Surface soil phytoliths as vegetation and altitude indicators: a study from the southern Himalaya

Xiaohong An1,2, Houyuan Lu2,3 & Guoqiang Chu2

Phytoliths represent one of the few available altitudinal vegetation proxies for mountain ecosystems. This study analyzed 41 topsoil phytolith samples collected from five altitudinal zones in the southern Himalaya as far as, and beyond, the timberline, from tropical forest (up to 1,000 m a.s.l.) to subtropical forest (1,000–2,000 m a.s.l.), to temperate forest (2,000–3,000 m a.s.l.), to subalpine forest (3,000–4,100 m a.s.l.) and finally to alpine scrub (4,100–5,200 m a.s.l.). The statistical results show a good correlation between phytolith assemblages and these five altitudinal vegetation zones: the five phytolith assemblages identified effectively differentiated these five altitudinal vegetation zones. In particular, coniferous phytoliths accurately indicated the timberline. Additionally, we tested the phytolith index Ic (a proxy for estimating the percentage of Pooideae vis-à-vis the total grass content) as a quantifier of phytolith variety versus altitude. Ic increased along altitude, as expected. An investigation of phytoliths provided an initial basis for the analysis of the composition of gramineous vegetation. Furthermore, redundancy analysis and discriminant analysis also suggested a significant correlation between phytolith assemblages and altitude. Our research therefore provides an up-to-date analogue for the reconstruction of changes to palaeovegetation and palaeoaltitude in mountainous areas.

Plateau palaeoaltitude has attracted considerable scientific attention, in particular because plateau uplift is an external expression of tectonic movements and a principal factor influencing climate change1,2, both of which are geoscientifically significant. However, there are few palaeoaltitude indicators for mountain ecosystems. Although pollen can be an effective indicator, and has been used in estimating palaeoaltitude in the Mercantour Massif, the eastern European Alps, the eastern Pyrenees, the Sila Massif, and the northern and central Apennines3, difficulties have arisen in distinguishing different species of the same plant family3,4, and in clarifying the complex transportation and deposition of pollen-spores3,5, which is vital for palaeoaltitude estimation. In contrast, phytoliths are more resistant than pollen grains to biogenic or physical forces during their deposition, demonstrating that phytolith analysis serve as both an efficient and a complementary route for the study of palaeoaltitude.

Phytoliths are hydrated silicon particles formed in plant growth and preserved in soils and sediments after plant tissues have decayed6. They have been proven to be reliable indicators in palaeovegetation and palaeoenvironment reconstruction7–9. In mountainous areas like the Himalaya, phytoliths and pollen grains may complement each other, because some deficiencies in pollen-spores can be countered by the presence of phytoliths in the study of the relation between montane vegetation and altitude. First,
most phytoliths are naturally resistant to strong weathering and are therefore well-preserved in terrestrial sediments, where pollen-spores can be easily destroyed. Second, even though phytoliths can be transported by the wind, gravity-aided in situ deposition remains their dominant modus operandi. Third, phytoliths are more reliable indicators for differentiating varieties of grass subfamily and genera than pollen-spores. Besides, as the study of phytolith morphology progresses, the potential of phytoliths to differentiate various microenvironments, especially near the timberline, has become increasingly apparent.

There has been some limited research focusing on morphology and phytolith types in mountainous areas, but, as far as we know, only a very few studies of the relation between phytolith composition and altitudinal vegetation belts, e.g. on tropical African mountains, have been published. More international research is required. We chose the Himalaya, specifically the southern Qinghai-Tibet Plateau (QTP), for our research.

The QTP averages >4,000 m a.s.l., rising to 8,848 m a.s.l. at Mount Everest (Qomolangma). With its typical mountain climate ecosystem, it is a key area in the scientific research into Asian, and even global, climate change. The uplift of the QTP is also crucial to a better understanding of global tectonics. The study of the QTP’s palaeoaltitude could be significant in both regards, but there remain no direct records of the local palaeoaltitude, palaeovegetation or palaeo-timberline.

The Himalayan Mountains form the southern margin of the QTP and the northern boundary of the Indo-Gangetic Plain (Fig. 1). The southern Himalaya provides environments for various vegetation types, ranging clearly from tropical rainforest to perpetual frost along the altitudinal gradient. It is an optimum area for the study of variations in vegetation, as well research into indicators of QTP palaeoaltitude. Several classification systems are already available for the vegetation of Nepal, which lies on the mountain range's southern flank. Notably, Dobremez et al. developed a six-category bio-climatic (altitudinal) classification of vegetation. The published data have focused on flora classification and the characterization of plants, and the area's flora and vertical vegetation zones are therefore well-documented. However, due to international borders and poor accessibility, no known work on indexing vegetation belts along the tropical rainforest to perpetual frost altitudinal gradient has been conducted, even if the vertical vegetation range and climate change gradient are most marked in this area. It is therefore imperative to establish a useful index for the reconstruction of both palaeovegetation and palaeoaltitude in this region.

We obtained a diverse collection of samples from a wide range of vegetation belts from the southern Himalaya. In this study, we aimed to explore variations in the composition of phytolith assemblages, and
verify our hypothesis that phytolith assemblages can indicate and differentiate vegetation zones along an altitude gradient, providing the basic data necessary for the reconstruction of palaeovegetation and palaeoaltitude in mountainous areas.

Results
Vegetation description and material. The Himalaya exhibit typical montane altitudinal vegetation belts. Based on previous work, mainly Dobremez’s altitudinal classification23, this paper classifies local forest vegetation vertically from bottom to top into six formations (Fig. 2).

Tropical moist lowland Indo-Malayan forest (<1,000 m a.s.l.) (Fig. 2a). Shorea robusta (Sal) is predominant in this belt. Acacia catechu and Dalbergia sissoo replace Sal in riverine forests. Other dominant broadleaved evergreen forest types include Cycas pectinata, Gnetum montanum, Calamus spp., Cythea spinulosa, Arecaceae, Magnoliaceae and Pandanus nepalenses, except for some coniferous forests (Pinus roxburgii) on southern slopes at higher altitudes. In the foothills of western Nepal, Sal forest is replaced by Terminalia/Anogeissus forest. This zone consists of ~2,000 species of flowering plants and ~80 species of Pteridophytes. The common understory grasses include Zingiberaceae, Acanthaceae, Commelinaceae in addition to tall bamboos and grass. Grassland emerges where rainforest has been cut down. Grassland is composed of the tall grasses Saccharum arundinaceum, Apluda mutica, Themeda Forssk, et al. Rice, maize and banana are cultivated in this assemblage21,26.

Subtropical forest (1,000–2,000 m a.s.l.) (Fig. 2b). This consists of species such as Schima wallichii, Castanopsis indica and Castanopsis tribuloides in relatively humid areas, and Pinus roxburghii forests in drier regions. >1,900 flowering plant species grow vigorously in this zone27. Sinarundinaria nitida accounts for a large proportion of the total shrub content. The grasses Carex changmuensis, Arisaema sp., Ophiopogon bodinieri, Elatostema surculosum, Miscanthus sp. et al. are also found in these forests26. Cultivation is common in this belt. In some places, natural forests have been almost entirely replaced by cultivated crops such as Musa paradisiaca and Chaenomeles sinensis. Owing to the hot and rainy summer, as well as the dry and warm winter, there are many paddy fields in this zone28.

Temperate forest (2,000–3,000 m a.s.l.) (Fig. 2c). This zone is dominated by plants such as Quercus lamillosa and Q. semicarpifolia, forming pure or mixed broadleaved evergreen forest. This latter category can be subdivided into lower temperate mixed broadleaved forest, upper temperate broadleaved forest, upper temperate mixed broadleaved and coniferous forest22. Bamboo grows in sunny areas in this
zone, and *Sinorundinaria nitida* is abundant\(^{28}\). *Ainsliaca latifolia*, *Ophiopogon intermedus*, *Pilea racemos*, *Arisaema*, *Zingiber* and *Thalictrum* are common understory grasses\(^{28}\).

**Subalpine forest (3,000–4,100 m a.s.l.)** *(Fig. 2d)*. This zone is inhabited by forest vegetation up to the timberline, with >1,400 flowering plants\(^{27}\). *Betula-Rhododendron campanulatum* and *Abies spectabilis* forests are representative of the vegetation of this zone. *Rhododendron* spp. forms mixed forest within *Abies* or *Betula* forest, or occurs as a component of areas of open shrub. *Juniperus* spp. grows in the drier forest areas of this zone. Herbaceous plants growing in this zone include *Fagopyrum dibotrys*, *Abies* spp., forms mixed forest within *Juniperus communis*, *Rhododendron anthopogon*, *Rhododendron lepidotum*, *Ephedra gerardiana* and *Hippophae tubetana*. *Primula* spp., *Gentiana* spp., *Corydalis* spp., *Saussurea* spp., *Carex* spp., *Ptilagrostis* spp., *Poa* spp., *Deyeuxia* spp., *Festuca* spp., *Danthonia* spp. and *Helictotrichon* spp. are common and important grasses in this zone\(^{28}\).

**Alpine shrub (4,100–5,200 m a.s.l.)** *(Fig. 2e)*. Alpine shrub is characterized by the presence of various stunted bushy shrubs, including *Juniperus* spp., *Rhododendron* spp., *Juniperus recurva*, *Juniperus indica*, *Juniperus communis*, *Rhododendron anthopogon*, *Rhododendron lepidotum*, *Ephedra gerardiana* and *Hippophae tubetana*. *Primula* spp., *Gentiana* spp., *Corydalis* spp., *Saussurea* spp., *Carex* spp., *Ptilagrostis* spp., *Poa* spp., *Deyeuxia* spp., *Festuca* spp., *Danthonia* spp. and *Helictotrichon* spp. are common and important grasses in this zone\(^{27}\).

**Perpetual snow (>5,200 m a.s.l.)** *(Fig. 2f, g)*. This zone is composed of permanent snowfields, rocks, glaciers and ice on the high Himalayan ranges to the north. The area is barren, with lichens on exposed rocky places and a few hardy flowering plants, such as *Stellaria decumbens*. The main vegetation type on the northern slopes of the Himalaya is grass, reflecting the cold-dry climate of the QTP. Forests can be found only in valleys below 4,100 m a.s.l.\(^{28}\).

Table S1 shows the coordinates, altitudes and the principal flora of the sampling sites.

**Phytolith types.** Of the 47 processed samples, 41 contained >300 phytoliths. These were identified and classified into the 27 common categories listed below, mainly following the classification system used by Lu et al.\(^{30}\), but with reference to the classification systems of Wang and Lu\(^{31}\), Kondo et al.\(^{32}\) and Twiss et al.\(^{33}\), and using the International Code for Phytolith Nomenclature 1.0\(^{34}\).

The woody phytolith types include globular, abbreviated stellate, cylindrical sclereid and Gymnosperm. Globular types were subdivided into globular echinate (Fig. 3a) and global granulate (Fig. 3b), the former being produced specifically in Palmae and the latter in tropical trees and shrubs (but principally in Palmae in China)\(^{14,35,36}\). Abbreviated stellate (jigsaw) phytoliths (Fig. 3c) are produced by evergreen broadleaved plants\(^{13,31,37,38}\). The cylindrical sclereid (Y-shape) phytolith (Fig. 3a) is a type particular to broadleaved plants\(^{31,38}\). Gymnosperm (Fig. 3w), as the term suggests, derives mainly from Pinaceae\(^{39,40}\).

A diversity of gramineous phytolith types were observed during the course of this study. Bilobate short cell (dumbbell)\(^{30}\) (Fig. 3d) and cylindrical polylobate (multilobe) phytoliths (Fig. 3f) are both representative of Panicoideae, which adapt to warm-humid conditions\(^{31}\). The cross-shaped\(^{41}\) phytolith (Fig. 3c) is typical of the bilobate short cell type; the form of this phytolith produced in maize can be differentiated from the type produced in wild grass by its mirror-image and greater width (usually >12.5 \(\mu\)m)\(^{42,43}\). The cross-shaped type referred to in this paper specifically represents maize, with other variants classed as bilobate short cell phytoliths. Square saddle (short saddle)\(^{30}\) phytoliths (Fig. 3h) are mainly found in Chloridoideae, with a small fraction occurring in Arundinoideae. Chloridoideae mainly grows in dry-hot conditions. Arundinoideae covers a broad range of southern China. The square saddle type is considered representative of C\(_4\) plants\(^{31,38}\). Oblong concave saddle (long saddle)\(^{40}\) phytoliths (Fig. 3l) are produced in Bambusoideae, which grow in hot, moist climates throughout southern China\(^{31,34}\).

Cuneiform bulliform cell (fan-shaped)\(^{30}\) (Fig. 3o–3q) and parallelepiped bulliform cell (square and rectangular)\(^{30}\) (Fig. 3u, v) phytoliths develop from motor cells produced in Panicoideae, Oryzoideae and Bambusoideae\(^{41}\). These plants flourish in the warm and humid climate of southeastern China. Sometimes an apparently parallelepiped bulliform cell can actually be the side-on view of a cuneiform bulliform cell\(^{31}\). Some Chloridoideae produce parallelepiped bulliform cells, but not cuneiform bulliform cells. Cuneiform bulliform cell-rice (Fig. 3o) and cuneiform bulliform cell-bamboo (Fig. 3p) phytoliths can be differentiated from other cuneiform bulliform cell phytoliths by the ornamentation of cracks and spines along their front edges, respectively\(^{31,45}\).

The hair cell (point)\(^{30}\) phytolith type (Fig. 3t) develops from spiny grass cells, which resist cold and drought. Hair cell phytoliths thrive in northern and northwestern China\(^{31}\), rondel (Fig. 3g), trapeziform (Fig. 3r, s) and *Stipa*-bilobate short cell (Fig. 3e) phytoliths are produced in Pooidae\(^{38}\), and are representative of cold climates and high altitudes within tropical regions. *Stipa*-bilobate short cell phytoliths can be differentiated from Panicoideae bilobate short cell phytoliths by their slim necks and differing opposite sides\(^{15,46,47}\).
In addition to the abovementioned, the Pteridophyte phytolith type (Fig. 3ab) is particular to ferns, and the sedge (papillae) type (Fig. 3m,n) is typical of Cyperaceae, a grass-like plant growing in wet places.

We identified some types with no current taxonomical significance: the one-horned tower (Fig. 3k) and two-horned tower (Fig. 3l) are small phytolith types, exhibiting a diversity of shapes; the elongate type, including elongate smooth (Fig. 3z) and elongate echinate (Fig. 3y) phytoliths, develop from long epidermis cells (these increase in quantity in China from south to north and from humid to dry regions); and the goblet (nubby-irregular shape) type, a non-gramineous phytolith, appears in arid areas in China. We have classified these difficult-to-categorize types as unknown and as-yet-uninvestigated (see Supplementary Fig. S5 online).

Supplementary Table S2 (online) is a summary of phytolith types, descriptions and their source plants and ecoenvironments. More plates of the aforementioned phytolith types can be found in Supplementary Figs S1-S5 (online).
Phytolith assemblages. Phytoliths were divided into the following five assemblages, according to phytolith type and percentage of total composition (Fig. 4).

Assemblage I, from tropical lowland evergreen broadleaved forest (<1,000 m a.s.l.), is characterized by a high percentage of cylindrical sclereid, globular and cuneiform bulliform cell phytoliths. The maximum contents of abbreviated stellate, cylindrical sclereid, globular and cuneiform bulliform cell phytoliths are 1.8%, 7.3%, 3.9% and 19.6% of the total phytolith content, respectively. The highest numbers of broadleaved types (cylindrical sclereid and abbreviated stellate) are found in this belt, where evergreen broadleaved plants predominate. The volumetric production of Palmaceae phytoliths implies tropical lowland, high temperature conditions. Parallelepipedal bulliform cell 1, bilobate short cell, and oblong concave saddle types account for 11.6%, 5.2%, 3.4% and 4.3% of the content total, respectively. The low contents of gramineous types such as bilobate short cell and oblong concave saddle phytoliths indicate a weak growth of grass and bamboo in low altitude areas with abundant tree cover. Sedge (2.5%) is at its maximum in this zone, implying a lowland, humid environment. There is also a high quantity of unknown types (4.8%); many varieties from this wide range of phytolith morphotypes have yet to be investigated. The emergence of cuneiform bulliform cell-rice (0.5%) and maize cross (0.5%) phytoliths implies that rice and maize were cultivated in this low altitude zone.

Assemblage II, from subtropical broadleaved forest (1,000–2,000 m a.s.l.), is characterized by cylindrical sclereid, bilobate short cell and cuneiform bulliform cell phytoliths. The higher percentages of bilobate short cells (15.8%) and oblong concave saddles (11.8%) vis-à-vis total content in this assemblage compared to Assemblage I suggests a greater quantity of grasses and bamboo. Frequent and repeated human cultivation has resulted in a high grass content. In contrast with gramineous types, Palmaceae (0.9%) and cylindrical sclereid (2.6%) phytoliths decrease markedly, and abbreviated stellate types decline sharply from 1.8% to 0.3% of total content, but cylindrical sclereid remains the principal woody type. Both falls in overall content are likely to be the result of a decrease in woody plants due to human cultivation.

Assemblage III, from warm-temperate mixed forest (2,000–3,000 m a.s.l.), is characterized by saddle, parallelepipedal bulliform cell and rondel phytoliths. The increase in rondel (2.5% to 5.8%) and trapeziform (0.8% to 2.4%) phytoliths, and the decrease in bilobate short cells (15.8% to 4%), probably reflects the change in climate from warm to cold, and in altitude from low to high. The high altitude type of Stipa-bilobate short cell phytolith appears. Cuneiform bulliform cell-rice and cross-shape phytoliths were not observed.
Assemblage IV, from subalpine cold-temperate needle-leaved forest (3,000–4,100 m a.s.l.), is characterized by gymnosperm-type, trapeziform and *Stipa*-bilobate short cell phytoliths. Gymnosperm-type phytolith content increases from 0.2% to 8.6%, *Stipa*-bilobate short cell phytolith content from 2.3% to 5.6%, and trapeziform phytolith content from 2.4% to 16.2% vis-à-vis Assemblage III, while oblong concave saddle phytoliths decrease from 16.9% to 1.3%, and parallelepipedal bulliform cell 1 types decline from 15.1% to 4.8%, indicating flourishing gymnosperm forest and understory Pooideae environments, both of which are indicative of high altitude conditions.

To summarize, an abundance of phytolith types, accompanied by clear variations in phytolith percentages vis-à-vis total phytolith content, occurs along the altitudinal gradient on the slopes of the southern Himalaya. Phytolith assemblages can be clearly differentiated (Table 1).

### Table 1. Correlation between phytolith assemblage and vegetation zone.

| Altitude       | Vegetation Zone                                         | Understory Grasses                                                                 | Phytolith Assemblage                                      |
|----------------|---------------------------------------------------------|-----------------------------------------------------------------------------------|----------------------------------------------------------|
| <1,000 m a.s.l.| Tropical lowland evergreen broadleaved forest          | Zingiberaceae, Acanthaceae, Commelinaeace, Saccharum arundinacum, Aphela mutica, Themeda Portik et al. | Cylindrical sclereid/globular/cuneiform bulliform cell assemblage |
| 1,000–2,000 m a.s.l. | Subtropical broadleaved forest                           | Carex changmuensis, Arisaema sp., Ophiopogon bodinieri, Elatostema auriculatum, Miscanthus sp. et al. | Cylindrical sclereid/bilobate short cell/cuneiform bulliform cell assemblage |
| 2,000–3,000 m a.s.l. | Warm-temperate mixed forest                             | Ainiluca latifolia, Ophiopogon intermedia, Fleea racemosa, Arisaema, Zingiber and Thalictrum et al. | Saddle/parallelepiped bulliform cell/rondel assemblage |
| 3,000–4,100 m a.s.l. | Subalpine cold-temperate needle-leaved forest           | Fagopyrum dibotrys, Carex lacta, Elatostema obtusum, Arisaema jasquemontii, Poa crymophila, Deyeuxia scabrescens, Chernoopodium sp., Oryza polyst安心, Arundinella nepalensis, Anthraxon sp., Pennisetum flaccidum et al. | Gymnosperm-type/trapeziform/Stipa-bilobate short cell assemblage |
| 4,100–5,200 m a.s.l. | Alpine shrub                                           | Primula spp., Gentiana spp., Corydalis spp., Saussurea spp., Kobresia spp., Carex spp., Phalacrostis spp., Poa spp., Deyeuxia spp., Festuca spp., Danthonia spp. and Helictotrichon spp. et al. | Trapeziform/rondel/gobbet assemblage |

Assemblage IV, from subalpine cold-temperate needle-leaved forest (3,000–4,100 m a.s.l.), is characterized by gymnosperm-type, trapeziform and *Stipa*-bilobate short cell phytoliths. Gymnosperm-type phytolith content increases from 0.2% to 8.6%, *Stipa*-bilobate short cell phytolith content from 2.3% to 5.6%, and trapeziform phytolith content from 2.4% to 16.2% vis-à-vis Assemblage III, while oblong concave saddle phytoliths decrease from 16.9% to 1.3%, and parallelepipedal bulliform cell 1 types decline from 15.1% to 4.8%, indicating flourishing gymnosperm forest and understory Pooideae environments, both of which are indicative of high altitude conditions.

Assemblage V, from alpine shrub (4,100–5,200 m a.s.l.), is characterized by trapeziform, rondel and gobbet phytoliths. The percentages of most types drop very low in this assemblage, but there are abundant rondel (5.6%), trapeziform (30.6%) and gobbet (6.9%) phytoliths, indicating the predominance of high altitude, cold climate-adaptable plants.

To summarize, an abundance of phytolith types, accompanied by clear variations in phytolith percentages vis-à-vis total phytolith content, occurs along the altitudinal gradient on the slopes of the southern Himalaya. Phytolith assemblages can be clearly differentiated (Table 1).

### Floristic composition (RDA results) 

Floristic composition (RDA results). 18 of the aforementioned 27 phytolith types account for a certain proportion of the samples and clearly indicate vegetation type. In elucidating the relation between phytolith assemblages and altitude, redundancy analysis (RDA) results (Fig. 5) show that the first and second axes account for 42% and 16% of the total variance, respectively, describing 58% of the information in total. The other axes exhibit very low values. This suggests that phytolith composition is controlled by environmental factors represented by the first two axes, and especially by altitude (the first axis).

As the RDA results reveal, low altitude-adaptable phytolith types are grouped in a positive direction toward the first axis, while those adapting to high altitude point in a negative direction. The former group includes globular, cylindrical sclereid, bilobate short cell, parallelepipedal bulliform cell, cuneiform bulliform cell, cuneiform bulliform cell-bamboo, oblong concave saddle and square saddle phytoliths; the latter is composed of gymnosperm-type, hair cell, elongate smooth, elongate echinate, rondel, trapeziform, *Stipa*-bilobate short cell and gobbet phytoliths (Fig. 5a). Low altitude samples t1–t20 are positively grouped toward the first axis, while high altitude samples t21–t41 exhibit a negative direction (Fig. 5b). This reflects the clustering of phytolith samples from higher altitudes around negative coordinates, in
In contrast to samples from lower altitudes, which are clustered around positive coordinates. This therefore demonstrates a correspondence between phytolith assemblages and the altitudinal distribution of vegetation.

**Floristic composition (DA results).** Discriminant analysis (DA) was applied to test whether the established surface phytolith assemblages described in this paper can reliably differentiate the vertical vegetation belts in the southern Himalaya. Using *a priori* groups, the 41 surface samples with the 18 phytolith types used for RDA were then classified to co-validate the classification of phytolith assemblages. 39 samples (95%) were correctly classified with respect to these *a priori* groups (Table 2). The first two discriminant function scores are illustrated herein: group centroids are distinctly separate (Fig. 6). The classification function coefficients are shown in supplementary Table S3 (online).

**Phytolith indices.** In order to present a direct and quantifiable relation between altitude and phytolith type, we applied the phytolith indices I_{c} and I_{ph}^{17}. I_{c} represents the proportion of short cell phytoliths from Pooidae relative to total short cell phytoliths from Pooidae, Chloridoideae and Panicoideae^{17}, thus:

\[
I_{c} = \frac{\text{rodel + trapeziform + Stipa-bilobate short cell}}{\text{rodel + trapeziform + Stipa-bilobate short cell + saddle + bilobate short cell + cross}} \times 100
\]

I_{c} (%) represents an index for the total percentage of Pooidae grass cover.

I_{c} has been shown to depend principally on altitude (SPSS curve estimation of R^2 = 0.8) (Fig. 7a). As a result, it is clear that, along with increasing altitude, Pooidae grasses increase gradually, while Chloridoideae and Panicoideae grasses decrease. The relation between altitude and I_{c} can be expressed using the equation:

---

**Table 2. DA results for the 41 surface samples extracted from the five altitudinal vegetation zones.** (95.1% of originally-grouped cases correctly classified).

| Actual Group                                      | Group No. | 1     | 2     | 3     | 4     | 5     | Total |
|---------------------------------------------------|-----------|-------|-------|-------|-------|-------|-------|
| Tropical lowland evergreen Broadleaved forest     | 1         | 10 (90.9%) | 1 (9.1%) | 0     | 0     | 0     | 11    |
| Subtropical Broadleaved forest                    | 2         | 1 (20.0%) | 4 (80.0%) | 0     | 0     | 0     | 5     |
| Warm-temperate mixed forest                       | 3         | 0     | 0     | 4 (100.0%) | 0     | 0     | 4     |
| Subalpine cold-temperate needle-leaved forest     | 4         | 0     | 0     | 0     | 15 (100.0%) | 0     | 15    |
| Alpine shrub                                      | 5         | 0     | 0     | 0     | 0     | 6 (100.0%) | 6     |

---

**Figure 5. RDA results for principal phytolith types (a) and sampling sites (b).**
where \( x \) represents \( I_c \) (%) and \( y \) represents elevation (m).

\( I_c \) is a climatic index for temperature, because a high \( I_c \) represents higher quantities of Pooideae, which adapt to the cool climates prevalent at high altitudes\(^4\). Altitude is the most influential factor vis-à-vis temperature in the Nepal Himalaya; mean annual temperature in general gradually decreases northward as altitude increases\(^5\). The \( I_c \) results therefore render an approximate, linear correlation between Pooideae amount, temperature and altitude\(^6\).

\( I_{ph} \) was also tested, but the result of the curve estimation was less useful, because the sig. = 0.109 for curve estimation (Fig. 7b).

**Discussion**

We have obtained samples from the world’s highest mountain range, the Himalaya, and have derived significant results from our analysis of the material. Although the composition of preserved phytoliths in soils is influenced by translocation and dissolution\(^7\), phytolith assemblages in soils reflect the composition of local vegetation\(^8\). First, the surface soil phytoliths described in this paper are mainly bleached mountain spodosols, dark-brown mountain earths and brown mountain earths\(^9\). These weakly acidic
soils are suitable depositional environments for phytoliths; pH ranges between four and six, and is therefore never high enough for the sustained dissolution of phytoliths. Even if dissolved, dissolution rates are similar for most phytolith morphotypes\(^5^3,5^4\). Second, small phytoliths may be dispersed by the wind\(^5^5\). However, the geographical scale of transfer versus study area is considered insignificant in this paper. Most probably, therefore, phytolith composition accurately reflects phytolith composition for each corresponding vegetation belt.

It should however be recognized that the percentage content of each phytolith vis-à-vis total phytolith content cannot be equal to the percentage content of the corresponding vegetation. Some plants yield abundant phytoliths, resulting in over-representation. So in describing vegetation assemblages, it is imperative we focus on the relative variation in each phytolith type versus altitude. Furthermore, the taphonomy of phytoliths in sediments should be considered when reconstructing palaeovegetation\(^1^0\).

In view of the reliability of our samples, the phytoliths analyzed in this paper can accurately describe vegetation zones. Phytolith percentages clearly illustrate that each vegetation zone has a distinctive phytolith assemblage and some typical phytolith types. In tropical forest at low altitudes (<1,000 m a.s.l.), the cylindrical sclereid/globular/cuneiform bulliform cell phytolith assemblage corresponds to an abundance of hot climate-adaptable woody plants, such as *Shorea*, *Cycas*, *Magnoliaceae*, *Palmae*, and *Panicoideae*, principally *Saccharum*, *Apluda* and *Themeda*. In the subtropical broadleaved forest belt (1,000–2,000 m a.s.l.), the cylindrical sclereid/bilobate short cell/cuneiform bulliform cell assemblage highlights the dominance of broadleaved woody plants and tall *Panicoideae* grasses (e.g. *Miscanthus*). In warm-temperate mixed forest (2,000–3,000 m a.s.l.), the saddle/parallelepiped bulliform cell/rondel phytolith assemblage corresponds to the mixture of warm and cold temperate zone grasses, as well as a mixture of broadleaved and coniferous plants. In the subalpine needle-leaved forest zone (3,000–4,100 m a.s.l.), the cylindrical bulliform/bilobate short cell/cuneiform bulliform cell assemblage suggests the dominance of Gymnosperm (*Abies* and *Pinus*) and low temperate gramineous types (*Deyeuxia* and *Oryzopsis*). In the alpine shrub zone (4,100–5,200 m a.s.l.), an abundance of rondel, trapeziform and goblet phytoliths implies an absence of woody plants and a predominance of high altitude-adaptable Pooideae, including *Carex*, *Festuca*, *Helictotrichon*, and especially *Ptilagrostis* (belonging to Stipeae).

To corroborate the increasing/decreasing trend in different phytoliths along altitude, typical bilobate short cell, cuneiform bulliform cell and trapeziform phytolith types are shown in boxplots, with emendations (Fig. 8). It is clear that the bilobate short cell and cuneiform bulliform cell types decrease with altitude, while the trapeziform type increases.

DA results, with their clearly-separated group centroids, also demonstrate that the five phytolith assemblages can accurately delineate vertical vegetation zones, as each vegetation belt is represented by a distinctive phytolith assemblage. Our classification of phytolith assemblages suggests that phytoliths can be used as valid proxies for subdividing montane vegetation in the QTP area, and can thus be used effectively in reconstructions of palaeo-vegetation and palaeo-altitude.

Notably, some phytoliths exhibit a high degree of sensitivity to environment and climate. In particular, the fluctuation of the timberline is crucial to montane phytolith study. It not only reflects variations in vegetational composition, but, more importantly, indicates changes in altitude. The upper altitudinal limit of montane needle-leaved plants is commonly considered to be the same as the timberline\(^5^6\). However, most conifer pollens have two or more sacs, e.g. pine pollen. Long-distance dispersal of pollens leads to their wide representation in assemblages. Investigations of modern pollens have shown that pine forest only grows where conifer pollens account for at least 30% of the total\(^4\), so the existence of conifer pollens does not necessarily accurately reflect the proportion of coniferous forests. Autochthonous or proximal deposition is characteristic of phytoliths\(^5^1,5^7\). Coniferous phytoliths from the southern slopes of the Himalaya appear only in the planting zone of needle-leaved plants, correspondent to the distribution of coniferous forests; phytoliths can thus be used as discriminants of the timberline and the altitude of montane vegetation zones. When researching montane palaeoenvironments, both the frequency of

![Figure 8. Boxplots of phytolith percentages.](image-url)
occurrence and the variability in the content of coniferous phytoliths can provide a good basis for deducing the historical position of the timberline, as well as palaeoaltitude.

Furthermore, crop phytoliths can reflect different types of farming activity\textsuperscript{43,45}, and indeed different crops grow at different altitudes. The cross-shaped and cuneiform bulliform cell-rice phytoliths produced in the surface soils of the Nepal Himalaya indicate the planting of maize and rice, in accordance with the area's relatively low altitude. This also implies that human disturbance has changed the composition of the local vegetation. The existence of cultivated phytolith morphotypes in sediments and/or archaeological remains would certainly indicate an historical development of this region by humans\textsuperscript{10}.

Based upon our qualitative analysis, we conducted a quantitative analysis, with good results. The mathematical transfer function can be ideal for providing a basis for palaeoenvironmental research, especially when substantial quantities of modern data are applied. In this study, although we did not have a huge number of samples, our Ic values expressed the functional relation between phytolith assemblage variety and altitude very well, demonstrating that Ic is a perfect index for representing gramineous composition\textsuperscript{17} in mountainous areas like the Himalaya. However, we adjusted its formulaic expression. In previous research, the Stipa-bilobate short cell phytolith was added to neither the numerator nor the denominator in the index's mathematical expression. However, Stipa belongs to Pooideae, and can survive in extremely cold and arid environments. Stipa grasses are widely distributed at high altitudes on the QTP, and are a vital component of the grass cover in our study area\textsuperscript{17}. When Stipa-bilobate short cells are taken into account, the curve estimation of $R^2 = 0.8$. However, $R^2 = 0.7$ without the inclusion of Stipa-bilobate short cells. Certainly, Stipa-bilobate short cells, as an individual phytolith type, should be distinguished from the sum of phytoliths during identification and this should be taken into account when using the Ic index. However, when samples are collected from soil surfaces or Cenozoic stratigraphic sections in cold and arid areas like the QTP, the presence of Stipa-bilobate short cells must be fully taken into consideration.

Ic can be used to reconstruct the Quaternary palaeoelevation of the QTP during periods of relatively stable climate. We noted that in the East African tropical mountains, Ic values are ~40, ~70 and ~98 at 2000 m a.s.l., 3000 m a.s.l. and 4000 m a.s.l., respectively\textsuperscript{17}. The corresponding values are ~50, ~70 and ~90 in our paper. This may imply that Ic has some universal applicability in low latitude areas, as both of the abovementioned areas are low latitude regions.

The humidity-aridity index Iph was also tested, but proved not obviously applicable to altitude. First, unlike temperature, precipitation in the Nepal Himalaya is affected by two major atmospheric circulation systems. The interaction of the complex topography with the monsoonal and westerly weather systems results in variations in the spatial distribution of rainfall\textsuperscript{50}, suggesting that the annual precipitation pattern is not dominated by altitude. Second, Iph was defined as the ratio of Chloridoideae (saddle) versus Chloridoideae and Panicoideae (saddle, cross-shaped and bilobate short cell)\textsuperscript{17}. The square saddle phytolith is typical type of Chloridoideae. Some Chloridoideae species, which have adapted to drought conditions, are prone to produce square saddle phytoliths. Other Chloridoideae species in humid environments can produce another short cell phytolith, \textit{i.e.} bilobate. Iph has been shown to be applicable to the Great Plains of North America as well as to tropical savannah\textsuperscript{17,58}, but both these environments have dry seasons which last several months. However, the northern Himalaya enjoys rainier weather. Square saddle phytoliths may therefore not indicate the presence of Chloridoideae in the area. Moreover, the presence of Bambusoideae may also render Iph inapplicable. Bambusoideae produce great quantities of oblong concave saddle phytoliths, which account for a certain proportion of all saddle phytoliths. Using Iph as a index, saddle phytoliths should present as Chloridoideae, but the saddle phytoliths in the northern Himalaya are derived from Chloridoideae, Bambusoideae and Arundinoideae. Bremond et al. (2008) also demonstrated that Iph was not a relevant proxy for Chloridoideae versus Chloridoideae and Panicoideae\textsuperscript{17}.

Less research has been conducted into woody phytoliths than gramineous phytoliths, most probably because of the great variety in the morphology of woody phytoliths and the consequent difficulties discriminating between them. If a more detailed identification were possible, the environmental marker function would become more precise. For example, palmaceae phytoliths from the montane subtropical evergreen belt provide good indices for discriminating palm tree planting. There is a fundamental need for systematic and detailed research into woody phytoliths. Such palaeoenvironmental research would provide a basic reference tool for identifying vegetation belts, phytocoenoses, edificators, and even typical species.

**Methods**

In this study, 47 topsoil samples were collected at altitudes between 100 m a.s.l. and 5200 m a.s.l. from the Sino-Nepal Himalaya (Fig. 1). No sample was collected from the perpetual snow zone, because the vegetation becomes extremely sparse above 5200 m a.s.l. After clearing away any loose debris, each sample was extracted from the top 0–2 cm of surface soil and put into a valve bag. A GPS receiver was applied to keep a record of the longitude, latitude and altitude of each sampling site. Samples were dried out in the laboratory before long-term storage in order to prevent clumping.

Phytoliths were extracted from each sample by conventional heavy liquid flotation based on the method of Piperno and Pearsall\textsuperscript{6,59}, but were not passed through a sieve, as some phytoliths in this paper (e.g. rondel and square saddle phytoliths) are very small and could be filtered off during the process.
All samples were treated in the Key Laboratory of Cenozoic Geology and Environment, Institute of Geology and Geophysics, Chinese Academy of Sciences. A subsample of ~1–3 g was taken from each sample, according to its particular composition. Each subsample was sequentially processed as follows: (1) organics were removed with 30% H2O2; (2) a tablet of Lycopodium spores (27637 spores per tablet) was added to determine phytolith concentration (the method of Piperno and Pearsall does not include this step); (3) carbonates were dissolved with 10% HCl; (4) flotation of phytoliths was accomplished using a ZnBr2 solution (density 2.35 g/cm3); and (5) after cleaning, each subsample was dipped in resinine to facilitate slicing. Morphotype counts were counted under a microscope; 41 samples were found to include >300 phytoliths.

Canoco5 software was used to perform RDA.60

We used SPSS 17.0 statistics software to build scatter diagrams of Ic and Iph along altitude, and make curve estimations. We also applied SPSS 17.0 to DA and to our boxplots.

References
1. Ramstein, G., Fluteau, F., Besse, J. & Joussaume, S. Effect of orogeny, plate motion and land-sea distribution on Eurasian climate change over the past 30 million years. Nature 386, 788–795 (1997).
2. Fauquette, S. et al. Reconstruction of the Northern and Central Apennines (Italy) palaeoaltitudes during the late Neogene from pollen data. Rev Palaeobot Palynol, doi: http://dx.doi.org/10.1016/j.revpalbo.2014.06.001 (2014).
3. Zheng, Z. H. et al. A study on surface pollen assemblage and relationship with vegetation from some vegetation types in Central North China. Geography and Geo-Information Science 24, 92–97 (2008).
4. Xu, Q. H., Li, Y. C., Yang, X. L. & Zheng, Z. H. Quantitative relationship between pollen and vegetation in northern China. Sci China Ser. D Earth Sci 50, 582–599 (2007).
5. Lu, H. Y. et al. Spatial pattern of Abies and Picea pollen distribution along the elevation gradient in the Qinghai–Tibetan Plateau and Xinjiang, China. Boreas 37, 254–262, doi: 10.1111/j.1503-8837.2007.00191.x (2008).
6. Piperno, D. R. Phytoliths: A Comprehensive Guide for Archaeologists and Palaeoecologists. (AltaMira Press, 2006).
7. Lu, H. Y., Wu, N. Q., Liu, K. B., Jiang, H. & Liu, T. S. Phytoliths as quantitative indicators for the reconstruction of past environmental conditions in China II: Palaeo-environmental reconstruction in the Loess Plateau. Quat Sci Rev 26, 759–772 (2007).
8. Calegari, M. R., Madella, M., Vidal-Torrado, P., Pessenda, L. C. R. & Marques, F. A. Combining phytoliths and 813C matter in Holocene palaeoenvironmental studies of tropical soils: An example of an Oxisol in Brazil. Quat Int 287, 47–55, doi: http://dx.doi.org/10.1016/j.quaint.2011.11.012 (2013).
9. Alexandre, A., Meunier, J. D., Lézine, A. M., Vincens, A. & Schwartz, D. Phytoliths: indicators of grassland dynamics during the late Holocene in intertropical Africa. Palaeogeogr Palaeoclimatol Palaeoecol 136, 213–229, doi: http://dx.doi.org/10.1016/S0031-0182(97)00039-8 (1997).
10. Garnier, A., Neumann, K., Eichhorn, B. & Lespez, I. Phytolith taphonomy in the middle- to late-Holocene fluvial sediments of Onejogou (Mali, West Africa). The Holocene 23, 416–431, doi: 10.1177/0959683612463102 (2012).
11. Petö, Á. Studying modern soil profiles of different landscape zones in Hungary: An attempt to establish a soil-phytolith identification key. Quat Int 287, 149–161, doi: http://dx.doi.org/10.1016/j.quaint.2012.02.049 (2013).
12. Lu, H. Y. & Liu, K. B. Phytolith assemblages as indicators of coastal environmental changes and hurricane overwash deposition. The Holocene 15, 965–972 (2005).
13. Carnella, A. L., Theurillat, J. P. & Madella, M. Phytolith types and type-frequencies in subalpine-alpine plant species of the European Alps. Rev Palaeobot Palynol 129, 39–65 (2004).
14. Mercader, J., Bennett, T., Esselmont, C., Simpson, S. & Walde, D. Phytoliths in woody plants from the Miombo woodlands of Mozambique. Ann Bot 104, 91–113 (2009).
15. Cordova, C. C. E., Poaceae and Restionaceae phytoliths as potential proxies for reconstructing winter rainfall in South Africa. Quat Int 287, doi: http://dx.doi.org/10.1016/j.quaint.2012.04.002 (2013).
16. Blinnikov, M. S., Bagent, C. M. & Reyerson, P. E. Phytolith assemblages and opal concentrations from modern soils differentiate temperate grasslands of controlled composition on experimental plots at Cedar Creek, Minnesota. Quat Int 287, 101–113, doi: http://dx.doi.org/10.1016/j.quaint.2011.12.023 (2013).
17. Bremond, L. et al. Phytolith indices as proxies of grass subfamilies on East African tropical mountains. Global and Planetary Change 61, 209–224, doi: http://dx.doi.org/10.1016/j.gloplacha.2007.08.016 (2008).
18. Liu, X. D. & Shen, B. D. Climatic warming in the Tibetan Plateau during recent decades. Int. J. Climatol. 20, 1729–1742, doi: 10.1002/1097-0088(20001130)20:14<1729::AID-JOC556>3.0.CO;2-Y (2000).
19. Pan, G. T. et al. Tectonic evolution of the Qinghai-Tibet Plateau. Journal of Asian Earth Sciences 53, 3–14, doi: http://dx.doi.org/10.1016/j.jseaes.2011.12.018 (2012).
20. Peterse, F. et al. Assessment of soil n-alkane δD and branched tetraterpane membrane lipid distributions as tools for paleoelevation reconstruction. Biogeosciences 6, 8609–8631 (2009).
21. Tibet Scientific Expedition Team, Chinese Academy of Sciences. Scientific Expedition Report on Mount Qomolangma Region: Physical Geography 1966–1968. 1–29 (Science Press, 1975).
22. Stanton, J. D. A. Forests of Nepal. (John Murray Publisher, 1972).
23. Dobremez, I. J. Le Nepal: Écologie et Biogéographie. (Centre National de la Recherche Scientifique, 1976).
24. Singh, J. S. & Singh, S. P. Forest vegetation of the Himalaya. Bot. Rev. 53, 80–192 (1987).
25. Baniya, C. B., Solhøy, T. & Vetaas, O. R. Temporal changes in species diversity in a trans-Himalayan landscape, Nepal. Herbaceous Plant Ecology 201, 383–399 (2008).
26. Scientific Expedition to the Qinghai-Xizang Plateau. vegetation of Xizang (Tibet). (science press, 1988).
27. Food and Agriculture Organization of the United Nations. FRA 2000, forest resources of Nepal-country report. (Rome, 1999).
28. Liu, D. S. World Geography Overview. (oriental publishing center, 1997).
29. Bhattarai, K. R., Vetaas, O. R. & Grytnes, J. A. Fern species richness along a central Himalayan elevational gradient, Nepal. J Biogeogr 31, 389–400 (2004).
30. Lu, H. Y. et al. Phytoliths as quantitative indicators for the reconstruction of past environmental conditions in China I: phytolith-based transfer functions. Quat Sci Rev 25, 945–959 (2006).
31. Wang, Y. J. & Lu, H. Y. The Study of Phytolith and its Application. (China Ocean Press, 1993).
32. Kondo, R., Childs, C. & Atkinson, I. Opal Phytoliths of New Zealand. (Manaaki Whenua Press, 1994).
33. Thwaites, P. C., Suess, E. & Smith, R. M. Division S—soil genesis, morphology, and classification (morphological classification of grass phytoliths). Soil Science Society of America Proceedings 33, 109–115 (1969).
34. Madella, M., Alexandre, A. & Ball, T. International Code for Phytolith Nomenclature 1.0. Ann Bot 96, 253–260 (2005).

www.nature.com/scientificreports/
Competing financial interests: The authors declare no competing financial interests.

How to cite this article: An, X. et al. Surface soil phytoliths as vegetation and altitude indicators: a study from the southern Himalaya. Sci. Rep. 5, 15523; doi: 10.1038/srep15523 (2015).

This work is licensed under a Creative Commons Attribution 4.0 International License. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in the credit line; if the material is not included under the Creative Commons license, users will need to obtain permission from the license holder to reproduce the material. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/