Sedimentary ancient DNA reveals a threat of warming-induced alpine habitat loss to Tibetan Plateau plant diversity

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Studies along elevational gradients worldwide usually find the highest plant taxa richness in mid-elevation forest belts. Hence, an increase in upper elevation diversity is expected in the course of warming-related treeline rise. Here, we use a time-series approach to infer past taxa richness from sedimentary ancient DNA from the south-eastern Tibetan Plateau over the last ~18,000 years. We find the highest total plant taxa richness during the cool phase after glacier retreat when the area contained extensive and diverse alpine habitats (14–10 ka); followed by a decline when forests expanded during the warm early- to mid-Holocene (10–3.6 ka). Livestock grazing since 3.6 ka promoted plant taxa richness only weakly. Based on these inferred dependencies, our simulation yields a substantive decrease in plant taxa richness in response to warming-related alpine habitat loss over the next centuries. Accordingly, efforts of Tibetan biodiversity conservation should include conclusions from palaeoecological evidence.
global change affects mountain biodiversity and alters ecosystem functioning, eventually threatening the provision of ecosystem services to human society. Whether ongoing glacier retreat, treeline rise, and land-use intensification will result in an increase or decrease of high-elevation plant richness is heavily debated. Studies of elevational transects locate the highest plant richness at intermediate elevations, which are mostly found within the forest belt in temperate regions. For example, plant diversity peaks at ~3,600 m a.s.l. (metres above sea level) on the south-eastern Tibetan Plateau (Hengduan Mountains, Fig. 1a, red dotted line) which harbours one third of the vascular plant flora of China. However, it is unclear whether this widely observed hump-shaped diversity pattern represents a sampling effect (because mid-elevations are often preferentially sampled), an area effect (because mid-elevation areas are mostly overrepresented and thus can support higher plant diversity), or a mid-domain effect (because species distributions overlap mainly at the geometric centre even without environmental gradients), or whether it is really the mild temperatures and/or ecotone effect of forests that support a high richness. Hence, the question remains as to whether plant diversity at high elevations will increase with temperature-driven treeline advance, as would be expected when simply projecting an upward movement of vegetation belts.

Alpine plants, which typically include many endemic taxa, are threatened by habitat loss when treelines rise and are therefore a focus of conservation considerations. It is uncertain whether their preferred habitats or a diversity of habitats should be conserved to protect richness. It is also unclear whether landscape diversity resulting from extensive and traditional land use can, to some extent, compensate for climate-change effects on alpine plant richness.

These uncertainties mainly originate from a lack of long-term biodiversity records. Most projected mountain diversity changes are based on knowledge obtained from samples taken across a spatial extent and include various artefacts. The advantages of a time-series approach over the traditional space-for-time approach are that the sampled site is constant (i.e., normalizing for the sampling effect), that sampling elevation always represents the same portion of the investigated mountain range (i.e., normalizing for the area effect) and is always placed at the same relative elevation (i.e., normalizing for the mid-domain effect). Hence, such an approach can well reflect the temporal biodiversity-environment relationship and as such increase the effects of relevant variables when predicting biodiversity change over time. Mountain lake sediments are historical archives of ecological change, but classical vegetation proxies, such as pollen or macrofossils, are not suitable indicators for plant diversity change. However, methodological advances in sedimentary ancient DNA (sedaDNA) metabarcoding now allow an assessment of biodiversity at higher taxonomic resolution than traditional approaches through time.

Here, we reconstruct the taxonomic richness of plants in the catchment area of Lake Naleng (Hengduan Mountains, south-eastern margin of the Tibetan Plateau; Fig. 1a) over the past ~18,000 years. We apply a sedaDNA metabarcoding approach with general plant primers (Methods) to 72 horizons from a lake sediment core. We consider temperature, habitable area, forest shifts, and human impact as potential drivers of changes in plant taxa richness. In particular, we investigate whether richness increases with forest invasion into alpine habitats, recapitulating the modern elevational richness gradient. Finally, we make inferences for plant taxa richness under future climate change by applying the inferred long-term diversity-environmental relationships and provide suggestions for a future plant diversity conservation planning. We find that the total plant taxa richness is highest during late glacial times, when the area was dominated by alpine meadows, and lowest during the early Holocene when forest extent was at its maximum. By analogy to the past, we infer that total plant-taxon richness could decrease in the future due to a warming-induced upward movement of the treeline.

**Results and discussion**

**Plant DNA and taxa richness changes.** After bioinformatic filtering of raw sequencing output (Methods), 6,021,603 sequence counts were obtained from 138 PCR (polymerase chain reaction) replicates of 71 investigated sediment horizons. They were assigned to 218 terrestrial seed plant taxa with 100% best identity (Supplementary Data 1). A few PCR replicates, mainly from the late-glacial period, had to be excluded from further analyses because they were without read counts (Supplementary Fig. 1a).

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Fig. 1 The Naleng lacustrine core was recovered from the centre of Lake Naleng (black bullet point), in the Hengduan Mountains, a designated biodiversity hotspot in East Asia. a Location of the Hengduan Mountains on the south-eastern Tibetan Plateau, China (top-right inset, yellow fill). Area-elevation relationship (grey bars), elevational species richness distribution (red dotted line), and forest zone (blue dashed line) are shown in the lower-right inset. b Lake Naleng catchment area is 128 km². The simulation of the glacier extent (Methods) indicates that the Lake Naleng catchment became ice-free by about 14 ka. The extent of alpine area per 100-m elevation is shown as white bars.
Fig. 2 Long-term history of plant sedaDNA recorded in Lake Naleng compared with time-series data. a Total plant richness (red line; n = 138 PCR replicates of 71 lake-sediments, bars indicate 95% confidence intervals), b–f Comparison of sedimentary ancient DNA abundance (in %, grey polygon) of the most common alpine plant families with corresponding within-family plant richness (red line; n = 138 PCR replicates of 71 lake-sediments, bars indicate 95% confidence intervals) for Polygonaceae, Ranunculaceae, Asteraceae, Orobanchaceae, and Saxifragaceae. g The Northern Hemisphere (30°–90°N) temperature anomaly record since last deglaciation based on multiple proxies57,58 (black line with points, Methods) and percentage of Picea sedaDNA (grey polygon). h Alpine habitat area (red polygon) within the Lake Naleng catchment is the sum of pixels above the treeline (4400 m a.s.l.) based on simulated total habitat area (black outline, Methods). i sedaDNA indicators of traditional land-use including Sanguisorba officinalis (grey polygon) and Nardostachys jatamansi (black line). j The Mg/Ca ratio of Lake Naleng indicates the soil development within the lake catchment29. k Vegetation types inferred from the pollen record19,20 and sedaDNA record. l Zonation (horizontal grey lines) according to a stratigraphically constrained cluster analysis (CONISS) based on relative read abundance. Data are presented as mean ± 95% confidence interval (error bars) in a–f. Source data are provided with this paper.

Apart from 5 outliers with high read numbers, PCR replicates yielded read counts of a similar order of magnitude (Supplementary Fig. 1b). We found no correlation between read counts and total plant taxa richness (df = 69, rho = 0.014, p = 0.908). From this we conclude that read count has no impact on the inferred diversity signal. Also, neither variations in plant taxa richness nor compositional signals differed between results from single PCR samples or from pooled-PCR samples (Supplementary Fig. 1c, d, Supplementary Table 1 and Supplementary Table 2). Accordingly, we assume that the variations of plant taxa richness over time can be reliably tracked by pooling results from PCR replicates of one horizon.

Overall the sedaDNA record reproduces the compositional vegetation changes (Fig. 2k) inferred from pollen data19,20 (Supplementary Fig. 2) and pollen-based vegetation change agrees with other pollen records from the Tibetan Plateau:21,22 alpine steppe dominated 18–14 ka, alpine meadow 14–10 ka, open Picea forest 10–3.6 ka and alpine meadow after 3.6 ka with the presence of typical land-use indicators such as Sanguisorba officinalis (Fig. 2i, grey area)20 and Nardostachys jatamansi (a traditional Tibetan medicinal plant23) (Fig. 2i, black line). The sedaDNA better captures the vegetation signals within the lake catchment than pollen as it is not impacted by upward plant material transport (Supplementary Fig. 2) and records more taxa at higher taxonomic resolution than pollen spectra (Supplementary Table 3). Accordingly, and because the lake catchment covers the most common elevations in the Hengduan Mountains (~4200 to ~4900 m a.s.l., Fig. 1a, b), we conclude that Lake Naleng archived the main signal of the south-eastern Tibetan alpine ecosystem. This reasoning aligns with a modern study that indicates a non-random vegetation composition in the alpine belt of the Hengduan Mountains and identified phylogenetic clustering of alpine plant taxa in connection with environmental filtering24.

Total plant taxa richness was relatively low before 14 ka, higher between 14 and 10 ka, low again between 10 and 3.6 ka and high after 3.6 ka (Fig. 2a). Similar trends were obtained for taxa richness within important alpine plant families (Fig. 2b–f). Proportional immigration of taxa dominated during 14–10 ka and after 3.6 ka compared to their respective previous time period (Supplementary Fig. 3, Methods). Considering the restricted entries in the taxonomic database (EMBL Nucleotide Database standard sequence release 127,25, we reid the sedaDNA analyses using a 95% best identity threshold for taxa assignment which yielded 984 unique sequence types (Supplementary Data 2) indicating the possibility of additional plant taxa. Plant taxa richness patterns based on the 95% best identity are similar to those with 100% best identity (Supplementary Fig. 4), which provides confidence in our results. Furthermore, a similar temporal pattern of total plant taxa richness was obtained when analysing the data before taxonomic assignment and the data containing all terrestrial seed plant sequences (Methods), as indicated by the highly significant correlations of these time-series with total plant taxa richness (Supplementary Table 1). Sample processing-related errors (e.g., PCR and sequencing) may have slightly inflated taxa richness. However, we assume that we rather underestimate taxa richness because of non-specificity of the marker and non-completeness of the reference database, which likely means that the sedaDNA detected taxa number is lower than the absolute taxa number recorded in the flora. Additionally, species-rich families26 in the flora including Asteraceae, Saxifragaceae, and Orobanchaceae have highest richness in our record. Thus, plant taxa richness (total plant taxa richness and taxa richness within dominant alpine families) can be regarded as a semi-quantitative proxy of taxa richness in our study.

Taken together, our applied plant sedaDNA metabarcoding identifies more plant taxa at lower taxonomic level than any other
Drivers of plant taxa richness changes. Total plant taxa richness shows contrasting correlations with multiple proxy-based temperature reconstructions (Fig. 2g, dotted line) for different periods of the record: a positive correlation for 18–10 ka (rho = 0.225) but negative correlations for 14–3.6 ka (rho = −0.728) and 10–0 ka (rho = −0.932; Table 1). This suggests that temperature is unlikely to be a direct driver of plant taxa richness but may instead trigger different environmental processes that lead to contrasting biodiversity-temperature correlations.

Our analyses reveal a weak positive but statistically non-significant correlation between total plant taxa richness and total habitat area (Fig. 2h, black line) in response to glacier retreat at the end of the last glacial in the catchment of Lake Naleng (rho = 0.257, alpha level = 0.25; Table 1). Thus, we find no evidence to support the idea that total plant taxa richness mainly depends on the available area. Underestimation of this dependency might be related to limitations in our glacier-extent modelling approach that uses a proxy-based temperature reconstruction averaged from the Northern Hemisphere rather than from our study area. Nevertheless, to our knowledge, our study is the first palaeo time-series approach that addresses the extensively debated relationship between taxa richness and area27. Aside from habitable area, further processes related to rapid glacier retreat negatively impacted total plant taxa richness changes during the late glacial (rho = −0.587, alpha level = 0.025; Table 1). This might be attributed to disturbances on unstable slopes restricting vegetation establishment 28. The increase in pedogenic minerals (as indicated by sedimentary proxy Mg/Ca ratio from the same record 29, Fig. 2j) may have promoted the increase of richness of some alpine families (e.g. Polygonaceae, alpha level = 0.025; Ranunculaceae, alpha level = 0.025; Orobanchaceae, alpha level = 0.05, Supplementary Table 4), supporting the idea that soil development contributes to the coexistence of a large number of plant species 30. However, it is not the key driver for the total plant taxa richness (alpha level = 0.25, Table 1).

We find a strong positive relationship between total plant taxa richness and the alpine habitat extent (Fig. 2h, red area; which itself negatively correlates with sedaDNA signals of Pinus) in the catchment area after 14 ka (alpha level < 0.05; Table 1). Hence, in contrast to our expectation from the modern elevational plant taxa richness gradients in the Hengduan Mountains that peaks in the upper forest belt (Fig. 1a), early Holocene forest expansion into the catchment of Lake Naleng did not result in a plant taxa richness increase but in a richness decrease (alpha level = 0.01; Table 1). Accordingly, the reconstructed and simulated late-Holocene forest retreat (i.e., alpine area extent) also correlates with a richness increase (alpha level = 0.0005, Table 1). We assume that the retreat of forests is related to late Holocene cooling and weakening of the Asian summer monsoon, not to human impact, and is supported by a lack of late Holocene forest burning20.

Interestingly, sedaDNA results show that relative abundance and plant taxa richness within high-alpine plant families such as Asteraceae, Orobanchaceae, and Saxifragaceae can differ substantially, such that we find high within-family richness but low relative abundance at 14–10 ka (Fig. 2d–f). Of course, the relationship between relative read abundance and relative abundance of the taxa in the vegetation is still poorly understood and previous studies indicate that biases originate from, for example, PCR setup (e.g. preference for short reads and Table 1 Summary of correlation coefficients between total plant taxa richness and the predictor variables.

| Variable               | 18–10 ka | 14–3.6 ka | 10–0 ka |
|------------------------|----------|-----------|---------|
| Total habitat          | 0.357    | 0.054     | 0.025   |
| Temperature            | 0.025    | 0.746     | 0.002   |
| Glacier retreat        | 0.057    | 0.728     | 0.739   |
| Forested area          | 0.374    | 0.427     | 0.25    |
| Mg/Ca ratio            | 0.381    | 0.088     | 0.025   |

rho: Spearman’s rank correlation coefficient adjusted for multiple comparisons with Bonferroni adjustment. alpha level: adjusted p-value for Spearman’s rank correlation coefficient at the 0.05 significance level. For each significant variable, the table shows Spearman’s rank correlation coefficient, adjusted p-value, and the number of samples (df) for each correlation.
reads with high GC content\(^{(31)}\). However, studies of modern lake sediments have also shown that the compositional differences among sites are preserved\(^{(32,33)}\). Similarity in compositional changes between Lake Naleng sedaDNA and the pollen record supports this finding (Supplementary Fig. 2). We speculate that simultaneous high diversity of alpine habitats and maximum alpine habitat extent during 14–10 ka in the catchment of Lake Naleng may have provided habitats for many different plant taxa thereby suppressing domination by a few taxa\(^{(34)}\), which could, in turn, have created microenvironments that facilitate novel taxa migration.

We assume that the signs for late Holocene grazing intensification in sedaDNA and pollen records (Supplementary Fig. 2) are related to human impact. They indicate that human impact started or substantially increased after 3.6 ka, which aligns with archaeological evidence\(^{(35)}\). Their positive correlation with total plant taxa richness in the palaeorecord (10–0 ka, \(r = 0.939\), alpha level = 0.0005, Table 1) is consistent with the findings from experimental studies that moderate land use can increase taxa richness\(^{(34)}\). However, we find that the positive effect is smaller compared to the negative effect of alpine habitat loss due to forest invasion (10–0 ka, \(r = 0.966\), alpha level = 0.0005, Table 1).

**Potential pattern of future plant taxa richness.** In summary, we identified alpine habitat extent as the best predictor variable for total plant taxa richness (explained deviance = 96.04%, Supplementary Table 5), which was then used in a generalized linear model (Fig. 3a). The simulations project extensive alpine habitat loss in the Lake Naleng catchment area over the next 250 years in response to a predicted 2.5 °C warming (Supplementary Fig. 5), leading to a pronounced decrease in total plant taxa richness (Fig. 3b) and restricting cold-adapted taxa to high-mountain regions (Fig. 3c). In particular, taxa of endemic-rich high-alpine families are likely to disappear from the catchment (Supplementary Fig. 6). Because similar habitats will be rare in the surrounding Hengduan Mountains and the amplification of warming in high-elevation central Tibetan areas\(^{(36)}\) is unfavourable for these taxa, these taxa may become extinct. An upward expansion of montane taxa and a loss of high alpine taxa in the study area agree with predictions from a comprehensive species distribution modelling approach for the Hengduan Mountains\(^{(4)}\).

Our approach has several shortcomings. It assumes that treeline change is sensitive to temperature change in the region. Although this assumption is supported by palaeoecological evidence showing that forests expanded into higher elevations under warming during the early- to mid-Holocene and retreated to lower elevations during the late Holocene cooling\(^{(22)}\), and by modern observations of an upward shifting treeline on the southeastern Tibetan Plateau in the past 100 years\(^{(37)}\), the pace of treeline response is observed to lag the temperature warming in some mountain regions due to a variety of processes. Such processes include interspecific competition, forest-shrub interactions, dispersal variations, or even extreme climate events\(^{(37,38)}\). So, our temperature-treeline-richness relationship may therefore be correct on a millennial time scale but may overestimate changes on shorter time scales. Furthermore, our approach considers plant richness as a whole or focus on certain alpine families. Therefore, the habitat gain and loss of individual taxa or specific functional groups cannot be evaluated. Hence, our simulated taxa loss in relation to shrinking alpine habitat extent should be treated as a potential pattern by analogy to the past. It requires

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**Fig. 3 Predicted total plant taxa richness under 2.5 °C climate warming between 2050 and 2300 based on the inferred past relationship between total plant taxa richness and alpine habitat area.** a The relationship between total plant richness and alpine habitat area was established by a generalized linear model (Methods). b Prediction of total plant richness loss based on predicted alpine area habitat loss as inferred from simulating the forest rise in response to warming. c Visualization of total plant richness in Lake Naleng’s catchment after 2.5 °C warming. Data are presented as mean ± 95% confidence interval (red area) in a and b.
confirmation from a more sophisticated species-specific approach that also considers realistic migrational lags.

Implications for conservation of future plant taxa richness. With respect to conservation efforts of the unique Hengduan Mountains diversity hotspot the following conclusions can be derived from our time-series approach (Fig. 4). First, to maintain high total plant richness, nature conservation should focus on alpine habitats. This contrasts with the conclusions based on elevational gradients which suggest prioritizing forests.9. Focusing on alpine areas will also help to protect richness within alpine plant families that contain many endemic taxa as a result of Quaternary glaciations and geographical isolation.40. Second, areas harbouring extensive alpine habitat and habitat diversity in the current upper alpine zone should be protected to provide space for warming-related upward plant migration. Third, any potential positive effects from grazing management are probably too weak to compensate for climate-change impacts on plant diversity in alpine habitats. Our study indicates that time-series investigations from palaeoecological investigations using sedaDNA can inform decision-making in nature conservation by revealing potential plant responses to changing environments and, when used alongside modelling studies of modern species distributions,93 create a fuller picture of plant dynamics.

Methods

Study site. Lake Naleng (31.10° N, 99.75° E, 4200 m a.s.l.) is fed mainly via a major river channel on the northern side of the lake and drains from the southern margin. Several small streams from the adjacent mountains (up to 4,900 m a.s.l.) drain directly into the lake.29 The basin of the lake was formed by glacial activity during the Last Glacial Maximum as indicated by erratic boulders and moraines.29 The study area is in a rocking shaker overnight under the aluminium foil protection. The subsequent extraction steps followed the manufacturer’s instructions of the kit and were completed on the second day and DNA was eluted in 1.6 mL C6 buffer. For amplification, we used plant universal primers g and h targeting the P6 loop of the chloroplast tRNA (UAA) intron (Supplementary Table 6).3 To distinguish the samples after sequencing, both primers were modified by adding a 6 bp tag with at least five different base pairs between each to the 5’ end and three additional NNNs for improving cluster detection on Illumina sequencing platforms.47 Altogether, 25 µL per PCR reaction were performed with the following reagents: Primers (forward: 5′-NNN(8 bp tag)GGGCAATCCTGAGCCAA-3′; reverse: 5′-NNN(8 bp tag)TTTGTCTGCTGACACTATG-3′), with the final concentration of 0.4 µM, 1× Platinum® Taq DNA Polymerase High Fidelity PCR buffer (Invitrogen, USA), 0.25 mM dNTPs, 0.8 mg Bovine Serum Albumin, 2 mM MgSO4 (Invitrogen, USA) and 3 µL of sedaDNA template. PCRs (polymerase chain reactions) were run in the Post-PCR area separate from the ancient DNA facilities at 94 °C for 5 min (initial denaturation), followed by 50 cycles of 94 °C for 30 s, 50 °C for 30 s, 68 °C for 30 s and a final extension at 72 °C for 10 min. A no template control (NTC) was included for each PCR batch which included nine DNA extractions and one extraction control. PCR set-ups were conducted under a dedicated UV working station in the detached ancient DNA laboratory physically separated from the workplace of Post-PCR, where we did the thermal cycling, purification, and pooling. Each PCR batch was replicated until obtaining two positive PCR replicates for each lake-sediment sample when the associated controls were negative. A qualified positive PCR product was considered only if it matched two conditions: (1) the gene band is evidently longer than that of negative controls; (2) the brightest staining was in the 100–200 bp range. Specifically, the thin/blurry products below 50 bp in corresponding controls are primer dimers and not expected PCR products. PCR products were visualized with 2% agarose gel electrophoresis, purified with the MiniElute PCR Purification Kit (Qiagen, Germany), and measured with the ds-DNA BR Assay and the Qubit® 2.0 Flurometer (Invitrogen, USA) using 1 µL of PCR product. All purified PCR products were equimolarly pooled and sent away for sequencing to Fasteris SA, which used the MetaFast library protocol prior to sequencing on an Illumina HiSeq 2500 sequencing platform with paired-end reads of 123 bp length with the mode HiSeq High Output Version 4 by applying the HiSeq SBS Kit v4. Our project was sequenced together with another unknown sequencing project on a full HiSeq 2500 lane and resulted in 9.5 Gb with 37,922,797 generated clusters ≥Q30.

Fig. 4 Visualization of total plant taxa richness and effects of abiotic factors on plant richness across four time-intervals. We calculated the statistical relationship between total plant richness and predictor variables (rounded rectangle) between consecutive periods of time (Methods). Alpha significance codes of Spearman correlation are ***0.0005, **0.01, and *0.025 according to adjusted degrees of freedom. The mean annual temperature anomaly is indicated by the thermometer. Positive and negative correlation is marked in red and blue font, respectively. The catchment sketches illustrate that disturbance in the glaciated landscape was likely of importance during the deglaciation period. Our results indicate that once the catchment became ice-free, alpine habitat extent is the main driver of total plant richness while land-use is only of secondary importance during the late Holocene. From the switch in correlation sign, we assume that temperature is likely not a direct driver of richness change.

DNA extraction, amplification, and high throughput next-generation sequencing. All DNA work was carried out in ancient DNA dedicated facilities at Alfred Wegener Institute, Helmholtz Centre for Polar and Marine Research, using strict ancient DNA precautions and protocols. Each extraction batch included nine samples (3–10 g sample) and one extraction control, which was treated with a partially modified protocol of PowerMax® Soil DNA Isolation kit (Mo Bio Laboratories, Inc. USA). The isolation of DNA was first processed by loading 15 mL PowerBead solution, 1.2 mL C1 buffer, 0.8 mg proteinase K (VWR International), 0.5 mL 1 M dithiothreitol (VWR International), and samples into PowerBead tubes. Then, all tubes were vortexed in 10 min and incubated at 56 °C in a rocking shaker overnight under the aluminium foil protection. The subsequent extraction steps followed the manufacturer’s instructions of the kit and were completed on the second day and DNA was eluted in 1.6 mL C6 buffer. For amplification we used plant universal primers g and h targeting the P6 loop of the chloroplast tRNA (UAA) intron (Supplementary Table 6).3 To distinguish the samples after sequencing, both primers were modified by adding a 6 bp tag with at least five different base pairs between each to the 5’ end and three additional NNNs for improving cluster detection on Illumina sequencing platforms.47 Altogether, 25 µL per PCR reaction were performed with the following reagents: Primers (forward: 5′-NNN(8 bp tag)GGGCAATCCTGAGCCAA-3′; reverse: 5′-NNN(8 bp tag)TTTGTCTGCTGACACTATG-3′), with the final concentration of 0.4 µM, 1× Platinum® Taq DNA Polymerase High Fidelity PCR buffer (Invitrogen, USA), 0.25 mM dNTPs, 0.8 mg Bovine Serum Albumin, 2 mM MgSO4 (Invitrogen, USA), 1 U Platinum® Taq High Fidelity DNA Polymerase (Invitrogen, USA) and 3 µL of sedaDNA template. PCRs (polymerase chain reactions) were run in the Post-PCR area separate from the ancient DNA facilities at 94 °C for 5 min (initial denaturation), followed by 50 cycles of 94 °C for 30 s, 50 °C for 30 s, 68 °C for 30 s and a final extension at 72 °C for 10 min. A no template control (NTC) was included for each PCR batch which included nine DNA extractions and one extraction control. PCR set-ups were conducted under a dedicated UV working station in the detached ancient DNA laboratory physically separated from the workplace of Post-PCR, where we did the thermal cycling, purification, and pooling. Each PCR batch was replicated until obtaining two positive PCR replicates for each lake-sediment sample when the associated controls were negative. A qualified positive PCR product was considered only if it matched two conditions: (1) the gene band is evidently longer than that of negative controls; (2) the brightest staining was in the 100–200 bp range. Specifically, the thin/blurry products below 50 bp in corresponding controls are primer dimers and not expected PCR products. PCR products were visualized with 2% agarose gel electrophoresis, purified with the MiniElute PCR Purification Kit (Qiagen, Germany), and measured with the ds-DNA BR Assay and the Qubit® 2.0 Flurometer (Invitrogen, USA) using 1 µL of PCR product. All purified PCR products were equimolarly pooled and sent away for sequencing to Fasteris SA, which used the MetaFast library protocol prior to sequencing on an Illumina HiSeq 2500 sequencing platform with paired-end reads of 123 bp length with the mode HiSeq High Output Version 4 by applying the HiSeq SBS Kit v4. Our project was sequenced together with another unknown sequencing project on a full HiSeq 2500 lane and resulted in 9.5 Gb with 37,922,797 generated clusters ≥Q30.

Dating and chronology. Dating and an age-depth model are described in detail in a previous publication.42 As macrofossils were absent throughout the core, sixteen samples of bulk organic carbon were selected for accelerator mass spectrometer (AMS) 14C dating at the Leibniz Institute Kiel. The determined lake reservoir effect of 1500 years was subtracted from each 14C date, prior to calibration to calendar years (cal yr BP) using CALIB 5.0.13,44.

Material. In total, 72 sediment samples were collected from the core. In the climate chamber at a temperature of −10°C, about 2 mm of the exposed sediment of frozen samples was removed with a single-use clean blade and the inner part used for ancient DNA isolation.
Sequence analysis and taxonomic assignment. The sequence data were analysed using OBITools software48. First, the paired-end DNA sequence reads were aligned using the illumina/pairedend program in order to assemble the forward and reverse sequence reads. With the program ngsfilter paired sequences were assigned to samples based on their unique tag combination used for each sample. Afterwards read counts were summarized for unique sequence types using obunigq, and obrepq was used to discard those sequences with a length <10 bp and a total count <10 reads in total. Subsequently, obclust was used to exclude sequence variants which are likely attributed to PCR or sequencing errors by determining the sequences into head, internal, and singleton based on sequence count and similarity within one sample. Finally, sequence types were taxonomically assigned using the obseq2tax program run on two reference databases: EMBL Nucleotide Database (standard sequence release 12729) and Arctic and Boreal vascular plant and bryophyte reference libraries49–51. The applied EMBL database was created by using an in silico PCR52 with the g/h primers allowing five mismatches between primer and the targeted sequences of the EMBL entries to increase the taxonomic breadth.

sedDNA data quality control. To further denote, only those sequence types that were assigned to terrestrial seed plants and have a best identity value greater than or equal to 0.95 were kept for the following data processing. Sequence counts <10 in each sample were replaced with 0 using R software53. Subsequently, we excluded assumed contaminations from fruits, cultivars and taxa not occurring in China, which were Musaceae, PACMAD clade, Lycopercon, BOP clade, Malaceae, etc. from the dataset. The extraction and PCR blanks were mostly without any contamination, only in a single NTC sample (sample ID: NTC6, Supplementary Data 1) were some plant DNA fragments detected. However, we did not remove these sequences from its controlled samples (Supplementary Discussion). In addition, sequences were only considered as good if the corresponding ice cover could be found on the study area (Hengduan Mountains), which was aided by an international open access database54. We collected all sequence types with the best identity value of 1 and a frequency ≥2, into a dataset named bestid1, while the bestid0.95 dataset included the sequences with best id ≥0.95 and frequency ≥2. Sample ESL024 was excluded from further statistical analysis because no reads were obtained (Supplementary Data 1 and Data 2).

Glacier-extant and habitat area simulation. Several previous studies have shown that the climate at a millennial time scale on the eastern Tibetan Plateau is strongly impacted by monsoons, particularly the East Asian summer monsoon, which tracks changes in the westerlies and continental warming that are largely a function of mid- to high-latitude changes55,56. Thus, we modelled the past glacier-cover changes first (Fig. 1b). We considered the catchment area that is available for plant colonization to be ice-free. Past ice extents were estimated with the numerical ice-flow model GC2D59. We ran simulations on the present-day topography, based on a 90-m resolution SRTM digital elevation model. Climate was imposed through a vertical mass-balance profile that we estimated from present-day conditions. Based on the spatially averaged mean elevation of present-day glaciers, we estimated a present-day equilibrium line altitude (ELA) of ~5200 m61. We estimated the maximum ice accumulation rate to be 0.25 m yr

Statistical analyses. All statistical calculations were carried out using the R software53. All correlations were computed using the cor.test method = “ spearman” in the psych package46. The Spearman Rank Correlation (rho), non-adjusted/probability values (p-value) were computed using the corr.test function and sample size (n) were obtained. Both p-values are helpful to check the significance of correlation under the unadjusted degrees of freedom, which is equal to sample size (n) minus 2.

As plant taxa richness increases with read counts, we rarefied the data bestid1 to equal counts for each sample based on the minimal total read count occurring in the entire sedDNA dataset (base count = 11,949) 100 times (Supplementary Code 1), as well as the data bestid0.95 (base count = 13,344). To investigate potential methodological biases of the sedDNA-based plant taxa richness, two additional datasets were set up: (1) metacoding data before taxonomic assignment (hereafter referred as non-ecotag); (2) all terrestrial seed DNA sequences without further sequence filtering (hereafter referred to as Seq data). Both datasets were rarefied to their respective minimal total read count (16,209; 14,645) 100 times. We investigated whether plant taxa richness is correlated to read counts for both datasets and whether it is correlated with total plant taxa richness of bestid1 dataset using cor.test(adjust = “none”). The correlation was conducted using the bestid1 dataset which collected the deeply sequenced PCR product for each lake sediment sample from the study area (hereafter referred as single data). Then, we rarefied these data 100 times based on the minimal read count (6,339) across all samples and calculated the plant taxa richness. The correlation between plant taxa richness from data bestid1 and single data was calculated using cor.test(adjust = “none”). Both correlation analyses and Proven analyses were applied to check whether sample scores and taxa scores of the first two PCA (principal component analysis) axes of the single data match those of the bestid1 dataset. Only those sequences with a maximum relative read abundance of 0.25% at least were kept. Double-square root transformation was applied before PCA analysis57. The procrustes(), test() and rd(=FALSE) are available in the vegan package48.

In order to calculate the plant taxa richness for each single taxonomic family, we divided sedDNA sequence data into subsets of taxonomic families and rarefied these subsets to a cut-off value of 100 total read counts to minimize the effect of rarefaction on taxonomic abundance. Such richness data is correlated with low read counts (e.g., Ranunculaceae) could be compared with rich taxon richnesses of families having higher read counts (e.g., Asteraceae). Furthermore, four distinct vegetation zones were classified using the cutclus method = “CONISS” in vegan package58 based on the rarefied relative abundance. We summed up the samples in each zone and computed the mean relative value of read counts per zone. Then, we rarefied the zonal data to its minimal total read count (40,377) 100 times. Afterwards, we computed the total vegetation turnover (beta diversity) using the turnover() function in the coden package49 based on the rarefied zonal data (Supplementary Code 2).

To identify the main drivers of plant taxon richness, we calculated the correlation coefficient between plant taxon richness and driver variables using cor.test(adjust = “Bonferroni”) (Supplementary Code 3 and Supplementary Data 3). We separated the complete richness-time series into three time-intervals each consisting of two consecutive vegetation zones (according to CONISS), i.e., 14–15 ka, 14–3.6 ka, 10–0 ka. This approach accounts for variation of driver importance throughout the record and can even reveal sign changes in the relationship between driver variables and plant taxa richness. We used the smoothed data for plant richness and two predictor variables (Mg/Ca ratio and land-use) in the processing of correlation calculation, so that we adjusted the degrees of freedom to get the effective independent value of the smoothed data. We used gam() for land-use data smoothing as it included a large number of zeros and lowess(span = 0.5) to smooth the Mg/Ca ratio and plant taxa richness. Consequently, to obtain the alpha level we compared rho values with exact critical values of Spearman’s rho according to adjusted degrees of freedom. Only an alpha level ≤0.05 is considered statistically significant. The results of correlation are summarized in Table 1 and Supplementary Table 4. The related code is available in Supplementary Code 4. A generalized linear model (GLM) was built using glm (family = “gaussian”) for total plant taxa richness and those families that are significantly correlated to the predictor variables (alpine habitat area and land-use indicator) during 10–0 ka (Supplementary Code 5). This period was selected as it covers the warming and most modern phase of the record. Moreover, the correlation between total plant taxa richness and alpine habitat is highest in this time interval. The temporal resolution of the correlated time-series was about 250 years. The proportion of deviance explained by the GLM was calculated using Deviance was run on the modelA package. We predicted the results of the within family richness for 2050–2300 C.E. in 50-year time steps using glm.predict() based on the most important predictor variable (alpine habitat area) in the GLM models (Supplementary Code 5). The variable importance was calculated using varimp() in caret package51. The within-family sedDNA data was analysed using the same data processing.
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