Late Pliocene vegetation turnover on the NE Tibetan Plateau (Central Asia) triggered by early Northern Hemisphere glaciation

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Abstract

To reconstruct the timing and underlying forcing of major shifts in the composition of terrestrial ecosystems in arid Central Asia during the late Cenozoic (past ~7 Ma), we carry out palynological analysis of lake sediments from the Qaidam Basin (NE
Tibetan Plateau, China). Our results show that the steppe/semi-desert biomes dominating the Qaidam Basin experienced marked turnovers at ~3.6 and 3.3 Ma. Most notably, the younger of these turnover events is characterized by a two- to three-fold expansion of *Artemisia* at the expense of other steppe/semi-desert taxa. This turnover event led to the replacement of the Ephedraceae/Chenopodiaceae-dominated and *Nitraria*-rich steppe/semi-deserts that were dominant in the Qaidam Basin during the Paleogene and abundant during the Miocene by *Artemisia*/Chenopodiaceae-dominated steppe/semi-deserts as they exist until today. The vegetation turnover events are synchronous with shifts towards drier conditions in Central Asia as documented in climate records from the Chinese Loess Plateau and the Central North Pacific Ocean. On a global scale, they can be correlated to early glaciation events in the Northern Hemisphere during the Pliocene. Integration of our palynological data from the Qaidam Basin with Northern Hemisphere climate-proxy and regional-scale tectonic information suggests that the uplift of the Tibetan Plateau posed ecological pressure on Central Asian plant communities, which made them susceptible to the effects of early Northern Hemisphere glaciations during the late Pliocene. Although these glaciations were relatively small in comparison to their Pleistocene counterparts, the transition towards drier/colder conditions pushed previously existing plant communities beyond their tolerance limits, thereby causing a fundamental reorganization of arid ecosystems. The *Artemisia* dominance since ~3.3 Ma resulting from this reorganization marks a point in time after which the *Artemisia*/Chenopodiaceae pollen ratio can serve as a reliable indicator for moisture availability in Central Asia.

**Keywords**
Aridification; Neogene; Northern Hemisphere glaciation; Qaidam Basin; Tibetan Plateau; vegetation dynamics
1. Introduction

As a result of anthropogenic climate change, considerable shifts in hydrological conditions are projected for Central Asia for the near future (IPCC, 2014). It is therefore crucial to obtain a better understanding of the climatic forcing on plant-community dynamics in the affected regions in order to extend the lead time for mitigation and adaptation. In this context, the study of past intervals in Earth history that were warmer than today and had atmospheric carbon dioxide concentrations \( (p\text{CO}_2) \) similar to those expected for the near future (Martínez-Botí et al., 2015; Burke et al., 2018) provides a promising avenue for predicting future developments. From late Miocene (~11.6–5.3 Ma) to Pliocene (~5.3–2.6 Ma) times, global temperatures were several degrees warmer and atmospheric \( p\text{CO}_2 \) was slightly higher than today (Burke et al., 2018; Holbourn et al., 2018). Thus, the late Neogene represents the most recent geological interval that can serve as an analogue for a future anthropogenic greenhouse world.

Central Asia was among the most arid regions on Earth during most of the Neogene and forms a key region for deciphering the evolution of ecosystems in arid environments (Guo et al., 2002; An et al., 2005; Dupont-Nivet et al., 2007). Changes in the carbon-isotope signatures of soils from the Chinese Loess Plateau during the past 7 Ma have documented a major shift in plant communities at ~3.6 Ma, associated with the expansion of drought-tolerant C\(_4\) plants at the expense of plants exploiting the C\(_3\) photosynthetic pathway (An et al., 2005). Palynological data are in agreement with these findings; they document that *Artemisia* emerged as the dominant taxon in arid regions across China during the late Pliocene (Wang, 2004; Miao et al., 2011b; Cai et al., 2012). This expansion occurred at the expense of Chenopodiaceae and other drought-tolerant taxa (e.g., Ephedraceae, *Nitraria*) that prevailed in Central Asia since the Paleogene (Hoorn et al., 2012; Han et al., 2016;
Miao et al., 2016). This shift in plant communities has been attributed predominantly
to the uplift of the Tibetan Plateau and its impact on regional monsoonal dynamics
(An et al., 2001; Molnar et al., 2010), but may also represent a response to global
late Neogene cooling (Zhang et al., 2001; Guo et al., 2002; Lu et al., 2010; Licht et
al., 2016). Because of the lack of continuous, highly resolved and well-dated
vegetation records, the signature, timing and duration of these vegetation transitions,
and, by extension, the underlying trigger mechanisms have remained poorly
constrained.

We here present a new palynological record from drillcores that were retrieved from
the Qaidam Basin (north-eastern Tibetan Plateau) and spans from ~7.3 until 0.1 Ma
ago (Figure 1). To warrant detailed analysis of previously recognized phases of
pronounced vegetation turnover in Central Asia, we have studied the mid-Pliocene to
late Pleistocene interval in high (millennial- to suborbital-scale) temporal resolution
(~3–4 ka). The remainder of the drillcore record was studied in orbital- to tectonic-
scale resolution (~10 to >30 ka) in order to reconstruct the long-term vegetation
development since the late Miocene. To better constrain the impacts of both the
Tibetan Plateau uplift and global climate change on the evolution of plant
communities in the Qaidam Basin, we compare our palynological data with
supraregional terrestrial (e.g., Chinese Loess Plateau) and marine (e.g., North
Pacific) climate-proxy records.

2. Regional setting

Situated close to the north-eastern margin of the Tibetan Plateau, the Qaidam Basin
ranges among the highest, most evaporative, and largest inland basins of Central
Asia (Fang et al., 2007; Yang et al., 2011). It lies at an average altitude of ~2800 m
above sea level and is surrounded by up to ~5000 m high mountain ranges (Figure
1). Basin formation started in the early Eocene as a result of orogenic shortening due
to the India-Eurasia continental collision (Wang et al., 2008; Yin, 2010). Further expansion and deepening prevailed until the mid-Miocene (Fang et al., 2007), but from the late Miocene onwards tectonic compression led to shrinking of the basin (Wang et al., 2012; Yuan et al., 2013). Today, the basin hosts one of world’s thickest fluvio-lacustrine Cenozoic sedimentary records, with a maximum thickness of ~15 km (Fang et al., 2007).

As most other regions of Central Asia, the Qaidam Basin has undergone gradual drying through the Cenozoic, with the Miocene being wetter than the Pliocene and Pleistocene, respectively (Wang et al., 1999; Fang et al., 2007). This drying has been attributed to a combination of tectonic activity in Central Asia, including the northward drift and uplift of the basin connected to Tibetan Plateau uplift, and global Cenozoic cooling (Wang et al., 1999; Fang et al., 2007). To date, the basin is characterized by a W–E precipitation gradient; for the western part of the basin where the study material comes from, mean annual precipitation is 15.8 mm and mean annual potential evaporation is 2967 mm (Lenghu meteorological station for the period 1957–2000; Chen et al., 2013).

Because of the prevailing (hyper)arid conditions, the modern vegetation in the Qaidam Basin represents a transition between the Mongolian desert and the alpine steppe of the Tibetan Plateau (Walter and Box, 1983). Specifically, four major biomes (i.e., alpine meadows, steppes, steppe/deserts, and deserts) occur in the Qaidam Basin and the surrounding mountains (Yu et al., 2001; Zhao and Herzschuh, 2009; Zhang et al., 2010); they consist predominantly of herbs and shrubs of the genera *Artemisia*, *Calligonum*, *Ephedra*, *Haloxylon*, *Kalidium*, *Nitraria*, *Reaumuria*, *Salsola*, *Sympegma*, and *Tamarix* (Zhao and Herzschuh, 2009; Zhao et al., 2007, 2010; Wei et al., 2011). Forests are confined to the surrounding mountains and mainly consist of *Betula*, *Pinus*, and *Picea* (Zhao et al., 2009; Wei et al., 2011).
3. Material and methods

Two long drillcores were recovered in 2008 and 2011 from paleolake sediments in the western Qaidam Basin (Figure 1). Core SG-1 (38°24′35″N, 92°30′33″E) was drilled in the depocenter of the Chahansilatu sub-basin to a depth of 938 m with a recovery of ~95% (Zhang et al., 2012a). Core SG-1b (38°21′9.46″N, 92°16′24.72″E) was recovered from the Jianshan anticline, ~20 km SW of core SG-1, and reached a depth of 723 m; recovery is ~93% with the exception of the topmost 60 m, where core loss was ~20% (Zhang et al., 2014). Lithologically, both cores consist predominantly of lacustrine clay-, silt- and calcareous mudstones with intercalated marl, halite and gypsum layers (Figure 2; Wang et al., 2012; Zhang et al., 2012b; Lu et al., 2015).

Figure 1: Locations of the SG-1 and SG-1b cores in Central Asia and within the Qaidam Basin (insert map).
3.1 Chronology

Age control for core SG-1 is based on the integration of magnetostratigraphy, optical stimulated luminescence (OSL) and U-Th dating, and cyclostratigraphy. The observed magnetic polarity zones were correlated with Chrons 1n-2An of the geomagnetic polarity time scale (Zhang et al., 2012a). The age model was further refined by cyclostratigraphic analysis of a high-resolution magnetic susceptibility dataset yielding a basal age of 2.69 Ma (Herb et al., 2013; Herb et al., 2015a). The core top has an OSL- and U/Th-based minimum age of ~77.8 ± 4.0 ka (Han et al., 2014). The chronology of core SG-1b is based on magnetostratigraphy yielding a late Miocene to early Pleistocene age (Chrons 2n-3Br; ~7.3 to 1.6 Ma; Zhang et al., 2014). Based on their age models, the SG-1 and SG-1b cores overlap between ~2.6 and 1.6 Ma.

3.2 Palynology

In total, 847 pollen samples were analysed from both cores. Specifically, 505 samples were newly processed and counted for this study; this dataset was augmented by another 342 samples previously published in Herb et al. (2015b) and Koutsodendris et al. (2018). All samples were processed following standard palynological techniques including freeze-drying, HCl (30 %) and HF (38 %) digestion, treatment with KOH, heavy-liquid density separation, and sieving through a 10 µm mesh. At least 250 pollen grains were counted for 95 % of the samples (mean: 333; range: 23–801 grains), excluding pollen from aquatic plants and spores. For the calculation of pollen percentages, pollen from aquatic plants and fern spores were excluded because they bear a strongly local vegetation signal. All palynological plots were generated using the C2 software.
3.3 Statistical evaluation

Principal component analysis (PCA) was carried out using the PAST software to examine the relationships among the dominant pollen taxa in the core material. PCA was applied on a set of selected steppe/desert taxa that are continuously present in the record; arboreal and aquatic taxa were included in the analysis as groups.

4. Results

4.1 Palynological analysis of cores SG-1 and SG-1b

All palynological samples from cores SG-1 and SG-1b are dominated by herbaceous and shrubby taxa, particularly Chenopodiaceae (mean: 30%; maximum: 51%), Artemisia (28% and 76%), Poaceae (12% and 40%), Ephedraceae (5% and 39%), Cyperaceae (5% and 25%), and Asteraceae (4% and 16%) (Figure 2). Several other herbs and shrubs with average abundances <1% are present continuously across the study interval; some of these taxa transiently reach higher abundances, such as Tamaricaceae (maximum: 13%), Elaeagnaceae (12%), Ranunculaceae (11%), Calligonum (7%), Caryophyllaceae (7%), and Nitraria (6%).

Arboreal pollen (AP) represent only a small percentage of the pollen sum, with conifers and broad-leaved deciduous trees on average making up for 5 and 2% of the assemblages, respectively (Figure 2). The main conifer taxa include Pinus (maximum: 54%), Cupressaceae (13%), Picea (13%), and Abies (3%); Cedrus, Larix, Podocarpus, and Tsuga occur in low amounts. The most common deciduous broad-leaved trees comprise Betula (maximum: 9%), Ulmus (7%), Alnus (5%), Quercus (4%), and Juglans (3%); low amounts of Carpinus, Corylus, Fraxinus, Salix, and Tilia were also registered. Finally, aquatic plants including Typha latifolia, T. angustifolia, and Potamogeton are continuously present, with average abundances of 3% (maximum: 22%) (Figure 2).
Figure 2: Pollen percentages of selected pollen taxa identified in the SG-1 and SG-1b cores spanning the past 7.3 Ma. Magnetostratigraphy and cores lithologies are adapted from Wang et al. (2012), Zhang et al. (2012b, 2014) and Lu et al. (2015).
While the spectrum of pollen-producing plants has remained remarkably stable in the Qaidam Basin since the late Miocene, closer inspection reveals substantial changes in the abundances of the dominant taxa. Whereas Chenopodiaceae percentages remained rather stable throughout the record (~30 %), the percentages of other dominant taxa such as *Artemisia*, Ephedraceae and Poaceae exhibit considerable variability (Figure 2). Specifically, *Artemisia* abundances increased from an average of 12 % (~7.3–3.3 Ma) to 30 % (~3.3–0.3 Ma), and Ephedraceae abundances decreased from an average of 18 % (~7.3–3.7 Ma) to 4 % (~3.7–0.3 Ma). In addition, Poaceae abundances show a quasi-cyclical behaviour with high values of 19–20 % on average during ~3.5–3.3 and ~3.1–2.3 Ma, and 14 % during ~1.2–0.7 Ma; instead, values <5 % occurred during ~3.3–3.1, 1.6–1.3, and 0.7–0.6 Ma (Figure 2). Among the arboreal taxa, the percentages of *Pinus* increased from ~3.6 Ma onwards at the expense of all other tree-pollen taxa (including deciduous trees and conifers other than *Pinus*) (Figure 2).

4.2 Principal component analysis

The PCA yields three major components that account for 80.2 % of the total variance of our pollen dataset (Table 1). The first component explains 38.4 % of the total variance and is bipolar; it is primarily driven by *Artemisia*, which is marked by highly negative loadings, and Poaceae, Chenopodiaceae and AP, which are characterized by positive loadings (Table 2; Figure 3). The second component is also bipolar and explains 27.4 % of the total variance. Whereas its negative pole is primarily driven by Chenopodiaceae and to a lesser extent by Ephedraceae, its positive pole is driven by Poaceae and AP. The third component, which explains 14.4 % of the total variance, is also bipolar (Tables 1 and 2). It is marked by highly positive loadings of AP and highly negative loadings of Poaceae.
Table 1: Total variance explained by the first three PCA components of the pollen dataset from the SG-1 and SG-1b cores.

<table>
<thead>
<tr>
<th>PC</th>
<th>Eigenvalue</th>
<th>% variance</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>115.245</td>
<td>38.4</td>
</tr>
<tr>
<td>2</td>
<td>82.2967</td>
<td>27.4</td>
</tr>
<tr>
<td>3</td>
<td>43.0543</td>
<td>14.4</td>
</tr>
</tbody>
</table>

Table 2: PCA loadings of the pollen dataset from the SG-1 and SG-1b cores.

<table>
<thead>
<tr>
<th></th>
<th>PC 1</th>
<th>PC 2</th>
<th>PC 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Arboreal pollen (AP)</td>
<td>0.168</td>
<td>0.364</td>
<td>0.675</td>
</tr>
<tr>
<td>Elaeagnaceae</td>
<td>0.024</td>
<td>0.035</td>
<td>-0.019</td>
</tr>
<tr>
<td>Tamaricaceae</td>
<td>0.022</td>
<td>-0.045</td>
<td>-0.006</td>
</tr>
<tr>
<td>Ephedraceae</td>
<td>0.049</td>
<td>-0.132</td>
<td>0.143</td>
</tr>
<tr>
<td>Artemisia</td>
<td>-0.901</td>
<td>0.039</td>
<td>-0.125</td>
</tr>
<tr>
<td>Chenopodiaceae</td>
<td>0.210</td>
<td>-0.800</td>
<td>-0.078</td>
</tr>
<tr>
<td>Apiaceae</td>
<td>0.002</td>
<td>-0.005</td>
<td>-0.004</td>
</tr>
<tr>
<td>Brassicaceae</td>
<td>0.006</td>
<td>-0.008</td>
<td>-0.012</td>
</tr>
<tr>
<td>Calligonum</td>
<td>-0.011</td>
<td>-0.010</td>
<td>-0.034</td>
</tr>
<tr>
<td>Caryophyllaceae</td>
<td>0.011</td>
<td>-0.012</td>
<td>0.022</td>
</tr>
<tr>
<td>Asteraceae</td>
<td>0.014</td>
<td>0.027</td>
<td>0.134</td>
</tr>
<tr>
<td>Cyperaceae</td>
<td>0.078</td>
<td>0.160</td>
<td>0.033</td>
</tr>
<tr>
<td>Poaceae</td>
<td>0.321</td>
<td>0.423</td>
<td>-0.693</td>
</tr>
<tr>
<td>Rosaceae</td>
<td>0.009</td>
<td>0.003</td>
<td>-0.005</td>
</tr>
<tr>
<td>Thalictrum</td>
<td>-0.001</td>
<td>0.004</td>
<td>0.011</td>
</tr>
<tr>
<td>Aquatics</td>
<td>0.058</td>
<td>0.003</td>
<td>0.028</td>
</tr>
</tbody>
</table>
5. Discussion

5.1 Timing and causes of major vegetation turnover in the Qaidam Basin

Our palynological data from cores SG-1 and SG-1b document that a steppe/semi-desert biome dominated the vegetation in the Qaidam Basin from the late Miocene to the late Pleistocene (Figure 2). The PCA results show that the distribution of the dominant taxa is controlled by different factors. The first PCA component primarily differentiates *Artemisia* from Chenopodiaceae and Ephedraceae, whereas the second component differentiates the latter taxa from Poaceae and AP (Figure 3). Moreover, the third component further differentiates Poaceae from AP. Considering the modern distribution of these taxa in the Qaidam Basin (Zhao et al., 2007, 2009; Zhao and Herzschuh, 2009), we explain the first two components to represent the differentiation between *Artemisia*- and Chenopodiaceae/Ephedraceae-dominated steppe/semi-deserts. By extension, the third component differentiates between local and distal pollen sources, considering that grasses predominantly grow in proximity to the lake shores in the Qaidam Basin whereas forests develop only in the surrounding mountains (Cour et al., 1999; Zhao et al., 2007, 2009; Wei et al., 2011).

A closer look reveals a series of changes in the composition of the steppe/semi-desert vegetation during the late Pliocene. A first vegetation turnover at ~3.6 Ma is characterized by an increase in *Artemisia*, Poaceae and Cyperaceae abundances, and a coeval decrease in Ephedraceae and Chenopodiaceae abundances (Figure 2). A second, fundamental turnover in the composition of the steppe/desert vegetation in the Qaidam Basin occurred at ~3.3 Ma. It is marked by a two- to three-fold increase in *Artemisia* percentages at the expense of other steppe/desert taxa including Ephedraceae, Asteraceae and Tamaricaceae (Figure 2). This interval is also marked by a shift in the composition of the forests that thrived on the mountain slopes surrounding the Qaidam Basin; the abundance of the cold- and drought-tolerant *Pinus* increased at the expense of warmth- and high-moisture-demanding
deciduous trees (Figure 2). These two abrupt turnover events refine the exact timing of the *Artemisia* expansion in the Qaidam Basin. Based on the neighbouring pollen records from cores SG3 and KC-1 from the western Qaidam Basin that discontinuously span the past 18 Ma, this expansion can be placed between 5 and 3 Ma (Miao et al., 2011a; Cai et al., 2012).

**Figure 3:** Principal Component Analysis (PCA) of the palynological dataset from the SG-1 and SG-1b cores. The first two components explain 65.8 % of the total variance.

The vegetation-turnover events coincide with sedimentological and geochemical changes in core SG-1b and suggest two major climate shifts at ~3.6 and ~3.3 Ma, respectively. Increases in the medium-coarse silt and sand fractions in the respective core intervals indicate lake-level drops, and by extension, a prevalence of drier conditions (Lu et al., 2015). This finding is further supported by the coeval deposition of evaporites in the SG-1b core (Fang et al., 2016). On a supraregional scale, synchronous transitions towards drier conditions in Central Asia are also documented for the Chinese Loess Plateau and the Pacific Ocean. On the Chinese
Loess Plateau, an increase in the size of quartz grains in the loess deposits at ~3.6 Ma indicates enhanced wind strength resulting from a stronger influence of the cold and dry air masses of the winter monsoon (An et al., 2001; Lu et al., 2010). The coeval increase of C₄ plants on the Chinese Loess Plateau also suggests a shift in plant communities in response to a stronger prevalence of dry conditions (An et al., 2005). The onset of drier conditions at ~3.6 Ma is further evidenced by enhanced dust delivery from Central Asia into the Central Pacific Ocean as documented at Ocean Drilling Program (ODP) Sites 885 and 886 (Rea et al., 1998; Figure 4). This pattern is coeval with the Artemisia percentage increase in the SG-1 and SG-1b cores from the Qaidam Basin (Figure 4).

Because Artemisia requires a higher moisture availability than Chenopodiaceae and Ephedraceae during the growing season (El’Moslimany, 1990; Cour et al., 1999), its expansion at the expense of other drought-tolerant steppe/semi-desert taxa against a background of an increasingly dry climate during the late Pliocene (Rea et al., 1998; An et al., 2001; Lu et al., 2010; Sun et al., 2010) cannot be explained by a prevalence of wetter conditions. Instead, it could be plausibly explained through a seasonal shift in moisture availability. Quantitative analysis of the annual distribution of modern pollen rain along the western border of the Qaidam Basin has shown that maximum abundances of Artemisia pollen occur in late summer and autumn; in contrast, maxima in Chenopodiaceae, Ephedraceae, Poaceae, and Tamaricaceae pollen abundances occur in early summer (Cour et al., 1999). This pattern suggests that the late Pliocene expansion of Artemisia could result from a seasonal shift of maximum precipitation from summer to autumn, with the shift in moisture availability giving the taxon a competitive advantage over Chenopodiaceae and Ephedraceae. Support for such a scenario comes from the decoupling of the winter and summer monsoon systems as evidenced in sedimentological data from the Chinese Loess Plateau (Sun et al., 2010). Whereas the Miocene and early Pliocene climates in
Central Asia were characterized by enhanced summer precipitation, late Pliocene and Pleistocene climates were marked by relatively low summer precipitation (Sun et al., 2010). These findings further support our hypothesis that a seasonal shift in precipitation distribution caused the vegetation turnover in the Qaidam Basin and, by extension, in Central Asia during the late Pliocene.

**Figure 4:** Climate records for the past 7.3 Ma from the SG-1 and SG-1b cores in the Qaidam Basin plotted against regional and global climate records. (a) LR04 global marine isotope stack (Lisiecki and Raymo, 2005); (b) East Asian summer monsoon index from the Lingtai section, Chinese Loess Plateau (Sun et al., 2010); (c) Dust accumulation rates at ODP Sites 885 and 886 in the North Pacific Ocean (Rea et al., 1998); (d) *Artemisia* pollen percentages, (e) sum of *Artemisia* and Chenopodiaceae (A+C) pollen percentages, and (f) *Artemisia*/Chenopodiaceae (A/C) ratio, in the SG-1 and SG-1b cores (this study). The A/C ratio is plotted against the 200 mm boundary for annual precipitation on the Tibetan Plateau as defined by an A/C value of 1 (Zhao et al., 2012). Turnover events are indicated with dashed lines.
The monsoonal variability in Central Asia since the Miocene has been predominantly attributed to the tectonic uplift of the Tibetan Plateau (An et al., 2001), which poses a physical obstacle to atmospheric flow and alters the temperature and pressure field immediately above it due to surface heating (see Molnar et al., 2010, for a review). Previous work has repeatedly invoked the increase in aridity (Rea et al., 1998; An et al., 2001; Sun et al., 2010) and the expansion of drought-tolerant plants in Central Asia (An et al., 2005) to Tibetan Plateau uplift. However, longer-term and gradual tectonic forcing appears unlikely to have caused the rather abrupt shifts at ~3.6 and ~3.3 Ma as evidenced in the palynological data from the Qaidam Basin. This is even more so considering that the Qaidam Basin and surrounding mountain ranges had reached elevations close to today already in the mid-Miocene (Wang et al., 2008; Yuan et al., 2013). Hence, the abrupt shifts in steppe/desert composition as documented by the *Artemisia* expansions at ~3.6 and ~3.3 Ma must have been caused by a factor other than Tibetan Plateau uplift.

Conspicuously, both vegetation turnover events in the Qaidam Basin coincide with the first major Northern Hemisphere glaciations of the Pliocene (de Schepper et al., 2014). Specifically, the turnover at ~3.6 Ma occurs synchronously with the onset of gradual cooling in the Northern Hemisphere as documented in a surface-water temperature decline in the North Atlantic (Lawrence et al., 2009; Naafs et al., 2010) and an ice-sheet expansion in the Arctic (de Schepper et al., 2014). Although the glaciation at ~3.6 Ma is not linked to a specific marine isotope stage (MIS), it is highly likely that it occurred during the Gi2 and/or Gi4 glacial (de Schepper et al., 2014). The turnover in plant communities at ~3.3 Ma as documented by our palynological data coincides with MIS M2, which stands out as the first glacial of the Pliocene when local glaciers and ice caps in the Northern Hemisphere merged to modern-like ice sheets (de Schepper et al., 2014). A quasi-global temperature decrease during the M2 glaciation is documented for both the terrestrial and marine realms. Evidence
for such a cooling includes a surface-temperature drop (Naafs et al., 2010) and the
deposition of ice-rafted debris (Kleiven et al., 2002) in the North Atlantic, a global
sea-level drop of at least 10 m (Naish and Wilson, 2009), and an expansion of tundra
vegetation at the expense of forests in Arctic Siberia associated with a \( \sim 5 \) °C
decrease in the mean temperature of the warmest month (Brigham-Grette et al.,
2013). Collectively, the timing and short duration of the vegetation shifts in the
Qaidam Basin support the view that early Northern Hemisphere glaciations during
the Pliocene were responsible for the reorganization of vegetation communities in
Central Asia. The cooler conditions resulting from cryosphere expansion in the high
latitudes of the Northern Hemisphere (de Schepper et al., 2014) would have
strengthened the Siberian High pressure system, thereby leading to colder and more
arid conditions on the Tibetan Plateau (An et al., 2001; Sun et al., 2010).

Our data indicate unequivocally that the fundamental change in Central Asian
vegetation substantially pre-dates the intensification of Northern Hemisphere
glaciation during the Plio/Pleistocene transition at \( \sim 2.6 \) Ma (Naafs et al., 2012; de
Schepper et al., 2014). It appears that the early glaciations of the late Pliocene,
particularly the M2 glaciation, pushed previously existing plant communities beyond
their tolerance limits. Notably, these glaciations were able to do so although they
were markedly less pronounced than their Pleistocene counterparts (Lawrence et al.,
2009; Naafs et al., 2012). The subsequent Pleistocene glaciations have only
decreased the vegetation cover and led to changes in the proportion of the dominant
taxa percentages without triggering any large-scale vegetation turnover
(Koutsodendris et al., 2018). We attribute this apparent discrepancy to the ecological
pressure that the uplift of the Tibetan Plateau since the late Miocene has exerted on
these plant communities, notably via changes in monsoon dynamics associated with
uplift (An et al., 2001; Sun et al., 2010). This ecological pressure made them
susceptible to the climate perturbations connected to the first Northern Hemisphere
glaciations, even if those were of relatively small magnitude only. Our findings from
the Qaidam Basin support the view that terrestrial ecosystem changes in Central
Asia were primarily the result of global climate change rather than Tibetan Plateau
uplift (Zhang et al., 2001; Guo et al., 2002; Lu et al., 2010).

5.2 Implications for the Artemisia/Chenopodiaceae ratio as a moisture indicator

Because *Artemisia* requires more humidity during the growing season than
Chenopodiaceae, the *Artemisia*/Chenopodiaceae (A/C) ratio has long been used as
a qualitative index for moisture availability in arid regions during the past decades. In
this ratio, high (low) A/C values indicate wetter (drier) conditions (El-Moslimani, 1990;
vvan Campo and Gasse, 1993, Cour et al., 1999). This notion has been confirmed
repeatedly by the coupling of modern palynological and meteorological data across
the Tibetan Plateau that have yielded increasing A/C values from desert to steppe
environments and a positive correlation with annual precipitation (Herzschuh, 2007;
Zhao et al., 2012; Ma et al., 2017). By extension, the A/C ratio has been used to
reconstruct changes between steppe and desert biomes (Herzschuh et al., 2004; Luo
et al., 2009; Zhao and Herzschuh, 2009; Wei et al., 2011), and to infer semi-
quantitative information on annual precipitation (Zhao et al., 2012).

The suitability of the A/C ratio for the semi-quantitative reconstruction of moisture
availability has also led to its application to fossil pollen records from arid regions.
This holds particularly true for the Tibetan Plateau, where the A/C ratio has been
used in various studies spanning the Holocene (van Campo and Gasse, 1993; Zhang
et al., 2010; Zhao et al., 2007, 2010; Chen et al., 2013). According to Zhao et al.
(2012), the A/C ratio provides reliable semi-quantitative estimates for the moisture
variability in steppe/desert environments during the Holocene if (i) annual
precipitation is <500 mm and (ii) the sum of *Artemisia* and Chenopodiaceae (A+C)
pollen grains exceeds ~45 % of the total pollen sum. In a number of studies, the A/C
ratio has also been applied to pre-Holocene pollen assemblages, such as the Pleistocene (Herb et al., 2015; Koutsodendris et al., 2018), Pliocene (Wang et al., 2006; Cai et al., 2012), and Miocene (Hao et al., 2012a).

Considering the marked differences in *Artemisia* and Chenopodiaceae abundances within Central Asian pollen assemblages since the Miocene (Miao et al., 2011a,b; Cai et al., 2012; Figure 2) and their differences in water demand regarding seasonal precipitation distribution (El’Moslimany, 1990; Cour et al., 1999), the question arises since when the A/C ratio in Central Asian pollen records provides reliable estimates for moisture availability. To answer this question, we apply the criteria of Zhao et al. (2012) in their investigation of the A/C signal and its fidelity based on modern pollen datasets from the Tibetan and Chinese Loess Plateaus on our pollen datasets from the Qaidam Basin. The A+C percentages in our samples from the SG-1 and SG-1b cores are consistently >45 % since the late Miocene (~6 Ma) (Figure 4), suggesting that the A/C ratio may be used as a regional moisture indicator from that time onwards. However, the A/C ratios remain consistently below 1 in all samples between ~6 and ~3.6 Ma (Figure 4). Taken at face value, the A/C ratio would hence suggest continually dry conditions for the late Miocene and early Pliocene. However, this inference would contradict a large set of proxy information from Central Asia that consistently document more humid conditions during that time than during the late Pliocene and Pleistocene (An et al., 2001, 2005; Sun et al., 2010). Hence, the A/C ratio on pollen data from the Qaidam Basin most likely yields erroneous results for samples older than ~3.3 Ma. This can be attributed to the fact that the pollen assemblages, although containing both *Artemisia* and Chenopodiaceae, were dominated by Chenopodiaceae pollen (~35 % on average), with *Artemisia* pollen typically occurring in low numbers only (~12 %; Figure 2). By extension, the A/C ratio should be applied to fossil pollen spectra of the Qaidam Basin only after the strong expansion of *Artemisia* at ~3.3 Ma.
The A/C values in the Qaidam Basin for the past 3.3 Ma show a cyclical behaviour that allows us to distinguish between drier and wetter intervals (Figure 4). These intervals broadly correlate with glacial–interglacial variability evidenced in sedimentological and biomarker data in core SG-1 during the Pleistocene (Koutsodendris et al., 2018), which is driven by the interplay of the summer and winter monsoon systems (An et al., 2001; Guo et al., 2009; Sun et al., 2010). Interestingly, the A/C maxima show an increasing trend throughout the late Pliocene/Pleistocene, with peak values during the mid-Pleistocene (Figure 4). This is in concert with climate reconstructions from the Chinese Loess Plateau, which document a gradually increasing influence of the Central Asian summer monsoon towards the youngest Pleistocene interglacials (An et al., 2001; Sun et al., 2010).

Most notably, the A/C peak at ~0.5 Ma in the Qaidam Basin falls within MIS 13. This interglacial stands out as a time when a maximum in the summer monsoon coincided with a minimum in the winter monsoon, resulting in the lowest inland aridity in Central Asia for the entire Pleistocene (Guo et al., 2009; Hao et al., 2012b). The excellent agreement between our data from the Qaidam Basin and independent proxy records from the Chinese Loess Plateau suggest that the A/C ratio can be applied with high confidence as an indicator for moisture availability in arid Central Asia after the establishment of *Artemisia*/Chenopodiaceae-dominated steppe/deserts at ~3.3 Ma.

6. Conclusions

A new, continuous pollen and spore record has been generated from drillcore material from the western Qaidam Basin (NE Tibetan Plateau), allowing insights into the timing and underlying forcing of major shifts in vegetation composition of Central Asia during the past ~7 Ma. We find that the terrestrial ecosystems experienced a fundamental turnover during the late Pliocene associated with the establishment of *Artemisia*/Chenopodiaceae-dominated steppe/semi-desert biomes that exist until
today. Comparison with regional terrestrial and marine records strongly suggests that this vegetation shift was synchronous with the onset of drier conditions in Central Asia as a response to Tibetan Plateau uplift and the expansion of ice sheets in the high latitudes of the Northern Hemisphere shortly before the Plio/Pleistocene transition. Most notably, the strong expansion of *Artemisia* at ~3.6 and ~3.3 Ma at the expense of other steppe/semi-desert taxa coincides with the first glaciales of the Pliocene. This suggests that plant communities in Central Asia were highly susceptible to the effects of the early Northern Hemisphere glaciations although the extent of these glaciations was relatively small when compared to their Pleistocene counterparts.

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