of PFT peaks at the intermediate elevation range (1000- 2000 m), likely due to geometric constraints on range overlap (Figures 6, 8, and 10). Overall, PFTs richness and diversity decline monotonically at higher elevation The overall species composition pattern showed a monotonic decrease in species richness and diversity with increasing elevation gradient, with higher heterogeneity and the presence of unique species adapted to extreme climatic conditions (Sekar et al., 2024). At the lower end of the gradient, with higher elevation characterized by fewer and more specialized PFTs adapted to extreme climatic conditions ~~more seasonal rainfall and dry conditions, the vegetation was characterized by lower evenness and higher competitive dominance, as PFTs relied on realised niche occupation~~ (Figure 8).

The difference in PFTs' performance and dominance along the gradient ~~depended~~ emerge from an interaction between the climatic condition and ~~their~~ trait-mediated trade-offs that regulate carbon allocation, growth form, and competitive ability under contrasting

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

~~climatic constraint, allometric relations.~~ Overall, tropical broadleaved raingreen (TrBRG) was the most productive, followed by temperate evergreen (TeIBE) and alpine evergreen (ABE) across the elevation gradient. This result is consistent with the national forest inventory report, which states ~~species contribution in overall productions varies with elevation~~that *Shorea robusta* (37.83 ton ha⁻¹), *Quercus* species (46.09 ton ha⁻¹), and *Rhododendron* species (11.22 ton ha⁻¹) are the most productive species in Nepal, with significant contributions from *Pinus* species (DFRS, 2015). ~~At lower elevations, PFTs with rapid growth rate, efficient light capture, and high carbon gain dominate through asymmetric competition for light and space, which is translated into competitive superiority, reinforcing realised niche occupation and hierarchical dominance structures. However, in higher elevation balance shifts from competitive suppression to persistence under environmental stress and overall ecosystem productive decreases. These variations in productivity among PFTs therefore reflect underlying trait trade-offs, particularly involving wood density, size-density relationships, growth allocation strategies and their niche~~ (Khanal et al., (2024)). ~~also concluded that species growth and carbon mass production depends on the wood density and size density relation. *Rhododendron* (ABE) demonstrated wide niche space, and *Shorea robusta* (TrBRG), with its strong climatic niche and role climax vegetation, illustrates how PFT performance is shaped by the climatic niche at the elevations in which the species occurs. PFTs characterised by dense wood and conservative growth strategies maintain biomass through structural investment and stress tolerance, whereas PFTs achieving large stem volume sustain biomass primarily through size expansion when climatic conditions permit. As these contrasting strategies converge under harsher alpine conditions, performance differences among PFTs diminish, competitive hierarchies weaken, and no single PFT consistently dominates (Naud et al., 2019), despite broad climatic niche breadth in taxa such as *Rhododendron*. The results show that competitive dominance was not visible in higher elevations, with no individual PFTs becoming dominant. Similar to our finding, Naud et al. (2019) also highlighted that no individual species becomes dominant with increased altitude when species richness decreases, which is similar in the Himalayas.~~

Formatted: Font: Italic

Formatted: Font color: Red

A range of mechanisms, including deciduousness, shade tolerance, and drought resistance, allometric relationships, and wood density collectively determine how PFTs allocate carbon, compete for resources, and persist under varying environmental conditions. ~~contributed to the presence and abundance of PFTs under different environmental conditions along the gradient in our simulations.~~ In lower elevations, deciduousness emerges as an adaptation to escape seasonal drought was dominant, enabling drought avoidance and rapid resource use under seasonal climates. ~~whereas, In contrast, in higher elevations favour evergreen cold and shade tolerant PFTs were abundant, reflecting their functional strategy to adapted~~ to stresses such as cooler temperatures, shorter growing seasons, and harsher growing conditions. Similarly, allometric traits such as crown dimension-DBH, height-DBH, and wood density affected competitive interaction and competitive dynamics. In lower elevations, PFT coverage in the crown showed a clear hierarchy with the differences in competitive dominance in productivity and FPC. ~~Previous studies have shown that traits shape the distribution of vegetation (Maharajan et al., 2021), and competition determines productivity under favourable conditions (Sauter et al., 2021).~~ Our study further emphasized that growth conditions, coupled with biotic and abiotic interactions, and trade-offs between growth and adaptation to multiple stresses drive the overall ecosystem functioning along the elevation gradient. Under ongoing global change, relaxation of thermal constraints at higher elevations may intensify competition in currently stress-filtered communities, while increased climatic variability may narrow the zone of higher functional diversity at mid-elevations, with consequences for ecosystem resilience and carbon storage. ~~This is reflected in differences in productivity and vegetation structure along the elevation gradient, with variation in species abundance and evenness according to environmental conditions.~~

Formatted: Font color: Red

Formatted: Font color: Red

5.1 Limitations

We used high-resolution climate data (3 km) as forcing data to capture heterogeneity in climatic conditions along the elevation gradient. ~~This approach successfully captured broad elevational patterns in the dynamics and composition of the ecosystem.~~ However, the model does not fully capture the variability in microclimatic conditions that creates distinct topoclimatic conditions which favour high biomass accumulation, ~~created by~~ small-scale topography. This may partially account for the underestimation of above-

Formatted: Font color: Red

Formatted: Font color: Red

845 ~~ground carbon stock at ecosystem level in our simulations compared to forested area of~~
846 ~~particular range especially in high elevations.~~ With the elevation increase, climatic
847 heterogeneity amplifies with more diverse climatic conditions (Guan et al., 2024).
848 Integrating slope and aspect in the model could enhance its ability to characterize
849 vegetation dynamics and composition dynamics, particularly in mountain regions where
850 variation in radiation, soil moisture, and temperature across slope orientations plays a
851 crucial role. ~~In addition, trait data from Maharjan et al. (2021) Regional species trait data~~
852 ~~were used for defining functional groups representing 31 dominant tree species along the~~
853 ~~gradient. While, this approach captures the major functional strategies governing~~
854 ~~ecosystem processes, some locally species-abundant or patch-dominant species that~~
855 ~~contribute to overall ecosystem functioning and structural heterogeneity may not be~~
856 ~~explicitly represented. As a result, fine-scale variation in biomass, functioning, structure~~
857 ~~and community composition may be smoothed in the simulations. This reflects a~~
858 ~~limitation of using generalized PFTs, which may not fully capture species-specific~~
859 ~~responses and local ecological variability. Incorporating an expanded trait database and~~
860 ~~higher function resolution could further enhance the representation of local biodiversity~~
861 ~~and ecosystem structural component across complex mountain ecosystem.~~

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

Formatted: Font color: Red

862
863 Even though dynamic vegetation models like LPJ-GUESS integrate competition for space,
864 light, and soil resources among neighbouring plants, this competitive framework may not
865 fully account for how trait-based plasticity alters competitive interactions under different
866 growth conditions. This can limit the model's ability to capture the full range of ecosystem
867 dynamics. Integrating the allometric relation defined based on growth (competition)
868 conditions helped to better characterise above-ground competition for light and space,
869 whereas competition for below-ground resources, i.e. water and nitrogen in the model,
870 largely follows proportionately with plant size. In reality, differential root profiles,
871 including deep water access via tap roots characteristic of certain tree taxa, and other
872 factors such as mycorrhizal associations or release of root exudates to promote nutrient
873 mineralisation and uptake are known to predict plant success in environments
874 characterised by below-ground resource limitations (Freschet et al., 2021). Dynamic root
875 allocation based on resource availability in different layers is not currently simulated in
876 LPJ-GUESS. Integrating root trait data, especially the distribution of fine roots in different
877 soil profiles, root and mycorrhizal association (De Paula et al., 2021), and interlinking

878 them with soil depth, may better capture below-ground competition processes. The
879 current version of LPJ-GUESS incorporates a climate-driven prognostic wildfire scheme
880 (SIMFIRE-BLAZE; Rabin et al., 2017) which impacts the composition, structure, and
881 dynamics of the vegetation. However, other forms of disturbance, such as wood cutting
882 and managed fires, herbivory, insect pest impacts, and wind-throw shape vegetation
883 composition and demography, compounding with the biophysical and ecological
884 mechanisms included in the model (Brewer, 2011; Grime, 1973; Hall et al., 2012; Laurent
885 et al., 2017).

886
887 **8.6. Conclusions**

888 ~~After incorporating trait data and allometric relations reflecting of regional PFTs into~~
889 ~~LPJ-GUESS model, this study provides a mechanistic representation of vegetation~~
890 ~~dynamics from our along the Himalayan elevation gradient. The model study region, we~~
891 ~~successfully reproduced spatial and temporal patterns generally consistent~~
892 ~~with observed in vegetation composition, structure, and productivity and capturing key~~
893 ~~ecological processes across contrasting climatic conditions along the elevation gradient.~~
894 Our ~~model results~~ shows that environmental conditions, biotic and abiotic interactions,
895 allometric relations, and associated functional trade-offs jointly shape ecosystem
896 processes and drive the competition patterns and adaptation mechanisms ~~of vegetation~~
897 ~~along the elevation gradient~~. At the productive end of the gradient, competitive
898 interactions among woody PFTs in crowded stands had a strong influence on PFT
899 performance and abundance. These interactions, together with the realised niches of
900 PFTs, led to reduced evenness in PFT distribution, as certain PFTs become dominant in
901 different climatic conditions. This did not translate into higher PFT richness in the way
902 predicted by classical niche theory, suggesting favourable environmental conditions may
903 buffer competitive exclusion and promote species coexistence despite competitive
904 dominance hierarchies. In contrast, under more stressful conditions, only a few PFTs
905 survive and grow, exhibiting higher evenness in composition and shorter stature. The
906 increase in evenness with elevation reflects reduced crowding and weak asymmetric
907 competition, with the surviving PFTs adapting and persisting by occupying their
908 physiological niches in response to prevailing abiotic stress, particularly cold and freezing
909 temperatures. Our findings highlight how variations in climatic conditions, resource
910 availability, the climatic niche of PFTs, and unique adaptation mechanisms interact with

Formatted: Font color: Red

Formatted: Font color: Red

911 PFT traits and adaptation strategies to shape vegetation composition patterns and
912 productivity, thereby acting as overall controls on ecosystem function along the gradient.
913 Further study, integrating heterogeneity in topographic conditions with different
914 disturbances, along with a representation of ~~below-under~~ground competition (especially
915 root profiles and groundwater dynamics), could enhance our understanding of ecosystem
916 responses to global changes and extreme events, as well as their adaptation mechanisms
917 in the central Himalayas.

918

919

920 **7. Code and Data Availability**

921 The customized LPJ-GUESS version used in this study has been archived in the **LPJ-GUESS**
922 **Zenodo community** [<https://doi.org/10.5281/zenodo.17214801>]. The forcing data,
923 simulated output, that reproduce the analyses presented in the manuscript have been
924 deposited in **Zenodo** [<https://doi.org/10.5281/zenodo.17214851>].

925

926 **8. Author Contribution**

927 **PP:** conceptualization and design (lead); data curation (lead); simulation (lead); formal
928 analysis (lead); writing – original draft (lead); writing – review and editing (lead). **SO:**
929 Supervision (supporting); writing – review and editing (supporting). **MT:** Supervision
930 (supporting); writing – review and editing (supporting); Supervision (supporting);
931 writing – review and editing (supporting). **MP:** Supervision (supporting); writing –
932 review and editing (supporting). **DM:** Supervision (supporting); writing – review and
933 editing (supporting). **BS:** Supervision (lead); conceptualization and design (supporting);
934 writing – original draft (supporting); writing – review and editing (equal).

935

936 **9. Competing Interest**

937 The contact author has declared that none of the authors has any competing interests.

938

939 **10. Financial support**

Formatted: Font: (Default) Cambria, Bold

Formatted: List Paragraph, Numbered + Level: 1 +
Numbering Style: 1, 2, 3, ... + Start at: 4 + Alignment:
Left + Aligned at: 0.25" + Indent at: 0.5"

Formatted: Font: (Default) Cambria

Formatted: Font: (Default) Cambria, Bold

Formatted: List Paragraph, Justified, Numbered +
Level: 1 + Numbering Style: 1, 2, 3, ... + Start at: 4 +
Alignment: Left + Aligned at: 0.25" + Indent at: 0.5"

Formatted: List Paragraph, Numbered + Level: 1 +
Numbering Style: 1, 2, 3, ... + Start at: 4 + Alignment:
Left + Aligned at: 0.25" + Indent at: 0.5"

Formatted: List Paragraph, Numbered + Level: 1 +
Numbering Style: 1, 2, 3, ... + Start at: 4 + Alignment:
Left + Aligned at: 0.25" + Indent at: 0.5"

Formatted: List Paragraph, Numbered + Level: 1 +
Numbering Style: 1, 2, 3, ... + Start at: 4 + Alignment:
Left + Aligned at: 0.25" + Indent at: 0.5"

940 This research has been supported by Western Sydney University as a PhD scholarship.
941 Stefan Olin was supported by the Modelling the Regional and Global Earth System
942 (MERGE).

943

944 **9.11. Reference**

945 Acharya, B. K., Chettri, B., and Vijayan, L.: Distribution pattern of trees along an elevation
946 gradient of Eastern Himalaya, India, *Acta Oecologica*, 37, 329–336,
947 <https://doi.org/10.1016/j.actao.2011.03.005>, 2011.

948 Ahmad, M., Luo, Y.-H., Rathee, S., Spicer, R. A., Zhang, J., Wambulwa, M. C., Zhu, G.-F.,
949 Cadotte, M. W., Wu, Z.-Y., Khan, S. M., Maity, D., Li, D.-Z., and Liu, J.: Multifaceted plant
950 diversity patterns across the Himalaya: Status and outlook, *Plant Divers.*, 47, 529–543,
951 <https://doi.org/10.1016/j.pld.2025.04.003>, 2025.

952 Araghi, A. and Martinez, C. J.: Evaluation of CRU-JRA gridded meteorological dataset for
953 modeling of wheat production systems in Iran, *Int. J. Biometeorol.*, 68, 1201–1211,
954 <https://doi.org/10.1007/s00484-024-02659-9>, 2024.

955 Asner, G. P., Anderson, C. B., Martin, R. E., Knapp, D. E., Tupayachi, R., Sinca, F., and Malhi,
956 Y.: Landscape-scale changes in forest structure and functional traits along an Andes-to-
957 Amazon elevation gradient, *Biogeosciences*, 11, 843–856, [https://doi.org/10.5194/bg-](https://doi.org/10.5194/bg-11-843-2014)
958 11-843-2014, 2014.

959 Avolio, M. L., Forrestel, E. J., Chang, C. C., La Pierre, K. J., Burghardt, K. T., and Smith, M. D.:
960 Demystifying dominant species, *New Phytol.*, 223, 1106–1126,
961 <https://doi.org/10.1111/nph.15789>, 2019.

962 Ayer, S., Yadav, B. K., and Bhatta, K. P.: What drives the regeneration dynamics in Central
963 Himalayan Mountain Forests of Nepal?, *Environ. Chall.*, 21, 101359,
964 <https://doi.org/10.1016/j.envc.2025.101359>, 2025.

965 Bertness, M. D. and Callaway, R.: Positive interactions in communities, *Trends Ecol. Evol.*,
966 9, 191–193, [https://doi.org/10.1016/0169-5347\(94\)90088-4](https://doi.org/10.1016/0169-5347(94)90088-4), 1994.

967 Bi, W. and Zhou, Y.: A global 0.05° dataset for gross primary production of sunlit and
968 shaded vegetation canopies (1992–2020) (18),
969 <https://doi.org/10.5061/DRYAD.DFN2Z352K>, 2022.

970 Brewer, J. S.: Disturbance-mediated competition between perennial plants along a
971 resource supply gradient: Disturbances and competition, *J. Ecol.*, 99, 1219–1228,
972 <https://doi.org/10.1111/j.1365-2745.2011.01846.x>, 2011.

973 Brooker, R. W. and Kikvidze, Z.: Importance: an overlooked concept in plant interaction
974 research, *J. Ecol.*, 96, 703–708, <https://doi.org/10.1111/j.1365-2745.2008.01373.x>,
975 2008.

Formatted: Font: Cambria

976 Bugmann, H., Fischlin, A., and Kienast, F.: Model convergence and state variable update
977 in forest gap models, *Ecol. Model.*, 89, 197–208, [https://doi.org/10.1016/0304-](https://doi.org/10.1016/0304-3800(95)00135-2)
978 3800(95)00135-2, 1996.

979 Colwell, Robert. K. and Lees, D. C.: The mid-domain effect: geometric constraints on the
980 geography of species richness, *Trends Ecol. Evol.*, 15, 70–76,
981 [https://doi.org/10.1016/S0169-5347\(99\)01767-X](https://doi.org/10.1016/S0169-5347(99)01767-X), 2000.

982 Coomes, D. A. and Allen, R. B.: Effects of size, competition and altitude on tree growth, *J.*
983 *Ecol.*, 95, 1084–1097, <https://doi.org/10.1111/j.1365-2745.2007.01280.x>, 2007.

984 De Frenne, P., Graae, B. J., Rodríguez-Sánchez, F., Kolb, A., Chabrierie, O., Decocq, G., De
985 Kort, H., De Schrijver, A., Diekmann, M., Eriksson, O., Gruwez, R., Hermy, M., Lenoir, J.,
986 Plue, J., Coomes, D. A., and Verheyen, K.: Latitudinal gradients as natural laboratories to
987 infer species' responses to temperature, *J. Ecol.*, 101, 784–795,
988 <https://doi.org/10.1111/1365-2745.12074>, 2013.

989 De Paula, M. D., Forrest, M., Langan, L., Bendix, J., Homeier, J., Velescu, A., Wilcke, W., and
990 Hickler, T.: Nutrient cycling drives plant community trait assembly and ecosystem
991 functioning in a tropical mountain biodiversity hotspot, *New Phytol.*, 232, 551–566,
992 <https://doi.org/10.1111/nph.17600>, 2021.

993 DFRS: STATE of NEPAL'S FORESTS, Department of Forest Research and Survey,
994 Kathmandu, <https://doi.org/978-9937-8896-3-6>, 2015.

995 Dhakal, B. P.: Traits and distribution along the altitudinal gradients of the Himalayas
996 gradient of the Himalayas , Nepal, 1–75 pp., 2018.

997 Díaz, S., Kattge, J., Cornelissen, J. H. C., Wright, I. J., Lavorel, S., Dray, S., Reu, B., Kleyer, M.,
998 Wirth, C., Colin Prentice, I., Garnier, E., Bönisch, G., Westoby, M., Poorter, H., Reich, P. B.,
999 Moles, A. T., Dickie, J., Gillison, A. N., Zanne, A. E., Chave, J., Joseph Wright, S., Sheremet'ev,
1000 S. N., Jactel, H., Baraloto, C., Cerabolini, B., Pierce, S., Shipley, B., Kirkup, D., Casanoves, F.,
1001 Joswig, J. S., Günther, A., Falczuk, V., Rüger, N., Mahecha, M. D., and Gorné, L. D.: The global
1002 spectrum of plant form and function, *Nature*, 529, 167–171,
1003 <https://doi.org/10.1038/nature16489>, 2016.

1004 Drollinger, S., Müller, M., Kobl, T., Schwab, N., Böhner, J., Schickhoff, U., and Scholten, T.:
1005 Decreasing nutrient concentrations in soils and trees with increasing elevation across a
1006 treeline ecotone in Rolwaling Himal, Nepal, *J. Mt. Sci.*, 14, 843–858,
1007 <https://doi.org/10.1007/s11629-016-4228-4>, 2017.

1008 Dyola, N., Sigdel, S. R., Liang, E., Babst, F., Camarero, J. J., Aryal, S., Chettri, N., Gao, S., Lu,
1009 X., Sun, J., Wang, T., Zhang, G., Zhu, H., Piao, S., and Peñuelas, J.: Species richness is a
1010 strong driver of forest biomass along broad bioclimatic gradients in the Himalayas,
1011 *Ecosphere*, 13, <https://doi.org/10.1002/ecs2.4107>, 2022.

1012 Earth Resources Observation and Science (EROS) Center: Shuttle Radar Topography
1013 Mission (SRTM) 1 Arc-Second Global, <https://doi.org/10.5066/F7PR7TFT>, 2000.

1014 Earth Resources Observation and Science (EROS) Center: Global 30 Arc-Second
1015 Elevation (GTOPO30), <https://doi.org/10.5066/F7DF6PQS>, 2017.

1016 Falster, D. S., Duursma, R. A., Ishihara, M. I., Barneche, D. R., FitzJohn, R. G., Vårhammar,
 1017 A., Aiba, M., Ando, M., Anten, N., Aspinwall, M. J., Baltzer, J. L., Baraloto, C., Battaglia, M.,
 1018 Battles, J. J., Bond-Lamberty, B., Van Breugel, M., Camac, J., Claveau, Y., Coll, L., Dannoura,
 1019 M., Delagrange, S., Domec, J.-C., Fatemi, F., Feng, W., Gargaglione, V., Goto, Y., Hagihara, A.,
 1020 Hall, J. S., Hamilton, S., Harja, D., Hiura, T., Holdaway, R., Hutley, L. S., Ichie, T., Jokela, E. J.,
 1021 Kantola, A., Kelly, J. W. G., Kenzo, T., King, D., Kloeppel, B. D., Kohyama, T., Komiyama, A.,
 1022 Laclau, J.-P., Lusk, C. H., Maguire, D. A., Le Maire, G., Mäkelä, A., Markesteijn, L., Marshall,
 1023 J., McCulloh, K., Miyata, I., Mokany, K., Mori, S., Myster, R. W., Nagano, M., Naidu, S. L.,
 1024 Nouvellon, Y., O'Grady, A. P., O'Hara, K. L., Ohtsuka, T., Osada, N., Osunkoya, O. O., Peri, P.
 1025 L., Petritan, A. M., Poorter, L., Portsmouth, A., Potvin, C., Ransijn, J., Reid, D., Ribeiro, S. C.,
 1026 Roberts, S. D., Rodríguez, R., Saldaña-Acosta, A., Santa-Regina, I., Sasa, K., Selaya, N. G.,
 1027 Sillett, S. C., Sterck, F., Takagi, K., Tange, T., Tanouchi, H., Tissue, D., Umehara, T., Utsugi, H.,
 1028 Vadeboncoeur, M. A., Valladares, F., Vanninen, P., Wang, J. R., Wenk, E., Williams, R., De
 1029 Aquino Ximenes, F., Yamaba, A., Yamada, T., Yamakura, T., Yanai, R. D., and York, R. A.:
 1030 BAAD: a Biomass And Allometry Database for woody plants: Ecological Archives E096-
 1031 128, Ecology, 96, 1445–1445, <https://doi.org/10.1890/14-1889.1>, 2015.

1032 Freschet, G. T., Roumet, C., Comas, L. H., Weemstra, M., Bengough, A. G., Rewald, B.,
 1033 Bardgett, R. D., De Deyn, G. B., Johnson, D., Klimešová, J., Lukac, M., McCormack, M. L.,
 1034 Meier, I. C., Pagès, L., Poorter, H., Prieto, I., Wurzbarger, N., Zadworny, M., Bagniewska-
 1035 Zadworna, A., Blancaflor, E. B., Brunner, I., Gessler, A., Hobbie, S. E., Iversen, C. M.,
 1036 Mommer, L., Picon-Cochard, C., Postma, J. A., Rose, L., Ryser, P., Scherer-Lorenzen, M.,
 1037 Soudzilovskaia, N. A., Sun, T., Valverde-Barrantes, O. J., Weigelt, A., York, L. M., and Stokes,
 1038 A.: Root traits as drivers of plant and ecosystem functioning: current understanding,
 1039 pitfalls and future research needs, New Phytol., 232, 1123–1158,
 1040 <https://doi.org/10.1111/nph.17072>, 2021.

1041 Friedlingstein, P., O'Sullivan, M., Jones, M. W., Andrew, R. M., Bakker, D. C. E., Hauck, J.,
 1042 Landschützer, P., Le Quéré, C., Luijkx, I. T., Peters, G. P., Peters, W., Pongratz, J.,
 1043 Schwingshackl, C., Sitch, S., Canadell, J. G., Ciais, P., Jackson, R. B., Alin, S. R., Anthoni, P.,
 1044 Barbero, L., Bates, N. R., Becker, M., Bellouin, N., Decharme, B., Bopp, L., Brasika, I. B. M.,
 1045 Cadule, P., Chamberlain, M. A., Chandra, N., Chau, T.-T., Chevallier, F., Chini, L. P., Cronin,
 1046 M., Dou, X., Enyo, K., Evans, W., Falk, S., Feely, R. A., Feng, L., Ford, D. J., Gasser, T., Ghattas,
 1047 J., Gkritzalis, T., Grassi, G., Gregor, L., Gruber, N., Gürses, Ö., Harris, I., Hefner, M., Heinke, J.,
 1048 Houghton, R. A., Hurtt, G. C., Iida, Y., Ilyina, T., Jacobson, A. R., Jain, A., Jarníková, T., Jersild,
 1049 A., Jiang, F., Jin, Z., Joos, F., Kato, E., Keeling, R. F., Kennedy, D., Klein Goldewijk, K., Knauer,
 1050 J., Korsbakken, J. I., Körtzinger, A., Lan, X., Lefèvre, N., Li, H., Liu, J., Liu, Z., Ma, L., Marland,
 1051 G., Mayot, N., McGuire, P. C., McKinley, G. A., Meyer, G., Morgan, E. J., Munro, D. R.,
 1052 Nakaoka, S.-I., Niwa, Y., O'Brien, K. M., Olsen, A., Omar, A. M., Ono, T., Paulsen, M., Pierrot,
 1053 D., Pocock, K., Poulter, B., Powis, C. M., Rehder, G., Resplandy, L., Robertson, E.,
 1054 Rödenbeck, C., Rosan, T. M., Schwinger, J., Séférian, R., et al.: Global Carbon Budget 2023,
 1055 Earth Syst. Sci. Data, 15, 5301–5369, <https://doi.org/10.5194/essd-15-5301-2023>,
 1056 2023.

1057 Gautam, K. H. and Devoe, N. N.: Ecological and anthropogenic niches of sal (*Shorea*
 1058 *robusta* Gaertn. f.) forest and prospects for multiple-product forest management – a
 1059 review, For. Int. J. For. Res., 79, 81–101, <https://doi.org/10.1093/forestry/cpi063>, 2006.

1060 Grime, J. P.: Competitive Exclusion in Herbaceous Vegetation, Nature, 242, 344–347,
 1061 <https://doi.org/10.1038/242344a0>, 1973.

1062 Guan, Y., Liu, J., Cui, W., Chen, D., Zhang, J., Lu, H., Maeda, E. E., Zeng, Z., and Beck, H. E.:
1063 Elevation Regulates the Response of Climate Heterogeneity to Climate Change, *Geophys.*
1064 *Res. Lett.*, 51, e2024GL109483, <https://doi.org/10.1029/2024GL109483>, 2024.

1065 Hall, A. R., Miller, A. D., Leggett, H. C., Roxburgh, S. H., Buckling, A., and Shea, K.:
1066 Diversity–disturbance relationships: frequency and intensity interact, *Biol. Lett.*, 8, 768–
1067 771, <https://doi.org/10.1098/rsbl.2012.0282>, 2012.

1068 Hickler, T., Smith, B., Sykes, M. T., Davis, M. B., Sugita, S., and Walker, K.: USING A
1069 GENERALIZED VEGETATION MODEL TO SIMULATE VEGETATION DYNAMICS IN
1070 NORTHEASTERN USA, *Ecology*, 85, 519–530, <https://doi.org/10.1890/02-0344>, 2004.

1071 Jackson, J. K.: Manual of Afforestation in Nepal, Second., Forest research and Survey
1072 Centre, Kathmandu, Nepal, 1994.

1073 Jucker, T., Fischer, F. J., Chave, J., Coomes, D. A., Caspersen, J., Ali, A., Loubota Panzou, G. J.,
1074 Feldpausch, T. R., Falster, D., Usoltsev, V. A., Adu-Bredu, S., Alves, L. F., Aminpour, M.,
1075 Angoboy, I. B., Anten, N. P. R., Antin, C., Askari, Y., Muñoz, R., Ayyappan, N., Balvanera, P.,
1076 Banin, L., Barbier, N., Battles, J. J., Beeckman, H., Bocko, Y. E., Bond-Lamberty, B., Bongers,
1077 F., Bowers, S., Brade, T., van Breugel, M., Chantrain, A., Chaudhary, R., Dai, J., Dalponte, M.,
1078 Dimobe, K., Domec, J. C., Doucet, J. L., Duursma, R. A., Enríquez, M., van Ewijk, K. Y.,
1079 Farfán-Rios, W., Fayolle, A., Forni, E., Forrester, D. I., Gilani, H., Godlee, J. L., Gourlet-
1080 Fleury, S., Haeni, M., Hall, J. S., He, J. K., Hemp, A., Hernández-Stefanoni, J. L., Higgins, S. I.,
1081 Holdaway, R. J., Hussain, K., Hutley, L. B., Ichie, T., Iida, Y., Jiang, H. sheng, Joshi, P. R.,
1082 Kaboli, H., Larsary, M. K., Kenzo, T., Kloeppel, B. D., Kohyama, T., Kunwar, S., Kuyah, S.,
1083 Kvasnica, J., Lin, S., Lines, E. R., Liu, H., Lorimer, C., Loumeto, J. J., Malhi, Y., Marshall, P. L.,
1084 Mattsson, E., Matula, R., Meave, J. A., Mensah, S., Mi, X., Momo, S., Moncrieff, G. R., Mora, F.,
1085 Nissanka, S. P., O'Hara, K. L., Pearce, S., Pelissier, R., Peri, P. L., Ploton, P., Poorter, L., Pour,
1086 M. J., Pourbabaie, H., Dupuy-Rada, J. M., Ribeiro, S. C., Ryan, C., Sanaei, A., Sanger, J.,
1087 Schlund, M., Sellan, G., et al.: Tallo: A global tree allometry and crown architecture
1088 database, *Glob. Change Biol.*, 28, 5254–5268, <https://doi.org/10.1111/gcb.16302>, 2022.

1089 Khanal, S. and Boer, M. M.: Plot-level estimates of aboveground biomass and soil organic
1090 carbon stocks from Nepal's forest inventory, *Sci. Data*, 10, 406,
1091 <https://doi.org/10.1038/s41597-023-02314-9>, 2023.

1092 Khanal, S., Nolan, R. H., Medlyn, B. E., and Boer, M. M.: Disentangling contributions of
1093 allometry, species composition and structure to high aboveground biomass density of
1094 high-elevation forests, *For. Ecol. Manag.*, 554, 121679,
1095 <https://doi.org/10.1016/j.foreco.2023.121679>, 2024.

1096 Khanal, S., Nolan, R. H., Medlyn, B. E., and Boer, M. M.: Topoclimatic factors create
1097 favourable conditions for carbon-dense forests in the Central Himalayas, *Sci. Rep.*, 15,
1098 35134, <https://doi.org/10.1038/s41598-025-19127-y>, 2025.

1099 Kumar, P., Kumar, A., Patil, M., Hussain, S., and Singh, A. N.: Factors influencing tree
1100 biomass and carbon stock in the Western Himalayas, India, *Front. For. Glob. Change*, 6,
1101 1328694, <https://doi.org/10.3389/ffgc.2023.1328694>, 2024.

- 1102 Lamarque, J.-F., Shindell, D. T., Josse, B., Young, P. J., Cionni, I., Eyring, V., Bergmann, D.,
1103 Cameron-Smith, P., Collins, W. J., Doherty, R., Dalsoren, S., Faluvegi, G., Folberth, G., Ghan,
1104 S. J., Horowitz, L. W., Lee, Y. H., MacKenzie, I. A., Nagashima, T., Naik, V., Plummer, D.,
1105 Righi, M., Rumbold, S. T., Schulz, M., Skeie, R. B., Stevenson, D. S., Strode, S., Sudo, K.,
1106 Szopa, S., Voulgarakis, A., and Zeng, G.: The Atmospheric Chemistry and Climate Model
1107 Intercomparison Project (ACCMIP): overview and description of models, simulations
1108 and climate diagnostics, *Geosci. Model Dev.*, 6, 179–206, [https://doi.org/10.5194/gmd-](https://doi.org/10.5194/gmd-6-179-2013)
1109 6-179-2013, 2013.
- 1110 Latombe, G., Burke, A., Vrac, M., Levavasseur, G., Dumas, C., Kageyama, M., and Ramstein,
1111 G.: Comparison of spatial downscaling methods of general circulation model results to
1112 study climate variability during the Last Glacial Maximum, *Geosci. Model Dev.*, 11, 2563–
1113 2579, [https://doi.org/10.5194/gmd-](https://doi.org/10.5194/gmd-11-2563-2018)
1114 11-2563-2018, 2018.
- 1115 Laurent, L., Mårell, A., Korboulewsky, N., Saïd, S., and Balandier, P.: How does disturbance
1116 affect the intensity and importance of plant competition along resource gradients?, *For.*
1117 *Ecol. Manag.*, 391, 239–245, <https://doi.org/10.1016/j.foreco.2017.02.003>, 2017.
- 1118 Li, H., Guo, Q., Yang, L., Quan, H., and Wang, S.: Seasonal Eco-Physiology Characteristics of
1119 Four Evergreen Rhododendron Species to the Subalpine Habitats, *Forests*, 13, 653,
<https://doi.org/10.3390/f13050653>, 2022.
- 1120 Liang, J., Ding, Z., Lie, G., Zhou, Z., Singh, P. B., Zhang, Z., and Hu, H.: Species richness
1121 patterns of vascular plants and their drivers along an elevational gradient in the central
1122 Himalayas, *Glob. Ecol. Conserv.*, 24, e01279,
1123 <https://doi.org/10.1016/j.gecco.2020.e01279>, 2020.
- 1124 Luitel, D. R., Jha, P. K., Siwakoti, M., Shrestha, M. L., and Munniappan, R.: Climatic Trends
1125 in Di fferent Bioclimatic Zones in the, *Climate*, 8, 1–18, 2020.
- 1126 Maharjan, S. K., Sterck, F. J., Dhakal, B. P., Makri, M., and Poorter, L.: Functional traits
1127 shape tree species distribution in the Himalayas, *J. Ecol.*, 109, 3818–3834,
1128 <https://doi.org/10.1111/1365-2745.13759>, 2021.
- 1129 Maletha, A., Maikhuri, R. K., Bargali, S. S., Sharma, A., Negi, V. S., and Rawat, L. S.:
1130 Vegetation dynamics and soil nutrient availability in a temperate forest along altitudinal
1131 gradient of Nanda Devi Biosphere Reserve, Western Himalaya, India, *PLoS ONE*, 17, 1–
1132 22, <https://doi.org/10.1371/journal.pone.0275051>, 2022.
- 1133 Måren, I. E., Karki, S., Prajapati, C., Yadav, R. K., and Shrestha, B. B.: Facing north or south:
1134 Does slope aspect impact forest stand characteristics and soil properties in a semiarid
1135 trans-Himalayan valley?, *J. Arid Environ.*, 121, 112–123,
1136 <https://doi.org/10.1016/j.jaridenv.2015.06.004>, 2015.
- 1137 MoFSC: Conservation Landscapes of Nepal, Ministry of Forests and Soil Conservation,
1138 Kathmandu, Nepal, 2016.
- 1139 Monteux, S., Blume-Werry, G., Gavazov, K., Kirchhoff, L., Krab, E. J., Lett, S., Pedersen, E. P.,
1140 and Väisänen, M.: Controlling biases in targeted plant removal experiments, *New Phytol.*,
1141 242, 1835–1845, <https://doi.org/10.1111/nph.19386>, 2024.

1142 Muñoz Mazón, M., Klanderud, K., Finegan, B., Veintimilla, D., Bermeo, D., Murrieta, E.,
1143 Delgado, D., and Sheil, D.: How forest structure varies with elevation in old growth and
1144 secondary forest in Costa Rica, *For. Ecol. Manag.*, 469, 118191,
1145 <https://doi.org/10.1016/j.foreco.2020.118191>, 2020.

1146 Murtagh, F. and Legendre, P.: Ward's Hierarchical Agglomerative Clustering Method:
1147 Which Algorithms Implement Ward's Criterion?, *J. Classif.*, 31, 274–295,
1148 <https://doi.org/10.1007/s00357-014-9161-z>, 2014.

1149 Myneni, R., Knyazikhin, Y., and Park, T.: MYD15A2H MODIS/Aqua Leaf Area Index/FPAR
1150 8-Day L4 Global 500m SIN Grid V006,
1151 <https://doi.org/10.5067/MODIS/MYD15A2H.006>, 2015.

1152 Naud, L., Måsviken, J., Freire, S., Angerbjörn, A., Dalén, L., and Dalerum, F.: Altitude effects
1153 on spatial components of vascular plant diversity in a subarctic mountain tundra, *Ecol.*
1154 *Evol.*, 9, 4783–4795, <https://doi.org/10.1002/ece3.5081>, 2019.

1155 Olson, D. M., Dinerstein, E., Wikramanayake, E. D., Burgess, N. D., Powell, G. V. N.,
1156 Underwood, E. C., D'amico, J. A., Itoua, I., Strand, H. E., Morrison, J. C., Loucks, C. J., Allnutt,
1157 T. F., Ricketts, T. H., Kura, Y., Lamoreux, J. F., Wettengel, W. W., Hedao, P., and Kassem, K. R.:
1158 Terrestrial Ecoregions of the World: A New Map of Life on Earth, *BioScience*, 51, 933,
1159 [https://doi.org/10.1641/0006-3568\(2001\)051\[0933:TEOTWA\]2.0.CO;2](https://doi.org/10.1641/0006-3568(2001)051[0933:TEOTWA]2.0.CO;2), 2001.

1160 Paudel, P., Olin, S., Tjoelker, M., Pontarp, M., Metcalfe, D., and Smith, B.: Impacts of Tree
1161 Allometry on Structure, Composition, Functioning and Competitive Interactions in
1162 Savanna Ecosystems on the Northern Australian Tropical Transect, *J. Biogeogr.*, 53,
1163 e70131, <https://doi.org/10.1111/jbi.70131>, 2026.

1164 Peng, S., Terrer, C., Smith, B., Ciais, P., Han, Q., Nan, J., Fisher, J. B., Chen, L., Deng, L., and
1165 Yu, K.: Carbon restoration potential on global land under water resource constraints,
1166 *Nat. Water*, 2, 1071–1081, <https://doi.org/10.1038/s44221-024-00323-5>, 2024.

1167 Pierce, S., Brusa, G., Vagge, I., and Cerabolini, B. E. L.: Allocating CSR plant functional
1168 types: the use of leaf economics and size traits to classify woody and herbaceous
1169 vascular plants, *Funct. Ecol.*, 27, 1002–1010, [https://doi.org/10.1111/1365-](https://doi.org/10.1111/1365-2435.12095)
1170 2435.12095, 2013.

1171 Pokhrel, S. and Sherpa, C.: Analyzing the relationship, distribution of tree species
1172 diversity, and above-ground biomass on the Chitwan-Annapurna landscape in Nepal, *Int.*
1173 *J. For. Res.*, 2020, <https://doi.org/10.1155/2020/2789753>, 2020.

1174 Poudel, A. S., Shrestha, B. B., Joshi, M. D., Muniappan, R., Adiga, A., Venkatramanan, S., and
1175 Jha, P. K.: Predicting the Current and Future Distribution of the Invasive Weed *Ageratina*
1176 *adenophora* in the Chitwan-Annapurna Landscape, Nepal, *Mt. Res. Dev.*, 40, R61–R71,
1177 <https://doi.org/10.1659/MRD-JOURNAL-D-19-00069.1>, 2020.

1178 Pugh, T. A. M., Arneth, A., Kautz, M., Poulter, B., and Smith, B.: Important role of forest
1179 disturbances in the global biomass turnover and carbon sinks, *Nat. Geosci.*, 12, 730–735,
1180 <https://doi.org/10.1038/s41561-019-0427-2>, 2019.

1181 Rabin, S. S., Melton, J. R., Lasslop, G., Bachelet, D., Forrest, M., Hantson, S., Kaplan, J. O., Li,
1182 F., Mangeon, S., Ward, D. S., Yue, C., Arora, V. K., Hickler, T., Kloster, S., Knorr, W., Nieradzik,
1183 L., Spessa, A., Folberth, G. A., Sheehan, T., Voulgarakis, A., Kelley, D. I., Colin Prentice, I.,
1184 Sitch, S., Harrison, S., and Arneth, A.: The Fire Modeling Intercomparison Project
1185 (FireMIP), phase 1: Experimental and analytical protocols with detailed model
1186 descriptions, *Geosci. Model Dev.*, 10, 1175–1197, [https://doi.org/10.5194/gmd-10-](https://doi.org/10.5194/gmd-10-1175-2017)
1187 1175-2017, 2017.

1188 Satyakam, Zinta, G., Singh, R. K., and Kumar, R.: Cold adaptation strategies in plants—An
1189 emerging role of epigenetics and antifreeze proteins to engineer cold resilient plants,
1190 *Front. Genet.*, 13, 909007, <https://doi.org/10.3389/fgene.2022.909007>, 2022.

1191 Scherstjanoi, M., Kaplan, J. O., and Lischke, H.: Application of a computationally efficient
1192 method to approximate gap model results with a probabilistic approach, *Geosci. Model*
1193 *Dev.*, 7, 1543–1571, <https://doi.org/10.5194/gmd-7-1543-2014>, 2014.

1194 Shah, S., Shrestha, K. K., and Scheidegger, C.: Variation in Plant Functional Traits along
1195 Altitudinal Gradient and Land Use Types in Sagarmatha National Park and Buffer Zone,
1196 Nepal, *Am. J. Plant Sci.*, 10, 595–614, <https://doi.org/10.4236/ajps.2019.104043>, 2019.

1197 Shrestha, K. B., Hofgaard, A., and Vandvik, V.: Recent treeline dynamics are similar
1198 between dry and mesic areas of Nepal, central Himalaya, *J. Plant Ecol.*, 8, 347–358,
1199 <https://doi.org/10.1093/jpe/rtu035>, 2015.

1200 Sigdel, S. R., Liang, E., Wang, Y., Dawadi, B., and Camarero, J. J.: Tree-to-tree interactions
1201 slow down Himalayan treeline shifts as inferred from tree spatial patterns, *J. Biogeogr.*,
1202 47, 1816–1826, <https://doi.org/10.1111/jbi.13840>, 2020.

1203 Sigdel, S. R., Liang, E., Rokaya, M. B., Rai, S., Dyola, N., Sun, J., Zhang, L., Zhu, H., Chettri, N.,
1204 Chaudhary, R. P., Camarero, J. J., and Peñuelas, J.: Functional traits of a plant species
1205 fingerprint ecosystem productivity along broad elevational gradients in the Himalayas,
1206 *Funct. Ecol.*, 383–394, <https://doi.org/10.1111/1365-2435.14226>, 2022.

1207 Sitch, S., Smith, B., Prentice, I. C., Arneth, A., Bondeau, A., Cramer, W., Kaplan, J. O., Levis,
1208 S., Lucht, W., Sykes, M. T., Thonicke, K., and Venevsky, S.: Evaluation of ecosystem
1209 dynamics, plant geography and terrestrial carbon cycling in the LPJ dynamic global
1210 vegetation model, *Glob. Change Biol.*, 9, 161–185, [https://doi.org/10.1046/j.1365-](https://doi.org/10.1046/j.1365-2486.2003.00569.x)
1211 2486.2003.00569.x, 2003.

1212 Smith, B. and Wilson, J. B.: A Consumer's Guide to Evenness Indices, *Oikos*, 76, 70,
1213 <https://doi.org/10.2307/3545749>, 1996.

1214 Smith, B., Prentice, I. C., and Sykes, M. T.: Representation of vegetation dynamics in the
1215 modelling of terrestrial ecosystems: Comparing two contrasting approaches within
1216 European climate space, *Glob. Ecol. Biogeogr.*, 10, 621–637,
1217 <https://doi.org/10.1046/j.1466-822X.2001.00256.x>, 2001.

1218 Smith, B., Wärlind, D., Arneth, A., Hickler, T., Leadley, P., Siltberg, J., and Zaehle, S.:
1219 Implications of incorporating N cycling and N limitations on primary production in an
1220 individual-based dynamic vegetation model, *Biogeosciences*, 11, 2027–2054,
1221 <https://doi.org/10.5194/bg-11-2027-2014>, 2014.

1222 Thakur, D. and Chawla, A.: Functional diversity along elevational gradients in the high
1223 altitude vegetation of the western Himalaya, *Biodivers. Conserv.*, 28, 1977–1996,
1224 <https://doi.org/10.1007/s10531-019-01728-5>, 2019.

1225 Thakur, G., Kumar, P., Bhardwaj, D. R., Prakash, P., and Poonam: Dynamics of
1226 aboveground vegetation biomass and carbon stocks along the altitudinal gradients and
1227 overstorey composition types in the temperate Himalayan region, *Trees For. People*, 16,
1228 100553, <https://doi.org/10.1016/j.tfp.2024.100553>, 2024.

1229 Thakur, R. B. and Phulara, N. K.: *A Compendium of Tree Species of Nepal*, Sarvottam
1230 Offset Printing Press (P) Ltd., 2014.

1231 Thorne, J. H., Choe, H., Dorji, L., Yangden, K., Wangdi, D., Phuntsho, Y., and Beardsley, K.:
1232 Species richness and turnover patterns for tropical and temperate plants on the
1233 elevation gradient of the eastern Himalayan Mountains, *Front. Ecol. Evol.*, 10, 1–15,
1234 <https://doi.org/10.3389/fevo.2022.942759>, 2022.

1235 Tito, R., Vasconcelos, H. L., and Feeley, K. J.: Mountain Ecosystems as Natural
1236 Laboratories for Climate Change Experiments, *Front. For. Glob. Change*, 3, 38,
1237 <https://doi.org/10.3389/ffgc.2020.00038>, 2020.

1238 Tsunoda, Y., Tsuda, T., and Ohno, Y.: Influence of stand structure on tree height: A
1239 comparative study across 46 species, *For. Ecol. Manag.*, 578, 122474,
1240 <https://doi.org/10.1016/j.foreco.2024.122474>, 2025.

1241 Whittaker, R. H.: Dominance and Diversity in Land Plant Communities: Numerical
1242 relations of species express the importance of competition in community function and
1243 evolution., *Science*, 147, 250–260, <https://doi.org/10.1126/science.147.3655.250>,
1244 1965.

1245 Zhu, L., Zhang, Y., Ye, H., Li, Y., Hu, W., Du, J., and Zhao, P.: Variations in leaf and stem traits
1246 across two elevations in subtropical forests, *Funct. Plant Biol.*, 49, 319–332,
1247 <https://doi.org/10.1071/FP21220>, 2022.

1248