Sporopollen evidence for Late Miocene stepwise aridification on the Northeastern Tibetan Plateau

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Abstract

Holding a climatically and geologically key position both regionally and globally, the northeastern Tibetan Plateau provides a natural laboratory for understanding the interactions between tectonic activity and the evolution of Asian aridification. Determining when and how the Late Miocene climate evolved on the northeastern Tibetan Plateau may help us understand better relations between tectonic uplift, global cooling and ecosystem evolution. Previous paleoenvironmental research has focused on the western Longzhong Basin. Late Miocene aridification data derived from sporopollen now requires corroborative evidence from the eastern Longzhong Basin. Here, we present a Late Miocene sporopollen record from the Tianshui Basin in the eastern Longzhong Basin. Our results show a two-stage stepwise aridification: a temperate forest with a more humid climate developed in the basin between 11.4 and 10.1 Ma, followed by a temperate open forest environment with a less humid climate between 10.1 and 7.4 Ma; and an open temperate forest-steppe environment with a relatively arid climate occupied the basin during 7.4 to 6.4 Ma. The vegetation succession demonstrates that Asian aridification occurred after ∼7–8 Ma, which is confirmed by other evidence from Asia. Furthermore, this persistent aridification on the northeastern Tibetan Plateau parallels the global cooling of the Late Miocene; the stepwise vegetation succession is consistent with the major uplift of the northeastern Tibetan Plateau during this time. These integrated environmental proxies indicate that global cooling may have been a potential driving force for Asian interior aridification, most likely enhanced by stepwise uplift of the Tibetan Plateau.

1 Introduction

As the latter stage of the global Cenozoic cooling, the Neogene was a critical period for northern hemispheric aridification, especially the marked aridification of the Asian interior. As the world’s largest arid region in a temperate zone, it not only provides dust
material for the Chinese Loess Plateau (CLP), but also influences modern Asian social development. Establishing when, and how, this process of aridification began and evolved is therefore vital for elucidating the interactions between tectonic uplift, global cooling and ecosystem evolution. Although there is compelling evidence for the aridification of the Asian interior, there is no consensus vis-à-vis its evolution and driving mechanisms. For instance, previous researchers have suggested that aridification of the Asian interior began in the Late Miocene, based particularly on biological and isotopic evidence (Eronen et al., 2012; Andersson and Werdelin, 2005; Wang and Deng, 2005; Zhang et al., 2012; Quade et al., 1989; Dettman et al., 2001; Cerling et al., 1997). However, others have argued that the process of Asian aridification may have begun in the Early Miocene (22 Ma) or even earlier (in the Late Oligocene), as inferred from Miocene or Oligocene eolian deposition (Guo et al., 2002, 2008; Qiang et al., 2011; Sun et al., 2010). The particular driving mechanisms of such aridification also remain enigmatic. Up until now, the tectonic uplift of the Tibetan Plateau (TP), global cooling, polar ice-sheet volumes and land–sea distributions etc. have been suggested as the major drivers (Kutzbach et al., 1993; An et al., 2001; Liu and Yin, 2002; Molnar et al., 2010; Gupta et al., 2004; Miao et al., 2012). However, the question of which one of these has principally controlled Asian climatic evolution is far from agreed. We focused on an ideal region – the northeastern TP – to explore the nature of the interactions between tectonics and the climate.

The geographically-extensive Longzhong Basin, consisting of series sub-basins, is located in the northeastern TP. These sub-basins present a continuous record of mammalian fossil-rich Cenozoic sediments, recording the effect of TP uplift on regional climates (GRGST, 1984; Fang et al., 2003, 2005; Li et al., 2006, 2014). On the other hand, it lies in the so-called monsoonal triangle zone, a transition zone from a warm-humid Asian monsoonal climate, to a dry-cold inland climate, to the alpine climate of the TP (Li et al., 1988, 2014) (Fig. 1a). Its particular geological and geographical characteristics make it sensitive to the accurate recording of the aridification of northern China through and the evolution of the Asian Monsoon. As an ideal field laboratory for
studying tectonic-climate interactions (Molnar et al., 2010; Tapponnier et al., 2001), the Longzhong Basin region might be the most promising for distinguishing TP uplift and any associated environmental change.

As a reliable paleoenvironmental proxy, sporopollen has been used to reconstruct past climates because of its abundance and excellent preservation within sediments. Previous research has demonstrated that the Tianshui Basin, as a sub-basin of the Longzhong Basin, exhibits a typical Late Miocene lacustrine-fluvial sedimentary succession containing abundant pollens. Here we reconstruct a high-resolution palynological record from the well-dated Yaodian Section, located in the southern part of the Tianshui Basin. Our results not only provide new evidence for the evolution of vegetation in the Late Miocene and climate change on the northeastern margins of the TP, but also shed new light on the aridification of the Asian interior through time. We also discuss the possible relation between tectonic activity and climate change.

2 Geological and geographical settings

The rhomboid-shaped Longzhong Basin, which is one of the largest intermountain and fault-controlled sedimentary basins on the northeastern TP, is geographically delineated by the left-lateral strike-slip Haiyuan Fault to the north, the Liupan Shan Fault to the east and northeast, the Laji Shan Fault to the southwest, and the Western Qinling Fault to the south (Fig. 1b). The Tianshui Basin, one of its sub-basins, is located in the southeastern part of the Longzhong Basin (Fig. 1b). It has witnessed the continuous deposition of mammalian, fossil-rich Cenozoic sediments from the surrounding mountains; these sediments record the interactions between mountain uplift, erosion and climate change (Li et al., 2006; Alonso-Zarza et al., 2009; Peng et al., 2012). At present, the East Asian Monsoon influences this region, engendering a semi-humid, warm temperate, continental monsoon climate, characterized by relatively hot, humid summers and cold, dry winters. The mean annual temperature and mean annual precipitation of this area are ~ 10.4 °C and 504 mm, respectively, with rainfall concentrated
mainly in the summer and autumn (Fig. 1c). The modern natural vegetation in this region is warm-temperature forest-grassland. Warm grasslands are distributed in the valleys, and consist mainly of *Arundinella hirta*, *Spodiopogon sibiricus* and *Themeda triandran*. Shrubs such as *Zizyphus jujube*, *Sophora viciifolia* and *Ostryopsis davidiana* are found on the hillsides. Trees, including *Quercus liaotungensis*, *Pinus tabulaeformis*, *P. armandi* and *Platycladus orientalis*, grow in the mountains (Huang, 1997).

The selected Yaodian Section (105°55′ E, 34°38′ N) is located in the southern part of the Tianshui Basin (Fig. 1d). The Neogene sequence in this section is capped by Quaternary loess and lies unconformably on top of the Paleogene Guyuan Group. It has been divided into the Ganquan Formation (Fm), the Yaodian Fm and the Yangjizhai Fm, in sequence upwards (Li et al., 2006). In this study, our research mainly focuses on the Late Miocene Yaodian Fm and Yangjizhai Fm. Based on a determination of lithology and sedimentology, the Yaodian Fm can be divided into three principal strata. The lower stratum consists of massive fine gravel sandstone, sandstone and brown silty mudstone, occasionally with thin brown mudstone or interbedded paleosols, which can be considered fluvial channel deposits (Fig. 2e). Abundant teeth of *Hipparion weihoense*, *Cervavitus novorossiae*, *Ictitherium* sp. and their bone fragments were excavated from this stratum. The middle stratum of the Yaodian Fm consists of the interbedding of siltstone or fine sandstone with mudstone intercalated with paleosols, overlying the fluvial channel deposits. This assemblage’s characteristics are typical of floodplain deposition (Fig. 2d). The upper stratum of the Yaodian Fm is characterized by rhythmic cycles composed of grey or brown mudstone or sandy marlite and intraclastic marl intercalated with brown siltstone and mudstone, and contains fossil algae and gastropods; this section is representative of shallow lake deposition (Fig. 2a and c). This upper stratum is common throughout the basin, and is analogous to the “Zebra Bed” stratum found in the Linxia Basin in the western Longzhong Basin (Li, 1995). The Yangjizhai Fm is principally composed of reddish brown mudstone or silty mudstone and yellow-brown calcrite or calcareous mudstone, with scattered sandstone or grey mudstone and marlite. These sediments were deposited under strong evaporative conditions in
distal floodplain to palustrine environments (Fig. 2b). Previous paleomagnetic investigations have indicated that the Yaodian Fm ranges from 11.67 to 7.43 Ma in age, and that the Yangjizhai Fm dates from 7.43 to 6.40 Ma, both these ranges being consistent with the formations’ biostratigraphic ages (Li et al., 2006).

3 Materials and methods

Most of the samples came from lacustrine mud deposits and fine grain size intercalations found in floodplain and fluvial channel deposits. Because the lower 10 m of the Yaodian Fm consists of coarse gravel sandstone, and it was difficult to find fine-grained sediments therein, this part of the formation was not sampled. A total of 200 samples were processed for sporopollen analysis. For each sample, >100 g of sediment was washed in 20% HCl, soaked in 39% HF and then treated with 10% HCl solution to enable fluoride dissolution. The chemical processing was followed by physical enrichment procedures such as ZnCl₂ separation, ultrasound sieving with a 10 µm filter, and storage in glycerin. Sporopollen grains were identified by comparing samples with published modern and fossil sporopollen plates (Wang, 1995; Song, 1999), as well as modern sporopollen slides preserved in the Laboratory of Sporopollen Analysis of the Geography Department of Lanzhou University.

4 Results

Only 126 of the 200 samples contained enough palynomorphs to provide reliable data; the remaining 74 possessed fewer than 300 identifiable grains. Most of these latter samples had been preserved under oxidizing conditions, or had abundant carbonate content. Approximately 80 different palynomorphs were identified into family or genus. In order to synthesize the information and highlight palynological trends, the major taxa, and their percentage contents vis-à-vis recognized taxa, were used to construct a sporopollen diagram (Fig. 3).
This diagram principally demonstrates that tree pollen declines stepwise as herbaceous pollen increases. Tree pollen consists mainly of *Pinus*, Cupressaceae and *Ulmus*, along with *Quercus* and *Betula*. Additionally, a number of subtropical plants such as *Liquidambar*, *Pterocarya*, *Carya* and *Euphorbiaceae* (which is no longer found in this area today), appear often, but their contents are lower. Herbaceous pollen is mainly from *Artemisia*, Chenopodioideae, Poaceae and Asteraceae. Pollen from extremely drought-tolerant plants, such as *Ephedra* and *Nitraria*, only appear sporadically in individual samples. In addition, this section also contains some ferns and *Pediastrum*. Stratigraphically-constrained cluster analysis (CONISS) yields three distinct zones in the section, described from the bottom up as follows:

### 4.1 Zone 1 (11.4–10.1 Ma)

Samples from this zone exhibit high percentages of tree pollen (averaging 75.13 %) and relatively less herbaceous pollen (averaging 23.42 %). Coniferous taxa are mainly *Pinus* (18.57 %) and Cupressaceae (17.62 %), with smaller amounts of *Picea* and *Cedrus*. *Ulmus* (19.71 %) is the most common broadleaf tree, accompanied by *Betula* (2.73 %), *Quercus* (2.17 %) and *Salix* (2.10 %). Other arboreal taxa are *Juglans* and *Castanea*, with < 2 % each. Herbaceous taxa mainly include *Artemisia* (7.41 %), Chenopodioideae (5.51 %) and Poaceae (2.44 %), along with small amounts of Asteraceae, Ranunculaceae and Rosaceae, with amounts < 2 % each. Aquatic plants, algae and some subtropical taxa are also present in this zone, though with low contents.

### 4.2 Zone 2 (10.1–7.4 Ma)

In samples from this zone, total tree pollen content decreases, averaging 53.98 %, while the percentage of herbaceous pollen (averaging 42.93 %) increases. Coniferous taxa are principally represented by *Pinus* (13.95 %), Cupressaceae (7.48 %), *Picea* (2.16 %) and *Cedrus* (1.49 %). Among broadleaf trees, the dominant taxa are *Ulmus* (averaging 8.04 %), *Quercus* (2.35 %), *Betula* (2.28 %), *Salix* (1.89 %) and *Juglans*
(1.42%). Herbaceous taxa are dominated by *Artemisia* (13.68%) and Chenopodioidae (8.83%), along with Poaceae (4.89%), Asteraceae (3.15%) and Ranunculaceae (2.94%). Aquatic vegetation appears successively and reaches the highest value found in the entire profile. There are also some subtropical taxa with low contents, such as *Liquidambar*, *Pterocarya*, *Carya* and Rutaceae. This zone is divided into two subzones, Zone 2-1 (10.1–8.6 Ma) and Zone 2-2 (8.6–7.4 Ma). Herbaceous pollen percentages are slightly higher in Zone 2-2 than in Zone 2-1.

### 4.3 Zone 3 (7.4–6.4 Ma)

The samples from this zone record a sudden decrease in tree pollen (with content averaging 39.20%), mirrored by an increase in herbaceous pollen (averaging 60.08%). Coniferous taxa are characterized by *Pinus* (7.24%) and Cupressaceae (5.26%). *Ulmus* (5.28%) dominates the broadleaf tree category, with *Quercus* and *Betula* accounting for 2.33 and 2.32%, respectively. Herbaceous taxa are composed of *Artemisia* (19.42%), Chenopodioidae (11.23%) and Poaceae (8.59%), together with Asteraceae (4.53%), Ranunculaceae (3.23%), Brassicaceae (3.18%) and Polygonaceae (1.86%). Aquatic plants and thermophilic species almost disappear.

### 5 Discussion

#### 5.1 Vegetation and climate reconstruction

When discussing the relation between the palynological spectrum and sedimentary facies in the Yaodian Section, it becomes apparent that these sedimentary facies have experienced four successive depositional stages: fluvial channel; floodplain; shallow lake; and distal floodplain to palustrine. Transitional ages can be dated to 10.4, 9.23 and 7.43 Ma, respectively (Li et al., 2006) (Fig. 2). Our palynological spectrum shows stepwise changes at 10.1 and 7.4 Ma, lagging slightly behind those evinced by the sedimentary facies. Another distinctive feature of the palynological spectrum is that the
green lacustrine deposits of fine grain size exhibit dense palynomorph concentrations, with higher tree pollen percentages. In contrast, the reddish floodplain deposits with coarse grain sizes possess sparse palynomorph concentrations, with higher herbaceous pollen percentages (Fig. 3). However, in the same sporopollen zones, we find that the palynomorph concentration clearly changes between different sedimentary facies, but that any variation in overall percentage content is minor. Interestingly, between different sporopollen zones, the palynomorph percentage changes greatly vis-à-vis the same sedimentary facies. We can therefore conclude that these changes in the palynological spectrum are caused by changes in regional vegetation, rather than differing preservation conditions. The paleoecological information inferred from the percentage change of sporopollen record can thus be considered reliable.

According to modern surface pollen studies, Pinus is often overrepresented in pollen records because of its abundant pollen production and the ease with which this pollen is transported over long distances by the wind. As a general rule, it can be assumed that there is/was no proximate pine forest if less than 25 to 30\% of Pinus pollen occurs in samples (Li and Yao, 1990). Higher percentages of Cupressaceae and Taxodiaceae coexistent with temperate tree, shrub and herbaceous pollen may reflect a warmer, wetter and more humid climate (Song, 1978). Nowadays, Ulmus is commonly distributed in the sub-humid temperate and warm temperate mountain foothills of northern China, but percentages of its pollen collected from CLP surface soils never exceed 1\%, even under broadleaved forests containing elm (Liu et al., 1999). In general, when their content exceeds 3–5\% of total arboreal pollen content, birch and oak can be considered to be/have been present in woodland (Liu et al., 1999). Salix produces very little pollen, and most of this pollen falls near the tree itself (Li et al., 2000). Modern Artemisia and Chenopodioidae are extensively distributed throughout the arid and semi-arid regions of China. Chenopodioidae are more drought-resistant than Artemisia. Artemisia being the dominant vegetation type may reflect a semi-arid grassland environment, while Chenopodioidae may reflect an arid desert environment. Surface pollen analysis shows that Artemisia and Chenopodioidae are greatly overrepresented in pollen
rain. Only when Chenopodioidae and *Artemisia* pollen content exceeds 30% of the total should their presence be considered as primarily local (Herzschuh et al., 2003; Ma et al., 2008). Poaceae pollen content is sparse, usually only 3–6%, even when it represents the dominant modern species (Tong et al., 1995).

Our record therefore indicates that, during the period when the Yadian Fm was being deposited, the study area was dominated by temperate forests and a warm and humid climate. Mixed deciduous forests, characterized by the dominance of *Pinus*, Cus- pressaceae, *Ulmus* and *Quercus*, were distributed within the basin and the low altitude hills surrounding it. Mid- and high-altitude vegetation, such as *Abies*, *Picea* and *Cedrus*, existed in the surrounding uplands. The river banks or lake margins were colonized by *Salix*, *Alnus*, *Fraxinus* and *Taxodiaceae*. Cyperaceae, *Typha* and *Myriophyllum* grew along the lake shores or in shallow water areas. Ranunculaceae, *Poaceae*, *Chenopodioidae* and *Artemisia*, principally occupied the forest understory, or were distributed in forest clearings. However, as indicated by our record, the environment was not static. During 11.4–10.1 Ma, temperate forest and a more humid climate developed in the basin. The growth of fluvial channel deposits and the presentation of a large number of mammalian fossils (Li et al., 2006) also support the theory that much denser vegetation capable of supporting large mammals such as rhinoceroses developed during this interval. Moreover, we know that the northern Tianshui Basin was dominated by temperate and warm-temperate deciduous broadleaf forest (Hui et al., 2011). Our result is also consistent with research into the climatic evolution of the Qaidam Basin, which found that the presence of $\delta^{18}$O values characteristic of large mammals indicated a warmer, wetter, and perhaps lower-altitude Qaidam Basin (Zhang et al., 2012). The early Late Miocene mammal fauna discovered in the Qaidam Basin also reflected a wooded environment, in which many streams with aquatic plants such as *Trapa* and *Typha* developed (Wang et al., 2007). From 10.1–7.4 Ma, the study area was dominated by a warm-temperate open forest environment and an arid climate, relative to the previous interval. Sedimentary facies become characteristic of shallow lake deposits (Li et al., 2006). Mammal fauna identified in the eastern Qaidam Basin also indicate that
a mixed habitat of open and wooded environments, with abundant freshwater streams, was predominant at that time (Wang et al., 2007). During this interval, after ~ 8.6 Ma, herbaceous plants also increased their presence in the Tianshui Basin, as confirmed by mammalian fossil records. In the northern Tianshui Basin at ~ 9.5 Ma, there is evidence of a sizeable rhinoceros population, which would have required a relatively moist woodland environment to sustain itself. However, the typical Hipparion fauna that presents ~ 8.0 Ma probably represents a relatively temperate climate with a more mixed vegetative landscape, i.e. an open forest environment rather than a vast, open landscape. Large mammals would still have been able to survive in such an environment (Zhang et al., 2013).

An open temperate forest-steppe environment clearly developed in the study region, indicating significant aridification after ~ 7.4 Ma. Grassland, composed principally of Poaceae, Artemisia and Chenopodioidae, developed in most of the basin, while shrinking areas of open forest, dominated by Cupressaceae, Ulmus and Quercus, existed in the surrounding mountains. Salix continued to grow in relatively humid environments such as riverbanks. Distal floodplain to palustrine deposits now characterize the study area (Li et al., 2006). A sudden increase in magnetic susceptibility after ~ 7.4 Ma may indicate the enhanced oxidation characteristic of an arid environment (Fig. 4b). In the northern part of the Tianshui Basin, drought-tolerant Artemisia predominates after 7.4 Ma, further confirming the presence of a drier climate (Hui et al., 2011) (Fig. 4c). Further, the growing presence of grazer mammalian species at the end of the Miocene in the Tianshui Basin suggests that the local environment was principally occupied by grassland, with some woodland, and even some desertification (L. P. Liu et al., 2011) (Fig. 4d). Furthermore, the gradual increase in eolian sediments after 7.4 Ma in the Linxia Basin would indicate a period of intense desertification in central China (Fan et al., 2006) (Fig. 4e). Biomarker evidence from the Linxia Basin also indicates a distinct change in the climate toward arid-cold conditions at ~ 8 Ma (Wang et al., 2012). The isotopic compositions of herbivorous fossil teeth and paleosols from the Linxia Basin (Wang and Deng, 2005) and southwestern China (Biasatti et al., 2012) also
indicate a shift to a drier, or seasonally drier, local climate. In the Qaidam Basin, *Hipparion teilhardi* fossils characterized by slenderer distal limbs, and dated to the end of the Miocene, imply an adaptation by this animal to the open steppe environment (Deng and Wang, 2004). Marine sediments also indicate that the climate changed at this time. For example, local seawater $\delta^{18}O$ reconstructions from ODP Site 1146 in the northern South China Sea suggest that the climate of East and South Asia shifted toward more arid conditions after $\sim 7.5$ Ma (Steinke et al., 2010) (Fig. 4f).

### 5.2 More arid conditions at the end of the Miocene and their possible causes

Based on the currently-available data from Late Neogene Chinese mammalian fossils, Zhang (2006) suggested that mammal communities in northern China were rather stable and uniform from $\sim 13$ Ma to the end of the Miocene ($\sim 7$–$8$ Ma), and that differentiation between the humid fauna communities prevalent in eastern China and the dry fauna communities identified in western China occurred after the end of the Miocene. The diversity in Bovidae fossils also increases significantly toward the end of the Miocene, with some genera even appearing in southwestern China (Chen and Zhang, 2009), indicating an expansion in grasslands and an aridification of the climate. Using macro- and microfloral quantitative recovery techniques to reconstruct the climate in northern China at the time, Y.-S. C. Liu et al. (2011) proposed that the west–east temperature and precipitation gradient pattern did not develop in northern China until the end of the Miocene. This corroborates the quantitative results gained from using mammalian fossils as a proxy for paleoprecipitation (Liu et al., 2009). A semi-quantitative reconstruction of Chinese Neogene vegetation also indicates that the aridification of western, central and northern China occurred during the Miocene–Pliocene transition (Jacques et al., 2013). Indeed, in order to adapt to the arid climate in northern China during the end of the Miocene, some plants and arthropods also evolved more arid-tolerant species, such as *Frutescentes* (Fabaceae) (Zhang and Fritsch, 2010), *Ephedra* (Ephedraceae) (Qin et al., 2013) and *Mesobuthus* (Buthidae) (Shi et al., 2013). This marked aridification has been well documented in other parts of Asia. For example,
dramatic changes in the carbon and hydrogen isotopic ratios of leaf waxes at ODP Site 722 indicate an increasing aridity at the end of the Miocene in continental source regions, including Pakistan, Iran, Afghanistan, and the Arabian Peninsula (Huang et al., 2007) (Fig. 4g). The isotopic compositions of herbivorous fossil teeth and paleosol carbonates also suggest that the climate became drier over the Indian Subcontinent, China, and Central Asia during toward the end of the Miocene (Quade et al., 1989; Cerling et al., 1997; Wang and Deng, 2005; Biasatti et al., 2012; Barry et al., 2002; Badgley et al., 2008; Zhang et al., 2009). The evidential synchronicity of these climatic events in Asia strongly suggests that a process of aridification began toward the end of the Miocene (∼7–8 Ma). The onset of such a marked aridification is further corroborated by the presence of red clay across much of the CLP (An et al., 2001).

In our study, the vegetation succession appears to be consistent with the general direction of global cooling, indicating that global cooling may have been a potential driving force for the aridification of the Asian interior (Fig. 4h). Because atmospheric moisture content decreases as air temperature cools, one would expect global cooling to lead to a gradual net global aridification (van Dam, 2006). However, drought occurs in a stepwise fashion. Although the most significant Late Cenozoic global cooling event occurred at ∼14 Ma (Zachos et al., 2001), only minor cooling events at ∼10.1 Ma and ∼7.4 Ma have been documented (Fig. 4h). The Late Miocene climate was much warmer and/or wetter than today in many regions (Pound et al., 2012). Evidently, factors other than global cooling also exert a strong effect on precipitation regimes, particularly paleotopography and continental tectonic configuration, which themselves affect monsoonal patterns.

The uplift of the TP is a potentially significant trigger of Asian aridification. First, although no consensus exists vis-à-vis the timing and degree of any uplift, many studies have suggested that the interval ∼8–10 Ma not only experienced rapid uplift of the TP (Molnar et al., 2010; Fang et al., 2003, 2005; Li et al., 2014; Enkelmann et al., 2006; Zheng et al., 2006, 2010; Wang et al., 2012, 2006; Lease et al., 2007) (Fig. 4i), but that this uplift achieved an altitude sufficient to block the penetration of moisture from
the Indian Ocean into western China (Dettman et al., 2001, 2003). Second, the main source of water vapor for arid northwestern China has been the Asian Summer Monsoon; the Asian Winter Monsoon promotes a cold and dry climate. Atmospheric and coupled atmosphere–ocean models show that such massive, continent-spanning orogen uplift may have generated, or intensified, Asian monsoon circulation (Kutzbach et al., 1993; An et al., 2001). Moreover, the effect of TP uplift on enhancement of the Asian Winter Monsoon was more significant than on the Asian Summer Monsoon (Liu and Yin, 2002). There is increasing evidence that the Asian Summer Monsoon gradually weakened after ~ 10 Ma (Wan et al., 2010; Clift et al., 2008), while the Asian Winter Monsoon gradually strengthened, particularly toward the end of the Miocene (An et al., 2001; Jacques et al., 2013; Clift et al., 2008; Sun and Wang, 2005; Jia et al., 2003). Therefore, global cooling, most likely enhanced by TP uplift, may be a potential driving force for the aridification of the Asian interior.

6 Conclusion

The Late Cenozoic basins in the northeastern TP document both the tectonic uplift process and its associated environmental changes. We investigated a Late Miocene sporopollen record from the Tianshui Basin in the northeastern TP. Our results indicated that a temperate forest, with a more humid climate regime (11.4–10.1 Ma), gave way to a temperate open forest environment with a less humid climate (10.1–7.4 Ma); this was in turn replaced by an open temperate forest-steppe landscape, accompanied by a relatively arid climate (7.4–6.4 Ma). This vegetation succession demonstrates that aridification of the Asian interior occurred after ~ 7–8 Ma, as corroborated by other studies of Asia. Our findings support the idea that global cooling may have been a potential driving force for aridification of the Asian interior, and that TP uplift probably enhanced this process. This study highlights these possible interactions between global cooling and tectonic uplift.
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Figure 1. Geographic setting of Yaodian Section. (a) The location of the Longzhong Basin. (b) The major tectonic faults of the Longzhong Basin. (c) Mean monthly temperature and mean monthly precipitation between in the Tianshui area, 1971–2000. (d) Geological map of the Tianshui Basin.
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nus mesolophidus, Mimomys teilhardi, Paenepetence zhudingi, Sinotamias sp., Ochotona gra-
cilis, O. lagreli, O. lingtaica, O. minor, O. plicodenta, O. sp., Ochotonoma sp., O. primitiva,
Trischizolagus mirificus, Hipparchion chiai, H. dermatorhinum, H. fossatum, H. plocodus, H. sp.,
H. weihooense, and Gazella sp.; grey triangles represent species which adapt to relatively hu-
mid environment, including Chleuastochoerus stehlini, Cervavitus novorossiae, Cervidae gen.
et sp. Indet., Palaeotragus microdon, P. sp., Samotherium sinense, S. sp., Rhinocerotidae ind-
et., Acerorhinus fuguebsis, Chiloitherium habereri, C. sp., C. wimani and Protanancus tobieni.
(e) Eolian sediment mass accumulation rates for the Linxia Basin, northeastern TP (Fan et al.,
2006); (f) South China Sea δ¹⁸Oseawater estimates from ODP Site 1146 (Steinke et al., 2010);
(g) Compiled global deep-sea δ¹⁸O values (Zachos et al., 2001); (h) Schematic model show-
ing major periods of TP uplift (Molnar et al., 2010; Fang et al., 2003, 2005; Li et al., 2014;
Enkelmann et al., 2006; Zheng et al., 2006, 2010; Wang et al., 2012, 2006; Lease et al., 2007).