The Miocene Climatic Optimum: evidence from ectothermic vertebrates of Central Europe

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Abstract

Data sets of Central European temporal distributions of thermophilic ectothermic vertebrates (Channidae, Varanidae, Chamaeleonidae, Cordylidae, Tomistomidae, Alligatoridae, giant turtles) and of North Alpine Foreland Basin (NAFB) distributions of ectothermic vertebrates adapted to dryer habitats (Albanerpeton inexpectatum, Salamandra sansaniensis, Bufo cf. viridis, Chamaeleo caroliqurti, Gekkonidae, Varanus hofmanni, Bransateryx sp.) are analysed. Two main migration events of thermophilic ectotherms at 20 Ma and 18 Ma in the Lower Miocene are discerned. They indicate the beginning of the Miocene Climatic Optimum in Central Europe (42–45°N palaeolatitude) with a lower limit of the mean annual temperature (MAT) of 17.4°C derived from the minimal MAT of their extant relatives. Furthermore, additional palaeobotanical data and records of bauxite point to a MAT of 22°C. This warm and humid optimum peaked at 18–16.5 Ma (Ottnangian, Karpatian), and is confirmed by the coexistence of all investigated thermophilic taxa. The following period (Early Badenian) is characterised by probably unchanged temperatures but a seasonality in precipitation with dry periods up to six months. Two major seasonal phases between 16.3 and 15.7 Ma (earliest Early Badenian) and between 14.7 and 14.5 Ma (Early/Middle Badenian transition) are indicated by an immigration of dry adapted taxa from the surrounding karst plateau to the NAFB. It is presumed that the tectonical reorganisation of the Central Paratethys realm had considerable influence on this regional humidity pattern. The warm period ended abruptly between 14.0 and 13.5 Ma (Middle/Late Badenian transition) with major regional extinction events of most of the thermophilic groups in Central Europe and a drop of the MAT of probably more than 7°C to temperatures around 14.8–15.7°C. This drop can be attributed predominantly to a decrease of more than 11°C of the minimum cold months temperature. This temperature decrease marked the beginning of a climatic zonation of the European continent and is also evidenced by a progressively southward disappearance of the crocodile Diplacynodon from 38–45°N palaeolatitude to 30–37°N during the Middle and earliest Late Miocene. The results correlate well with palaeobotanical data from the mid-latitudes of Europe and North America, and the deep-sea temperature curve generated from oxygen isotope ratios.

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1. Introduction

The complex history of global climate evolution

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(M. Böhme).
during the Cenozoic is being reconstructed in more and more detail, which is especially due to the high-resolution deep-sea δ¹⁸O isotope records, and the establishment of an ‘astronomically calibrated’ geological time scale (Zachos et al., 2001). One of the most remarkable episodes is the Mid-Miocene Climatic optimum, between 17 and 15 Ma. According to Flower and Kennett (1994), this climax of Neogene warmth was followed by short-term variation in the East Antarctic Ice Sheet volume, sea level, deep ocean circulation and in global climates (between ~16.0 and 14.8 Ma). A major and permanent cooling step occurred between 14.8 and 14.1 Ma, associated with increased production of cold Antarctic deep waters and a major growth of the East Antarctic ice sheet. As a result, the meridional temperature gradient increased, the boundaries between climatic zones strengthened and the aridification of the mid-latitudes increased (Flower and Kennett, 1994).

The climatic zonation of the world is reflected by the zonal distribution of vegetation types. But, in contrast to plants, animals have looser affinities to distinct climatic zones (Ostrom, 1970; Walter and Breckle, 1983). Many ectothermic vertebrates are physiologically adapted (e.g. due to dormancy) to seasonal climatic change. Thus, many reptiles of the tropics occur also in subtropical conditions (Haller-Probst, 1997). Therefore, tropical climate (daily thermal gradient > annual thermal gradient) cannot be recognised with certainty based on ectothermic vertebrates only, because in most cases only the lower limit of temperature is practically useful. Other proxies, such as palaeobotanical or geochemical data can help to improve the analysis.

Many recent studies of climate/evolution paradigms have focused on mammals. E.g. Alroy et al. (2000) claim that the global climate change had only unpredictable effects on the Cenozoic mammalian evolution in western North America. In contrast to mammals, however, fishes, amphibians and reptiles are ectothermic and their distribution is highly dependent on temperature (Markwick, 1998). Contrary to mammals, evolutionary rates of ectotherms are low, and for example within the Neogene only a few speciation events are recognisable (especially in amphibians – most Neogene species belong to extant species or species groups). The most important effects on the distribution and regional diversity of ectothermic vertebrates are migration and extinction events (Tihen, 1964; Böhme et al., 2001). Vertebrate migration events can generally be attributed to palaeogeographic (Rögl, 1999) and climatic events (Pickford and Morales, 1994). In the absence of physical barriers, climatic factors, such as temperature and humidity, may cause vertebrate migration (Van der Made, 1992). Several mammal (Van der Made, 1997; Rögl, 1999) and herpetofaunal (Böhme, 2001; Ivanov, 2001) migration events within and between the Old World continents of Africa, Asia and Europe have been detected within the Miocene. To investigate the influence of the Miocene global and regional climatic changes on migration and extinction events of ectothermic vertebrates, I have compared the temporal and spatial distribution of thermophilic and dry adapted ectotherms in Central Europe with data sets on Atlantic benthic foraminiferal oxygen isotope ratios.

2. Materials and methods

The studied record of Central European (palaeolatitude 42–45°N, see Meulenkamp et al., 2000) Miocene ectothermic vertebrates comes from 230 outcrops (mostly the author’s data, for complete occurrence matrix, see Appendix 1 in the online version of this paper) having ages ranging from 23 to 10 Ma ago (chronostratigraphical and geochronological data after Reichenbacher et al., 1998; Steininger, 1999; Daxner-Höck, 2001). Intensive sampling during the last few decades yielded about 111 taxa of a herpetological fauna. Most of the localities (113; mainly sampled by Kurt Heissig, Munich) belong to the ‘Older Series’ and ‘Middle Series’ of the Bavarian part of the North Alpine Foreland Basin (NAFB, palaeolatitude 42–43°N). Within the time interval 17–14.6 Ma ago, eleven sedimentary cycles (OSM 0–10, Heissig, 1997; Böhme et al., 2001) and six mammal units (OSM A–F, Heissig, 1997) have been recognised. Therefore, a time resolution of less
than 0.3 Ma is available for most of these localities. The remaining outcrops represent fissure fillings located immediately north of the NAFB (karstic plateaus of the Swabian and Franconian Alb, \( n = 38 \)) and stratified outcrops in surrounding areas such as the Swiss and Austrian parts of the Molasse Basin, the North Bohemian Brown Coal Basin, the Lower Rhine Embayment, the Allier Basin, and the Vienna Basin. The age uncertainty of these localities based on the mammal biostratigraphy and magnetostratigraphy is 0.3–0.5 Ma and only in some cases up to 1.0 Ma.

This large data set contains localities representing a wide range of environments (fissures, \( n = 48 \); rivers, \( n = 112 \); floodplains, \( n = 31 \); lakes, \( n = 22 \); ponds, \( n = 11 \); swamps, \( n = 4 \); coastal marine environments, \( n = 2 \); Böhme, 2000). The fossil material is disarticulated and is represented by isolated bones and teeth. In most cases their preservation is unweathered and unabraded. Assemblages from fluvial environments are found in channel-fill deposits and can be interpreted as generally autochthonous to their environment of deposition (Behrensmeyer, 1975, 1988). Especially the karstic and limno-fluvial environments (fissures, rivers, floodplains) are represented by a continuous record through the investigated interval, which gives evidence that the data set is comparable in terms of depositional and taphonomic environment.

I extracted the number of thermophilic species of ectothermic vertebrates as a proxy for temperature. I classified an animal as thermophilic if its extant relatives live exclusively or mostly in tropical or subtropical climates. These groups are: the snakehead fishes (Channidae), the monitor lizards (Varanidae), the cordylid lizards (Cordylidae), the chameleons (Chamaeleonidae), the crocodiles (Tomistomidae, Alligatoridae), and the giant turtles (Geochelone s.l.). The geographical distribution of modern representatives of these groups is shown in Fig. 1. As a proxy for seasonality in precipitation, I extracted ectotherms which are mainly adapted to areas with a lower groundwater table and dryer environment. An animal is characteristic of these conditions if its abundance maxima are within the contemporary fissure fillings of the surrounding karst plateau (approximately 50 km north) and occur with significantly lower frequencies and abundances in the limno-fluvial molasse deposits. Such animals (here called ‘dry adapted’) are the albanerpetontid (Albanerpeton inexpectatum), the salamander (Salamandra sansaniensis), the green toad (Bufo cf. viridis), the big chameleon (Chamaeleo caroliquarti), the gekkos (Gekkonidae), the monitor lizards (Varanus hofmanni), and the sand boas (Bransateryx sp.).

3. Results and analysis

The temporal distribution of the selected thermophilic taxa in Central Europe is shown in Fig. 2. The First Occurrence Datum (FOD) for the snakehead fishes is early MN4 (Neogene Mammal units after Mein, 1975) in the fissure filling at Hermaringen 1. The beginning of the MN4 is dated at 18 Ma (Steininger, 1999; Daxner-Höck, 2001). In the younger part of MN4 the Channidae are widespread in Western and Central Europe (e.g. La Romieu, France; Langenau, North Alpine Molasse Basin; Reichenbacher, 1993). The Last Occurrence Datum (LOD) of this group occurs between 13.2 and 12 Ma at the early–middle MN8 (Laimering 1b, Kleineisenbach, both Bavarian Molasse Basin; Anwil, Swiss Molasse Basin). The family Cordylidae is common in Eocene deposits and temporarily disappeared in Europe at the Eocene/Oligocene boundary (Auge, 1987). The next record is known from the terminal Oligocene locality Rott (Böhme and Lang, 1991) belonging to the Mammal Palaeogene unit MP30 (23.8–24.2 Ma, after Kempf et al., 1997). The FOD for the genus Palaeocordylus correlates to the earliest MN4 (fissure filling Petersbuch 2, 18 Ma). The LOD of the Cordylidae occurs during the early MN5 (latest Karpatian) at Puttenhausen.
Fig. 1. Geographical distribution of selected extant thermophilic taxa (from Haller-Probst, 1997, and unpublished data).
Fig. 2. Miocene time scale (after Berggren et al., 1995; Steininger, 1999; Daxner-Höck, 2001), litho- and biostratigraphical units of the Bavarian part of the Upper Freshwater Molasse of the NAFB (Böhme et al., 2001), distribution of thermophilic ectothermic vertebrates in Central Europe, the supposed evolution of the MAT (this paper, Fig. 5), and the global deep-sea oxygen isotope record (after Zachos et al., 2001). The shaded intervals indicate periods of higher seasonality in precipitation (this paper, Fig. 4).

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<tr>
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* Ries-Impact 14.47 ± 0.36 Ma; Soszynski et al. 1995; FOD, LOD;
and Obergänserndorf (Böhme, 2002; 16.5–16.7 Ma, after Daxner-Höck, 2001). Thus, the cordylid lizards have a short temporal distribution in Central Europe, probably not exceeding 1.5 Ma.

The FOD of the family Chamaeleonidae correlates to early MN3 (20–19.5 Ma). Its oldest representatives are described from Merkur (Fejfar and Schleich, 1994) and Wintershof-West (Moody and Rocc, 1980). The last unambiguous chamaeleonids from the Molasse Basin are to be found at the Swiss locality Ornberg (Bolliger, 1992, end of MN6, 13.7 Ma after Kempf et al., 1997), so that the LOD of Chamaeleonidae in Central Europe is at about 13.7 Ma.

The taxonomic status and the phylogeny of the European giant turtle *Geochelone* sensu latu is still unclear (De Broin, 1977) and needs a revision. This large animal (more than 1 m carapax length) is definitively present in Central Europe from the late MN3 (locality Stubersheim 3, NAFB) to the end of MN6. The youngest record of *Geochelone* s.l. is from the Bavarian part of the NAFB (locality Kirrberg), with an approximate age of 13.5–14.0 Ma.

Two crocodiles, the genera *Gavialosuchus* (sensu Buñetart et al., 1984, = Tomistoma of Antunes, 1994) and *Diplocynodon* occur in the Miocene of Central Europe. *Gavialosuchus* had its FOD in early MN3 at Eggenburg and its LOD in the Upper Marine Molasse of Southwest Germany (Bodensee area, locality of Bodman, Rossmann et al., 1999). The regression of the Upper Marine Molasse is diachronous (from east to west) and reached the East Switzerland/Bodensee area during late MN4 (Schlunegger et al., 1996; Kälin, 1997). Therefore, the minimum date for the LOD of *Gavialosuchus* is about 17 Ma. The regional extinction of *Gavialosuchus* can be correlated more likely with habitat shifting (regression of the Upper Marine Molasse – most of *Gavialosuchus* remains have been found in marine and littoral sediments, Buñetart et al., 1984; Antunes, 1994) than with climatic deterioration.

The genus *Diplocynodon* is known from Europe since the Middle Eocene (Berg, 1966; Ginsburg and Bulot, 1997). The LOD in Central Europe is recorded in the locality Kirrberg (late MN6) from the Bavarian part of the NAFB. The LOD of *Diplocynodon* in Central Europe has been estimated at 13.5–14.0 Ma, i.e. contemporaneous with that of the chamaeleonids and giant turtles. Based on the big hamster *Cricetodon* cf. *sansaniensis* (Rummel and Kälin, 2003), the aforementioned localities (Ornberg, Kirrberg) are well dated.

In this context, a single crocodile tooth from the late Middle Miocene locality Anwil (Swiss Molasse Basin, early MN8, 13.2 Ma after Kälin and Kempf, 2002) is surprising. Since reworking can be excluded (B. Engesser, Basel, pers. commun., 2001), a somewhat longer occurrence in the southeastern part of the NAFB (41.5°N palaeolatitude) could be possible. In France, *Diplocynodon* disappears just as in Central Europe at the end of MN6 (LOD at the locality Sansan, Ginsburg and Bulot, 1997); on the Iberian Peninsula its record reaches to the lowermost Upper Miocene (Antunes, 1994; MN9, 9.7–11.1 Ma). In Southeast Europe (Bulgaria) *Diplocynodon* also reaches the Upper Miocene at different localities (open cast mines) of the West-Mariza Brown Coal Basin (von Huene and Nikolov, 1963, probably MN9). Thus, it is clear that the disappearance of the crocodile (*Diplocynodon*) shifted southward (Fig. 3) from Central Europe and France (palaeolatitude 38–45°N, end of MN6, 13.5–14.0 Ma), to Southwest and Southeast Europe (palaeolatitude 30–37°N, end of MN9, about 10 Ma).

Fig. 4 shows the distribution of dry adapted ectothermic vertebrates, as a proxy for seasonality in precipitation, within the Bavarian part of the NAFB. Two time spans (shaded areas on Figs. 2 and 4) display a frequent occurrence of five of the seven selected taxa. The first one is represented by sedimentary cycles 3–5 (OSM 3–5), characterised by the coexistence of *Albanerpeton inexpectatum, Salamandra sansaniensis, Bufo* cf. *viridis, Chamaeleo caroliquarti* and *Gekkonidae*. The second one comprises cycle 9 (OSM 9) with *Salamandra sansaniensis, Bufo* cf. *viridis, Chamaeleo caroliquarti, Varamus hofmanni* and *Bransateryx* sp. The other sedimentary cycles yielded no records of dry adapted taxa (cycles 1, 2, 6) or only one taxon (cycles 7, upper part of 8, 10) or two taxa (cycle 0). This high frequency of dry adapted animals in the cycles 3–5 (16.3 to ~15.7 Ma, base of Early
Badenian) and cycle 9 (14.7 to ~14.5 Ma, Early/Middle Badenian transition) can be explained by an immigration from the northern karst plateau during periods with higher seasonality in precipitation, e.g. longer dry seasons.

4. Discussion and conclusions

The temporal distribution of selected thermophilic ectothermic vertebrates (Fig. 2) shows two subsequent migration events: chamaeleonids and *Gavialosuchus* at 20 Ma and channids, varanids and cordylids at 18 Ma. These are followed by two regional extinction events: cordylids at 16.5–16.7 Ma and chamaeleonids, alligatorids and giant turtles at 14.0–13.5 Ma. Two immigrant groups, the Cordylidae (Auge, 1987) and the Chamaeleonidae (Raxworthy et al., 2002) are of African origin. The Chamaeleonidae entered Europe via an African–Eurasian landbridge (the so-called *Gomphotherium* landbridge), which was established since approximately 20 Ma, leading to important intercontinental faunal exchange (Van der Made, 1997; Rögl, 1998; Agusti et al., 2001). The immigration of chamaeleonids was induced by both climatic (warming of the mid-latitudes) and palaeogeographic changes. In contrast, the cordylids appear later. The remaining immigrants (*Gavialosuchus*, channids, varanids) are of Asian origin. A European–Asian migration corridor, north of the Paratethys, was open since the Eocene/Oligocene boundary (Haq et al., 1987) and no physical barrier existed at that time. Therefore, these migration events within the Early Miocene can be explained by climatic change towards higher temperature, whereas the extinction of *Gavialosuchus* can be correlated with environmental change (regression of the Upper Marine Molasse – most of *Gavialosuchus* remains are to be found in marine and littoral sediments, Antunes, 1994). The regional extinction events of chamaeleonids, alligatorids and giant turtles can be explained by climatic deterioration. Less thermophilic herpetological groups (e.g. Lacertidae, Anguidae) are not affected by the thermal optimum and form an ever-present background (see Appendix 1 in the online version of this paper).

The dotted lines on Fig. 5 show the lower limits of the mean annual temperature (MAT), the minimal warm months temperature (mWMT), and the minimal cold months temperature (mCMT) during the investigated time span. The lines are constructed using the lowest temperatures derived from the extant distribution of the nearest living relatives of the investigated thermophilic taxa (see Table 1). *Gavialosuchus* is disregarded because of the relict distribution of its nearest living relative *Tomistoma schlegeli* (see Fig. 1). I consider the two migration events at 20 Ma and 18 Ma as indicative of a warming trend towards the Miocene Climatic Optimum. The time before the first migration event is characterised by a MAT not below 15.7°C, a mCMT not below ~1.7°C, and a mWMT not below 18.3°C. Thereafter, the lower limit for MAT must have exceeded 17.4°C lasting from 20 Ma to the time interval with the main extinction events between 14.0 and 13.5 Ma. During this warm episode the limit for the mWMT remained stable but the mCMT exceeded 8.0°C. This coincides with the findings of wood and leaf remains with tropical affinities in the Ottnangian
### Karpalian to Middle Badenian

<table>
<thead>
<tr>
<th>Karpalian</th>
<th>Early Badenian</th>
<th>Middle Badenian</th>
<th>Central Paratethys stages</th>
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<tr>
<td>Older Series</td>
<td>Middle Series</td>
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<td>OSM units Böhme et al. 2001</td>
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<tr>
<td>OSM I</td>
<td>OSM J</td>
<td>OSM K</td>
<td>OSM L</td>
</tr>
</tbody>
</table>

- Fine-grained sediment (clay, mud)
- Medium to coarse sand (silts, gravels)
- Pebble to boulder horizon

**Gekkonidae**
- *Chamaeleo caroliquarti*
- *Bufo cf. viridis*
- *Salamandra sansaniensis*
- *Varanus hoffmanni*
- *Brantsateryx sp.*
- *Albanerpeton inexpectatum*
and Karpatian from South Germany and in the Badenian from Romania (Heliconiaceae, Bombacaceae, Euphorbiaceae, Flacourtiaeae, Dipterocarpaceae, Meliaceae, Tiliaceae, Arecaeeae; see Selmeier, 1999; Schweigert, 1992, 1998; Iamandei and Iamandei, 2000). These palaeobotanical proxies indicate a paratropical rainforest with a MAT of 20°C or higher (Schweigert, 1992). The occurrence of bauxites and laterites in the Vogelsberg volcanic area (Schwarz, 1997) provides additional information. The estimated ages for the volcanic rocks vary between 16.2 and 15.5 Ma. Proxy models according to Price et al. (1997) indicate that modern bauxite formation requires a MAT higher than 22°C, six or less dry months (less than 60 mm rainfall) and a rainy season resulting in an annual mean precipitation of more than 1200 mm. This corresponds excellently with the distribution maxima of dry adapted ectotherms (Fig. 4), indicating an increasing seasonality of precipitation in the Badenian between 16.3 and 15.7 Ma (base of Early Badenian) and between 14.7 and 14.5 Ma (Early/Middle Badenian transition).

These periods with high seasonality antedate two main hiatuses within the Upper Freshwater Molasse section, between the sedimentary cycles OSM 5 and 6 and above cycle OSM 10 (Fig. 4). Both hiatuses can be attributed to minor tectonic movements of the NAFB (Reichenbacher et al., 1998). But during the Early Badenian no significant uplift in the Eastern Alps has been documented (Kuhlemann and Kempf, 2002). Thus a direct correlation between periods of higher seasonality in precipitation and Alpine uplift is rather unlikely.

A better explanation for the increased seasonality at the beginning of the Badenian can be inferred from the reorganisation of the Central Paratethys realm. The subduction of the North European platform under the Carpathian–Pannonian block and a back-arc extension associated with diapiric uprise of the asthenospheric mantle led to extensive explosive–effusive volcanism (Vass et al., 1988; Kováč et al., 2000) and to the tectonically-controlled sea-level changes in the Paratethys (Hudáčková et al., 2000). Especially during the Early Badenian, andesitic and rhyodacitic–rhyolitic volcanism culminated in the West Carpathian and Slovakian Basins (Vass et al., 1988) and gave rise to a belt of stratovolcanoes up to 4000 m or even higher palaeoaltitude (J. Lexa, Bratislava, pers. commun., 2001). This belt immediately east of Central Europe at palaeo-latitude 43–44°N most probably influenced the regional atmospheric circulation, especially if the dominant wind direction was from the east as proposed by Utescher and Bruch (2000, fig. 9).

Table 1
Range of the MAT, the mWMT and the mCMT within the extant distribution of selected thermophilic taxa (after Haller-Probst, 1997).

<table>
<thead>
<tr>
<th>Taxon</th>
<th>MAT°C (°C)</th>
<th>mWMT°C (°C)</th>
<th>mCMT°C (°C)</th>
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<tr>
<td>Varanidae</td>
<td>14.8–28.1</td>
<td>13.9–26.1</td>
<td>−3.9–19.4</td>
</tr>
<tr>
<td>Cordylidae</td>
<td>17.0–30.1</td>
<td>15.0–31.0</td>
<td>3.0–23.0</td>
</tr>
<tr>
<td>Geochelone ssp.</td>
<td>17.0–28.7</td>
<td>15.0–26.1</td>
<td>3.0–22.2</td>
</tr>
</tbody>
</table>

Fig. 4. Composite section of the Older and Middle Series of the Upper Freshwater Molasse (Bavarian part of the NAFB; thickness about 150 m) and the distribution of dry adapted ectothermic vertebrates (n refers to the number of studied localities from each cycle). See Appendix 1 in the online version of this paper for locality data. The age of the Ries-boulder horizon (marked by asterix) is 14.87±0.36 Ma (Storzzer et al., 1995), the age of the bentonite of OSM cycle 9 is 14.6±0.8 Ma (Storzzer and Gentner, 1970). The shaded areas indicate periods of higher seasonality in precipitation.
This interpretation implies that moist air masses originating in the Eastern Paratethys probably had no major influence on Central European climate. Coevally, this effect could have interfered with sea-level falls in the West Carpathian–North Pannonian Basins at the Karpatian/Badenian and the Lower Badenian/Middle Badenian boundaries (Hudáčková et al., 2000) which supposedly resulted in a decrease of moist air mass production.

Combining herpetological, palaeobotanical and bauxite formation data, it can be concluded that the Miocene Climatic Optimum in Central Europe ranged from about 18 to 14.0–13.5 Ma (Ottnangian to Early/Middle Badenian) and was characterised by a high MAT exceeding 17.4°C to at least 20–22°C (dashed line in Fig. 5). A period of high precipitation during the Ottnangian and Karpatian was followed by a period with stronger seasonality in precipitation (up to six dry months) in the Lower Badenian.

During the interval from 14.0 to 13.5 Ma, a rapid climatic deterioration occurred. This cooling event, which led to the contemporary regional (Central European) extinction of alligators, chameleons and giant turtles, is characterised by a decrease of the mCMT of more than 11°C and the possible occurrence of frost periods (Fig. 5). The MAT probably dropped at least 7°C to levels between 15.4°C and 14.8°C. This climatic deterioration resulted in a stronger temperature gradient in Europe and marks the beginning of a more distinct climatic zonation on the continent. The thermophilic ectotherms survived only in southern regions (south of 37°N palaeolatitude) as is shown for the crocodile Diplacynodon (Fig. 3). The record of ectothermic vertebrates suggests that the winter temperatures formed the major driving factor in the Miocene thermal evolution of the mid-latitudes, thereby confirming palaeobotanical results from both Europe (Utescher and Bruch, 2000, p. 147) and North America (Wolfe, 1994, p. 201). Additionally, the palaeobotanical investigations in the mid-latitudes of western North America (Wolfe, 1994) show the same trends (increasing MAT between 20 and 18 Ma, decreasing MAT around 13 Ma). The significantly lower values of MAT (15.4–16.6°C) are probably caused by higher palaeoaltitudes in this region.

The results show further good agreement with the record of global deep-sea oxygen isotope ratios (Fig. 2). This record indicates an earliest Miocene cooling event (Mi-1 glaciation at 23.0 Ma), followed by a warming peaking 17–15 Ma in the Mid-Miocene Climatic Optimum. Renewed cooling and reestablishment of the major ice-sheets on Antarctica led to a decline of the deep-sea temperature between 14 and 15 Ma (Zachos et al., 2001, fig. 2). The investigation presented here shows that at least in continental deposits of the mid-latitudes of Europe this cooling event can be dated somewhat later, between 14.0 and 13.5 Ma.

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