Taxon-specific variability of leaf traits in three long-ranging fossil-species of the Paleogene and Neogene: Responses to climate?

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ABSTRACT

Data of climate-sensitive leaf traits, which are usually collected and analyzed for entire fossil leaf assemblages, also include intraspecific responses to environmental conditions. Intraspecific correlations between climate and leaf traits represent plastic responses on the individual level as well as plasticity caused by genetic differences between disjunct populations of a species. Plasticity is taxon-specific, as documented by various studies on extant plants. Data on plasticity in fossil plants are, however, rare. In this study, the plasticity of climate-sensitive leaf traits of three long-ranging species, each covering an extended time interval from the late middle Eocene to the late Oligocene or to even the early Miocene, were tracked by using material from 16 sites located in Austria, Czech Republic and Germany. Selected taxa are Daphnogene cinnamomifolia, Eotrigonobalanus furcinervis and Platanus neptuni. Leaf size-related data (lamina size, length, width) as well as leaf shape-related data (centroid, length-to-width ratio and two parameters for “roundness”) were considered. All three considered fossil-species show various site-specific and significant differences for leaf size-related traits as well as for leaf shape-related traits. Data from allochthonous marine deposits show the highest plasticity, probably due to the accumulation of heterogeneous plant material from different growing sites. For the Oligocene and Miocene, the results are mostly consistent with palaeo-temperature data. This is particularly the case for “roundness” data, confirming the suitability of this trait as an indicator for climate. The high variability of various traits found for the Eocene is, however, difficult to attribute to temperature alone. Rather, the considerable variability of Eocene trait data may be explained by environmental instability during climate transition, such as changing precipitation patterns.
INTRODUCTION

Leaf traits are an important source of palaeoclimate information. A classic and popular leaf trait-climate relationship is the positive correlation between mean annual temperature (MAT) and the frequency of species with toothed margins, which is utilized as a proxy approach known as Leaf Margin Analysis (LMA) (Wilf, 1997; Su et al., 2010). Another approach based on a single leaf trait is the use of fossil leaf size as an indicator for humidity (Wilf et al., 1998; Jacobs and Herendeen, 2004). This is based on the observation that leaf size and precipitation tend to be positively correlated (Givnish, 1984). A widely used multivariate tool, CLAMP, utilizes the correlations between leaf traits and climate parameters (Wolfe, 1995; Spicer et al., 2009; Yang et al., 2015). Recently, some new trait-climate correlations were described. The number and area of leaf teeth was shown to be negatively correlated with MAT, which is part of a quite general response of leaf circularity to climate (Huff et al., 2003; Peppe et al., 2011; McKee et al., 2019). Circularity describes the degree of “roundness”, ranging from 1 (a perfect circle) to 0 (a line), and declines with the degree of leaf dissection, with the latter decreasing significantly with MAT (Peppe et al., 2011).

Data for climate-sensitive leaf traits are usually collected and analyzed for entire leaf assemblages of the local vegetation. The total leaf trait data of a specific site consist not only of the site-specific taxonomic composition, but also of phenotypic variability within the studied taxa. This intra-specific phenotypic variability is caused by plasticity, the ability of organisms to change their phenotype according to environmental conditions. In fact, plasticity is acknowledged as an important factor for the trait-climate relationships revealed by CLAMP data sets (Yang et al., 2015). Royer (2008) and McKee (2019) found for various species that 1) leaves tend to become more dissected when the plants are growing in colder climate, 2) leaf area responds negatively to temperature, while 3) circularity increases with increasing temperature. Intra-specific plasticity can also be caused by genetic differences between disjunct populations of a species, termed ecotypes, representing in many cases adaptations to regional climate (Larcher, 2003). To identify genetic differences between populations, “common garden experiments” are conducted, meaning the cultivation of individuals from different locations at one site. It is then tested whether population-specific trait differences are expressed at the cultivation site, which is taken as evidence of genetical differences between populations (Hufford and Mazer, 2003).

For various leaf traits, plasticity, ecotype evolution and interspecific evolutionary trends appear follow similar directions, as suggested by various data. For example, studies conducted by McKee et al. (2019) and Royer (2008, 2009) showed that plastic responses of leaf dissection and leaf circularity to temperature were largely consistent with the global correlations found by Peppe et al. (2011). Another example is provided by temperate heteroblastic species of Viburnum, in which the sequence of stronger dissected leaves emerging during spring are followed by less dissected sum-
mer leaves. This parallels the evolution of leaf types in different climate zones (Spriggs et al., 2018). Data provided by Mc Kee et al. (2019) for four deciduous woody species (Quercus rubra L., Acer negundo, Carpinus caroliniana and Ostrya virginiana (Mill.) K.Koch) indicate that leaf size responses to temperature are taxon-specific. Plasticity itself is controlled and regulated by genes (Chitwood and Sinha, 2016) and interspecific differences among species are therefore probable. For example, in the case of Quercus kelloggii, no response of leaf traits to temperature could be detected (Royer et al., 2008).

Generally, correlations between climate and leaf traits are considered to be the result of convergent evolution with the leaf traits representing adaptations to environmental conditions. Often, however, the concrete benefits of a single trait have not yet been clarified. For instance, the positive correlation between leaf size and humidity could be explained by water relations, because large leaves require sufficient water availability and small leaves should therefore be advantageous in drier habitats. In fact, however, leaf size is also affected by other environmental parameters, such as temperature (Dolphins and Dilcher, 1980) or light (Scoffoni et al., 2015). On a global scale, leaf size tends to be positively correlated with both temperature and humidity (Wright et al., 2017). The responses of leaf traits to environmental conditions are complex, because individual characters may have more than one function, and many climate parameters are correlated with one another. In fact, it has been shown that leaf traits are often intercorrelated (Yang et al., 2015).

Although intraspecific variability is to be expected to be present in fossil plants, detailed information on intraspecific variability is sparse. Taxon-specific changes in the leaf traits of Rhodomyrtophyllum reticulatum from the middle to late Eocene and Platanus neptuni from the late middle Eocene to early Miocene were reported and discussed recently by Moraweck et al. (2019), considering leaf size and also various micromorphological traits found for leaf cuticles. Besides showing distinct site-specific variances, the results also indicated a possible influence of depositional setting caused by habitat conditions (Moraweck et al., 2019). For various taxa from two Oligocene sites of the Paleogene Basin of Hungary, which show similar age and taxonomic composition, intraspecific differences in leaf size and ratio of leaf length to width could be demonstrated (Tamás and Hably, 2009). For the fossil species Eotrigo- nobalanus furcinervis, which is one of the three study taxa herein, evidence for a significant increase in leaf area between two Eocene sites of almost identical environmental settings was found by Toumoulin et al. (2020).

More information on palaeo-variability is needed because trait-climate correlations are the result of responses on three different levels: 1) habitat preferences of taxa bearing climate-sensitive traits, 2) intraspecific plasticity (ecotypes) and 3) sub-individual phenotypic plasticity. In this study, climate-sensitive leaf traits of three long-ranging species, each covering an extended time interval from the late middle Eocene to the late Oligocene or to even the early Miocene, were tracked. Besides leaf size-related data, such as lamina perimeter, also leaf shape-related data, such as circularity data, were collected. Selected fossil-species are Daphnogene cinnamomifolia, Eotrigo- nobalanus furcinervis and Platanus neptuni. Note that the aims of the present study do not include a contribution to trait-climate correlations, which are utilized in leaf-based tools for reconstructing palaeoclimate, such as Yang et al. (2015). Rather, the aim of this study was to investigate plasticity of fossil species, by addressing the following questions: 1) What is the degree of plasticity in the considered fossil-species, as compared to extant plasticity data? 2) What is the “plasticity behavior” over time (is there a possible evolutionary change)? 3) Are there significant site-specific differences in leaf traits exhibited by the three taxa? 4) Are these differences consistent with palaeoclimate, palaeogeography and stratigraphic trends? 5) Are there any effects detectable which are due to depositional settings?

**TAXA, SITES AND METHODS**

**Taxa**

Platanus neptuni (Ettingsh.) Bůžek, Holý & Kvaček (Platanaceae) occurred under warm-temperate to ‘subtropical’ climates from the late middle Eocene to the early Miocene in central Europe. For NW Romania, fossil material exists for P. neptuni dating back to the late Miocene (Kvaček and Manchester, 2004). It was also recovered from the Oligocene of Oregon (Manchester and Kvaček, 2010). P. neptuni was a thermophilous tree species, originally described as deciduous (Kvaček and Manchester, 2004), quite tolerant with respect to environment, and was able to grow in different habitats (Walther, 1985). P. neptuni thrived in zonal communities and riparian forests, in both mixed
mesophytic forests and evergreen broadleaved forests. For the present study, data for *P. neptuni* considered by Moraweck et al. (2019) were used, together with additionally collected data. 

*Eotrigonobalanus furcinervis* (Rossm.) Walther & Kvaček (Fagaceae) occurred in Europe and western Asia from the middle Eocene to the uppermost Oligocene. Other occurrences of leaves have been reported from the Danian of western Greenland (Grimsson et al., 2016), and from the Oligocene of Greece (Velitzelos et al., 2014). Based on its rich occurrences at many Eocene to Oligocene sites, the supposedly evergreen *E. furcinervis* represented a quite common element both of riparian and swamp forests (azonal settings) and of the zonal evergreen broadleaved vegetation in the Eocene. Later, in the early Oligocene, it probably represented more of an accessory element of the zonal mixed mesophytic forests but still preferred azonal settings such as swamp and riparian forests. In the late Oligocene, the preferred habitats of *E. furcinervis* were either swamp environments (Kvaček and Walther, 1989; Mai and Walter, 1991) or lacustrine settings (Walther, 1999). Mai and Walther (1985) and Mai (1995) concluded that evergreen Fagaceae including *E. furcinervis* were common in late Eocene zonal lowland Fagaceae–Lauraceae forests. Its occurrence appears to be quite independent of facies settings, suggesting a wider range of habitats suitable for *E. furcinervis* (Kriegel, 2001).

*Daphnogeton cinnamomifolia* (Brongn.) Unger (Lauraceae) represents another frequent fossil species with a wide stratigraphic range, recorded from the Paleogene and Neogene vegetation (Kvaček and Walther, 1998). *D. cinnamomifolia* was evergreen, with coriaceous leaves and preferred zonal communities. With respect to taxonomy, *D. cinnamomifolia* represents a special case, compared to *Eotrigonobalanus furcinervis* and *Platanus neptuni*. Although taxonomic identification of fossil leaves is often challenging, this is particularly the case for Lauraceae leaves, and *D. cinnamomifolia* may represent a morphotype which in fact encompasses more than one species (Kvaček and Knobloch, 1967). It was shown that leaf morphometric traits of the extant genus *Cinnamomum*, which includes many species similar in leaf morphology to *D. cinnamomifolia*, are climate-sensitive and little constrained by phylogeny (Tanrattana et al., 2020). Therefore, *D. cinnamomifolia* was considered in this study with the caveat that leaf trait variability may also include some interspecific variability. As will be shown, leaves of *D. cinnamomifolia* occupy a quite clearly delimited realm in the leaf morphospace.

All leaf specimens are included in the database Morphyll (http://www.morphyll.naturkundemuseum-bw.de/index.php), which is publicly accessible and described in Traiser et al. (2018). The data set used for this study can be found in Appendix 1.

**Sites**

Material from 16 fossil sites in central Europe was considered in the study (Table 1, Figure 1), including locations from the Leipzig Embayment and Kleinsaubernitz [described in detail by Moraweck et al. (2019)], Rauenberg (hitherto all situated in Germany), as well as sites from the Bohemian Massif [summarized in Teodoridis and Kvaček (2015)] (please note that the ages of the various sites were determined by different methods, see Table 1; age values which were not obtained radiometrically are approximate). Additionally, the sites Linz and Eferding, Austria, were included. The material belongs to various collections housed in four institutions (Senckenberg Natural History Collections Dresden, National Museum Prague, State Museum of Natural History Stuttgart and Naturhistorisches Museum Wien) (Table 1).

Rauenberg, close to the city of Wiesloch (Germany), is located in the upper Rhine Valley and the Mainz Basin. The Rauenberg site is a former clay pit harboring fossil-rich marine sediments of Rupelian age [described by Maxwell et al. (2016) and Kovar-Eder (2016)]. The fossil-bearing sediments of Kleinsaubernitz are quite close to fossil coastal lowlands like the Leipzig Embayment. They are, however, lacustrine and represent maar sediments containing a quite rich “volcanic” fossil flora from the late Oligocene (Walther, 1999). Various sites close to the city of Linz and Eferding (both Austria) are located in the North Alpine Foreland Basin and contain marine sediments from the Central Paratethys of the lower Miocene (Grunert et al., 2010; Grunert et al., 2015). The fossil floras were described by Kovar-Eder and Berger (1985) and Kovar (1982).

The ages of the sites range from the late middle Eocene to the lower Miocene and therefore cover a number of major global climate transitions recorded by marine data, including the late Eocene cooling with the distinct cooling event at the Eocene-Oligocene Transition (EOT; Hutchinson et al., 2020), a warming trend in the late Oligocene and the Oligocene-Miocene boundary (Pälike et al., 2006; Zachos et al., 2008; Eldrett et al., 2009;
TABLE 1. Summary of the considered sites. Collections: aSenckenberg Natural History Collections Dresden, bNational Museum Prague, cState Museum of Natural History Stuttgart, dNaturhistorisches Museum Wien. Absolute ages are – if not radiometrically determined - estimated from information available for the stratigraphic position of the sites. Abbreviations for dating: RM: radiometric. BS: biostratigraphy. LS: lithostratigraphy. The literature sources for dating and further information on the sites are indicated with superscript letters: 1Krutzsch (2011), 2Bellon et al. (1998), 3Knoebl et al. (1996), 4Rojik (2004), 5Moraweck et al. (2019), 6Standke et al. (2010), 7Krutzsch et al. (1992), 8Grimm et al. (2002), 9Standke (2008), 10Cajz (2000), 11Cajz (1999), 12Kvaček and Walther (2003), 13Teodoridis, and Kvaček (2015), 14Pers. comm. Miroslav Radon, Regional Museum of Teplice, Czech Republic, 15Goth et al. (2003), 16Rupp and Ćorić (2012b), 17Grunert et al. (2015), 17Rupp and Ćorić (2012a). Abbreviations for depositional setting: F: fluviatile. M: marine. V: volcanic. Abbreviations for Taxa: EF: Eotrigonobalanus furcinervis. DC: Daphnogene cinnamomifolia. PN: Platanus neptuni. Numbers in brackets indicate the number of available leaves. MAT and Std: Mean annual temperature and standard deviation, determined by CLAMP. GSP and Std: Growing season precipitation and standard deviation, determined by CLAMP. CLAMP results from Moraweck et al. (2019), with the exception of sites marked with an asterisk, which were taken from Teodoridis and Kvaček (2015). N.a.: data not available.

| Site                        | Site ID | Age*                        | Depositional setting | Leaf material | MAT ± Std | GSP ± Std |
|-----------------------------|---------|-----------------------------|----------------------|---------------|-----------|-----------|
| Profen-Schwerzau 1ua        | 1       | Bartonian 1 40 – 37.8 (BS)  | F                    | PN (7)        | 18.0 ± 2.1| 2106.3 ± 317|
| Kučlínb                     | 2       | Priabonian 2 38.3 ± 0.9 Ma (RM) | V                    | DC (21)  PN (21) | 16.8 ± 1.3*| 1267 ± 497|
| Klausaa                      | 3       | Priabonian 1 37.8 - 34.8 (BS) | F                    | DC (17)       | 18.6 ± 1.3| 700 ± 497 |
| Svatava – Na Pískáčhb        | 4       | Priabonian 3, 4 37.8 – 34 Ma (BS – Stare Sedlo Formation) | F                    | DC (6)         | n.a.         | n.a. |
| Český Chloumekb              | 5       | Priabonian 3, 4 37.8 – 34 Ma (BS – Stare Sedlo Formation) | F                    | DC (10)  EF (20) | n.a. | n.a. |
| Schleenhain-2o-3u             | 6       | Priabonian 1, 5, 6 36.4 – 34.8 (LS, BS) | F                    | EF (18)        | 19.4 ± 2.1| 2200.7 ± 317|
| Schleenhain-HCa              | 7       | Rupelian 1, 5, 6 33.9 – 33.0 (LS, BS) | F                    | PN (12)        | 14.2 ± 1.3| 654.5 ± 497|
| Haselbach-HCa                | 8       | Rupelian 1, 5, 6, 7 33.9 – 32 (LS, BS) | F                    | EF (7)         | 11.6 ± 1.3| n.a. |
| Rauenbergc                   | 9       | Rupelian 8 32.4 – 30.5 (BS)  | M                    | DC (48)  PN (41) | 18.0 ± 1.3| 1323.0 ± 497|
| Seifhennersdorfb             | 10      | Rupelian 2, 9 30.7 ± 0.7 Ma(RM) | V                    | PN (7)         | 10.0 ± 1.3| 849.4 ± 497|
| Markvartice u Děčínab         | 11      | Rupelian 10, 11, 12, 13, 14 30 - 28 (BS, LS, pers. comm.) | V                    | PN (32)        | 11.9 ± 1.3*| 1425.7 ± 497|
| Kleinsaubernitzb             | 12      | Chattian 1, 15 27.5 – 26.5 (BS) | V                    | EF (18)        | 13.4 ± 1.3| 850.4 ± 497|
| Borna-Ost/ Bockwitz-TC (BOB)a | 13      | Chattian 1 25.5 – 24.5 (BS)  | F                    | PN (25)        | 10.1 ± 1.3| 878.4 ± 497|
| Witznitz-WCa                 | 14      | Chattian 1 24.5 – 23 (BS)    | F                    | EF (14)  PN (17) | 13.8 ± 1.3| 791.4 ± 497|
| Eferdingd                    | 15      | Chattian-Aquitanian 27 – 23 Ma (BS) | M                    | DC (11)         | n.a. | n.a. |
| Linzd                        | 16      | Aquitanian 17, 18 23 – 22 (BS) | M                    | PN (16)        | 16.0 ± 1.3| 1147.5 ± 497|
The depositional settings of the considered sites are different and include fluvial, lacustrine ("volcanic") and marine sediments. The lacustrine sediments originate mostly from maar lakes or other lakes, shaped by tectonic and/or volcanic activity, and will be termed "volcanic" throughout the rest of the text (Akhmetiev et al., 2009). All sites and information with respect to dating and depositional setting are summarized in Table 1. To evaluate the results with respect to palaeoclimate, climate data available for the sites were compiled. These comprise various data based on CLAMP and on the Coexistence approach for the considered area (Moraweck et al., 2015, 2019; Teodoridis and Kvaček, 2015; Kunzmann et al., 2016).

**Selected Leaf Traits**

For the fossil leaves, the following parameters were considered: 1) leaf size-related traits: lamina area, perimeter, length, width and 2) leaf shape-related traits: lamina circularity, lamina roundness and lamina centroid. In the present study, effective leaf width was determined by QGIS (see below), based on the "bounding box". The bounding box is a rectangle in which the leaf fits completely, i.e., without any part of the lamina crossing the sides.

Lamina circularity and roundness can be derived as follows. Lamina circularity (LC), also termed shape factor, is defined by ($A_{\text{lamina}}$ = leaf lamina area, and $P$ = leaf perimeter):

$$\text{LC} = \frac{4}{\pi} \cdot \frac{A_{\text{lamina}}}{P^2}$$

Another parameter describing the similarity of the lamina shape with a circle is lamina roundness (LR) (with $D_{\text{cal}}$ = maximum caliper diameter):

$$\text{LR} = \frac{4}{\pi} \cdot \frac{A_{\text{lamina}}}{(D_{\text{cal}})^2}$$

$D_{\text{cal}}$ is calculated as the longest distance between any two points on the lamina outline. The advantage of lamina roundness is that it is less dependent on lamina perimeter, which often rises substantially when small but densely packed leaf teeth are present. Both LC and LR can adopt values between 0 and 1. The smaller the value, the larger is the difference to a circle, and 1 corresponds to a perfect circle.

Lamina centroid (LCE) means the distance (D) of the area-based centre of gravity from the lamina base, expressed relative to leaf length (L), in percent:

$$\text{LCE} = \frac{D \cdot 100}{L}$$

The LCE indicates the "ovateness" of a leaf: ovate leaves show a low value while obovate leaves show a high value. For example, a value of 50% corresponds to a centre of gravity exactly in the middle length of lamina whereas a value of 60% indicates a more obovate shape due to an apical shift of the centre of gravity.

Additionally, the length-to-width ratio (LWR) was included.

$$\text{LWR} = \frac{\text{lamina length}}{\text{lamina width}}$$

**Image Acquisition and Digitalisation**

The fossil leaves were photographed by using a copy stand and a Nikon 5100, or a Canon EOS 6D camera with a Canon 100 macro lens or scanned using a Canon CanoScan 5600F showing a high depth of field. The digital images were then rotated in order to align the main parts of the primary (central) leaf vein vertically by using Adobe Photoshop 10.0, 11.0 and CS5.1 (http://adobe.com). Finally, the images were size calibrated using ImageJ 1.46a (http://rsb.info.nih.gov/ij/). The leaf outlines visible in the images were digitized with QGIS 2.0 - 2.8 (http://qgis.org), an open source geographic information system (GIS). For this purpose, the images of fossil leaves were georeferenced to the uppermost part of the leaf petiole (if preserved in the fossil) which represents the "origin" of all leaf images. The resulting GIS shapefiles
of lamina outlines were used to calculate leaf morphometric parameters by spatial SQL-queries. For this purpose, the shapefiles were imported into an object-relational PostgreSQL 9.1.19 database (http://www.postgresql.org) with PostGIS 2.0.4 (http://postgis.org) extension, which adds support for spatial queries to be run in SQL. Further details are described and discussed in Roth-Nebelsick et al. (2017) and Traiser et al. (2018).

Fossil leaves are often fragmented. Therefore, incompletely preserved leaves were included in the study when reconstruction of the total lamina outline was possible from the preserved leaf area, on the basis of the morphology of the considered species. This was quite feasible in many cases for *Daphnogene cinnamomifolia* and also *Platanus neptuni*, and much more difficult in *Eotrigonobalanus furcinervis*, due to differences in taxon-typical size and shape (see Toumoulin et al., 2020). For detailed explanation and discussion of the applied procedures see Traiser et al. (2018) and Moraweck et al. (2019). The number of suitable leaves per site is provided in Table 1.

**Plasticity Index**

The plasticity index PI, introduced by Valdářes et al. (2000), is a measure of the degree of variability. It is calculated from measured trait data as PI = [(maximum value − minimum value)/maximum value] x 100.

**Leaf Mass per Area**

LM<sub>area</sub>, meaning the ratio between leaf dry weight and lamina area, is an important parameter in leaf economics (Wright et al., 2004). Due to its significant correlation with leaf longevity, it can be utilized to estimate leaf life span of fossil plants (Royer et al., 2007; Harris and Arens, 2016; Lowe et al., 2018). On the basis of petiole width (PW) and lamina area, leaf mass per area (LM<sub>area</sub>) can be determined approximately, according to Royer et al. (2007):

\[
\text{Log}(\text{LM}_{\text{area}}) = a + b \times \text{log}(\text{PW}^2/\text{A}_{\text{lamina}})
\]

with \( a \) and \( b \) as constants (\( a = 3.070, b = 0.382 \)) (Royer et al., 2007).

**Statistics**

For the three taxa, statistical significance of differences in the considered leaf traits was evaluated between sites within age groups. The considered age groups comprise age intervals: late Eocene, early Oligocene, late Oligocene and early Miocene. All statistical procedures were conducted by using SYSTAT 13.1, Systat Software, Inc.

Statistical significance was determined by ANOVA (for more than two groups) or with the t-test (in the case of two groups). Both tests require a normal distribution and equal variances. Normal distribution and equality of variances were tested with the Shapiro-Wilk test and the Levene test for homogeneity. Raw data sets, which did not show a normal distribution were log₁₀-transformed or – if necessary – the Box-Cox transformation was applied (Sokal and Rohlf, 2012). Often, homoscedasticity was markedly improved by data transformation. As PostHoc Test, the Bonferroni test was applied in the case of normal distribution and equal variances. In the case of non-equal variances, the Games-Howell test was selected. In rare cases, a normal distribution could not be achieved, and a distribution-free test, the Kruskal-Wallis test, was then used, with a PostHoc test (Dwass-Steel-Chritchlow-Fligner). For all tests, significance level was at least \( \alpha = 0.05 \).

To evaluate the morphospace of the different taxa with respect to all considered leaf traits, a classical discriminant analysis was conducted for the Eocene and Oligocene. For the Miocene, not enough data were available to conduct a meaningful discriminant analysis.

**RESULTS**

**Leaf Mass per Area**

The material allowed for the approximate calculation of LM<sub>area</sub>, with 64 suitable leaves for *Platanus neptuni* and 56 leaves for *Daphnogene cinnamomifolia*. Based on the equations provided by Royer et al. (2007), LM<sub>area</sub> = 95.5 g/m<sup>2</sup> ± 25.1 for *P. neptuni* and for *D. cinnamomifolia* LM<sub>area</sub> = 104.7 g/m<sup>2</sup> ± 29.6. For *Eotrigonobalanus furcinervis*, only six leaves were available for determining LM<sub>area</sub>. Here, the result was distinctly lower, amounting to 64.3 g/m<sup>2</sup> ± 14.4. All results are significantly different from each other.

Whereas the data confirm the general assumption of *Daphnogene cinnamomifolia* representing an evergreen taxon, the result for *Platanus neptuni* is close to the boundary between deciduousness and evergreeness (87 g/m<sup>2</sup>) as suggested by Royer et al. (2007). A deciduous leaf habit was concluded by Kvaček and Manchester (2004) for *P. neptuni*, based on the position of leaf buds, which are covered by the petioles of the mature leaves. Development of new leaves would
therefore require the shedding of the older leaves. Furthermore, Kvaček and Manchester (2004) argue that mass occurrences of leaves of Platanus neptuni would further support deciduousness. The terms “evergreen” and “deciduous” originally apply to the whole plant, meaning whether the canopy is green throughout the year or leafless during a certain time interval (Kikuzawa and Lechowicz, 2011). Within this context, the term “semi-evergreen” (sometimes also called “semi-deciduous” and “brevideciduous”) describes a situation in which the abscission of old leaves and emergence of new leaves follow immediately one after the other. The plant is therefore continuously green while the leaf longevity is often about one year. From the “leaf perspective”, a leaf longevity of 12 months is usually considered “evergreen” (Givnish, 2002). As already pointed out by Moraweck et al. (2019), it is quite probable that Platanus neptuni represented such a semi-evergreen taxon, a situation which would be perfectly consistent with the conclusions of Kvaček and Manchester (2004). Additionally, reconstructions of palaeoclimate for fossil sites, which contain material of P. neptuni often indicate a rather long growing season (Moraweck et al., 2019), supporting the notion of a semi-evergreen habit.

Evergreenness of Eotrigonobalanus furcinervis was concluded by several authors (e.g., Kvaček and Walther, 1989) based on the supposed botanical affiliation of the genus within the trigonobalanoid clade of Fagaceae, which consists of evergreen extant species only. However, Grimsson et al. (2016) place a group of extinct trigonobalanoids as a separate entity as sister to the extant clade in their phylogenetic analysis of Fagaceae. For E. furcinervis, the data do not support the interpretation of the leaves as evergreen. Rather, the results indicate deciduousness. With six leaves suitable for allowing calculation of LMarea, the data set was, however, limited. The data do not provide a clear-cut answer to the question whether E. furcinervis was deciduous or not, and the topic requires further analysis.

In short, the data for LMarea obtained in this study support the current evidence for foliar habit in Daphnogene cinnamomifolia and Platanus neptuni. In the case of Eotrigonobalanus furcinervis, there are possibly not enough data available. It should, however, be noted that data relating leaf life span and LMarea show a considerable scatter (for example, Enrique et al., 2016), hampering a clear-cut distinction between evergreen leaves and deciduous leaves based on LMarea. Furthermore, there is a certain species-specific increase of LMarea with rising CO2 (about 20% increase on average for 800 ppm compared to the extant level, Poorter et al., 2009). However, LMarea is also affected by various other environmental parameters, such as water availability, temperature and also irradiation (Poorter et al., 2009). Environmental effects on LMarea of fossil leaves are therefore difficult to pinpoint under changing climate conditions.

**Plasticity Index**

Plasticity is highest for data related to leaf size, lamina area, perimeter, length and width (Figure 2). Leaf centroid is much less plastic, while circularity and roundness show medium plasticity (Figure 2). There are, however, considerable sitespecific differences in plasticity. These are partially caused by the number of available leaves per site, because a certain amount of data is necessary to capture trait variability. As indicated by the data set, about 10 leaves can provide a reasonable approximation for plasticity within the fossil material, because from around 10 data points a relatively stable average value is established. The extremely low plasticity in the case of Profen-Schwerzau, for which only Platanus neptuni leaves were available for this study, may at least partly be caused by a low number of available leaves (Figure 2). It is, however, also possible that low PI reflects environmental conditions (see discussion section). From the entire data set (Figure 2), it seems that PI tends to higher values for marine deposits (Figure 2).

**Leaf Size-Related Traits**

Site-specific leaf size data vary particularly strongly for the Eocene for all taxa, with mostly significant differences between sites (Figures 3–4). This high variability in leaf size-related data is distinctly shown by Daphnogene cinnamomifolia, which provided the largest amount of material for the Eocene sites (Figure 3E–F and Figure 4E–F). In Eotrigonobalanus furcinervis, perimeter and length show statistically significant differences for the Priabonian sites, in contrast to lamina area and width, indicating that leaves from the Český Chloumek site are shorter than at the other sites.

Variability of leaf size during the early Oligocene and late Oligocene appears to be much lower for all taxa. Platanus neptuni shows some significant site-specific differences in leaf length and leaf width data from the early Oligocene. For the early Miocene, P. neptuni leaf size data indicate a substantial rise, compared to the Oligocene data,
whereas leaf size of *Daphnogene cinnamomifolia* does not show considerable differences between the Eferding site (early Miocene group) and the early Oligocene data.

**Leaf Shape-Related Traits**

With respect to circularity and roundness (Figure 5), site-specific differences between Eocene sites are all significant for *Eotrigonobalanus furcinervis* and *Daphnogene cinnamomifolia*. Circularity and/or roundness show a trend towards lower values from the Eocene and early Oligocene to the late Oligocene for *Platanus neptuni* (Figure 5A–B). For the early Oligocene, there appears to be no clear trend with respect to circularity and roundness in *P. neptuni* (Figure 5A–B). A statistically significant difference is, however, shown between circularity values of the Chattian sites, followed by a higher value for the Aquitanian (Figure 5A). In *D. cinnamomifolia*, circularity and roundness tend to
lower values from the Eocene to the early Oligocene, although quite low Eocene values do also exist (Figure 5E–F). Also, circularity is low for the Chattian-Aquitanian in *D. cinnamomifolia* (Figure 5E). For *E. furcinervis*, no statistically significant differences could be detected for circularity and roundness data of the Oligocene sites (Figure 5C–D).

Despite low plasticity in LCE, the centroid data also show statistically significant differences for the Eocene (Figure 6A, 6C, 6E) and here also for *Eotrigonobalanus furcinervis* (Figure 6C). For the early Oligocene, LCE data show no significant

**FIGURE 3.** Site-specific data for lamina area and lamina perimeter plotted for each taxon. 3A: Lamina area for *Platanus neptuni*. 3B: Lamina perimeter for *P. neptuni*. 3C: Lamina area for *Eotrigonobalanus furcinervis*. 3D: Lamina perimeter for *E. furcinervis*. 3E: Lamina area for *Daphnogene cinnamomifolia*. 3F: Lamina perimeter for *D. cinnamomifolia*. The boxes span the 50% interquartile. The horizontal lines within the boxes indicate the median values. The “whiskers” mark the highest and lowest values. Outliers located at a distance of up to 1.5 times the quartile span outside the whiskers are drawn as asterisks, and extreme outliers are drawn as circles. Different minuscule letters indicate statistically significant differences among sites. Colors indicate deposit type. Green: fluviatile. Red: volcanic. Blue: marine. Lines delimit age groups. LE: Late Eocene. EO: Early Oligocene. LO: Late Oligocene. EM: Early Miocene. For site numbers and dating see Table 1.
differences for *Platanus neptuni* (Figure 6A) and *Daphnogene cinnamomifolia* (Figure 6E). However, centroid data are significantly different for both late Oligocene sites in *P. neptuni* (Figure 6A). Both *P. neptuni* (Figure 6A) and *D. cinnamomifolia* (Figure 6E) show low values for centroid at the Chattian-Aquitanian and the Aquitanian site. All in all, centroid data of *P. neptuni* appear to show a declining trend with decreasing age, and therefore to less ovate leaves (Figure 6A). However, the range of the centroid values is quite narrow, and therefore the change in leaf shape is minor.

As visible from Figure 5 and Figure 6B, 6D and 6F, and as expected, data in circularity and
roundness tend to be inversely related to LWR data. No relationship could be detected between LCE and circularity and/or roundness (data not shown).

Discriminant analysis (Figure 7) shows that the three taxa form consistent groups which are quite well separated, particularly for the Eocene, despite some overlapping. For the Oligocene, overlapping is stronger, meaning that the considered leaf traits become more similar for the three taxa. The reasons for this result are unclear. Possibly, the niches occupied by the three taxa over-
lapped more during the Oligocene, but this is purely conjectural.

DISCUSSION

Trait Variability and Possible Drivers

Intraspecific variability of morphometric leaf traits can be high, particularly for leaf size-related traits, and PI values found in this study are similar.
to values found for extant taxa (Günthardt-Goerg et al., 2013; Richardson et al., 2013) (Figure 2). Intraspecific variability, expressed as PI, varied substantially between sites, which may be caused by site-specific environmental conditions (Günthardt-Goerg et al., 2013). Data availability does, however, also affect PI because a certain minimum number of data are required to assess PI. Sites with fewer leaves are therefore expected to provide a more limited insight into variability than sites with a higher number of leaves. Despite this caveat, in this study sites with a lower number of leaves also show PI values which are quite similar to modern data. Although PI is quite high for many traits, the trait spaces appear to be confined to taxon-specific morphospaces during the whole considered time frame, as is also indicated by Figures 3–7. Just for leaf size-related data of *Platanus neptuni* data from the Aquitanian are outside the range of the older data sets: *P. neptuni* leaves from the Aquitanian site Linz tend to definitely larger leaves compared to all other sites. However, also in this case, leaf shape does not change, as is indicated by LWR and LCE data. This indicates that size-related parameters, lamina area, perimeter, width and length, change mostly into the same direction, which is also valid for the other two taxa. The site Profen-Schwerzau is exceptional, showing an extremely low PI. It is, however, quite possible that the Profen-Schwerzau PI is not just an artifact due to low data availability, as will be discussed in the last section.

Intraspecific and sub-individual plastic responses to environment are often considered as having an adaptive value itself, because the individual – or the ecotype – is able to respond adequately to immediate circumstances (Dudley, 2004). Large leaves supposedly signal high humidity levels due to their high water demand. However, other factors are also involved. Plants with large leaves tend to occur preferentially not only in more humid habitats, but are also more frequent under warmer conditions. These plants also require sufficient nutrient availability and are also more common under lower irradiation (Givnish, 1984; McDonald et al., 2003; Nicotra et al., 2011; Peppe et al., 2011; Wright et al., 2017). The high scatter of global data, such as that collected by Wright et al. (2017) indicates that the influence of environment on leaf size is even more complex. For example, Traiser et al. (2005) found for data for woody plants of Europe, that large angiosperm leaves are not uncommon at higher latitudes, possibly to facilitate leaf heating during a cool spring to promote photosynthesis in deciduous taxa (Okajima et al., 2012). Additionally, low angles of incident light may promote the occurrence of larger leaves as was suggested for the Paleogene Arctic mixed broadleaved forests (Herman, 1994).

For evergreen leaves, the lamina should be small and xerophytic in climates with winter freezing, because the common formation of substantial amounts of extracellular ice within tissues of frost hardy plants strongly dehydrates living cells (Hacker and Neuner, 2007; Konrad et al., 2019). Also, freezing-induced water stress can occur because it is not possible for plants to absorb water from frozen soils (Tranquilini, 1982). In fact, Reichgelt et al. (2017) interpreted smaller leaf sizes in Miocene floras from New Zealand, compared to
extant vegetation, as possibly being caused by lower temperatures and higher probability for winter freezing.

Intraspecific responses of leaf size to environment are also not clear-cut. For example, temperature-caused changes in leaf size found by McKee et al. (2019) for different extant species were quite heterogeneous. Xu et al. (2012) reported a significant increase in lamina size for experimentally warmed *Betula utilis*, while four extant *Nothofagus* species studied by Richardson et al. (2013) revealed a slightly positive correlation between leaf size and temperature which was, however, statistically not significant. Trait variability in response to environment can be complex because different environmental parameters acting upon one trait can elicit changes in different directions. The combined total response as a result of two and more factors cannot be derived from the isolated responses to each single factor (Valladares et al., 2007). Additionally, leaf traits are often correlated with each other (Yang et al., 2015).

Leaf size, for instance, is affected by a number of environmental factors, with complex outcomes (Dolph and Dilcher, 1980; Fonseca et al., 2000; McDonald et al., 2003; Carlson et al., 2015; Tozer et al., 2015; Wright et al., 2017). For example, a study showed that leaf size of *Fagus sylvatica* in a temperate forest increased with declining precipitation along a humidity transect (Meier and Leuschner, 2008). This result was contrary to expectation, because leaf expansion is strongly negative affected by water stress and leaf size is therefore expected to decline under lower humidity (Bradford and Hsiao, 1982). These observed site-specific differences in leaf size, however, were shown to be driven by conditions in spring when leaves expanded and when humidity was not substantially limited (Meier and Leuschner, 2008). Since the sites with higher summer drought were also warmer during spring, the positive correlation between leaf size and spring temperature was causal while the seemingly negative correlation of leaf size with annual precipitation was a side effect (Meier and Leuschner, 2008). Also, nutrient availability has to be considered. Schneider (2004) reported a lower average leaf size in fossil *Magnolia liblarense* from Miocene lignites in eastern Germany compared to leaves of the same species from fluvial assemblages, suggesting lower nutrient availability as possible factor leading to smaller leaves in the former habitat.

In previous studies, the correlation between circularity and temperature was reported to be positive, when floras from different areas were compared (Huff et al., 2003), and the same was observed for a global data set (Peppe et al., 2011). Circularity was also demonstrated to show a plastic response to temperature (Huff et al., 2003; Royer et al., 2008; Royer et al., 2009; Peppe et al., 2011; McKee et al., 2019). For circularity, possible adaptive benefits are even less clear than for lamina size. For lobed and toothed leaves, the rounding of lobes and leaf teeth obviously contribute substantially to the plastic increase in circularity (Royer et al., 2008; Royer et al., 2009; McKee et al., 2019).

None of the taxa considered in this study is lobed, and one taxon is entire-margined, but all showed significant variation in circularity. Obviously, changes in circularity of the considered taxa are connected to changes in LWR, as shown by Figure 8, which shows the taxon-specific circularity data plotted against LWR. Clearly, an increasing LWR leads to lower circularity (Figure 8). In Figure 8, also the relationship between circularity and length-to-width ratio of an ellipse is included, which is quite closely tracked by the taxon-specific data collected in this study. This is particularly the case for *Daphnogene cinnamomifolia*, whereas the data of *Platanus neptuni* and *Eotrigonobalanus furcinervis* are located somewhat below the ellipse graph. This is very likely due to the leaf teeth of *P. neptuni* and *E. furcinervis* and to their more spindle-shaped lamina, reducing their lamina area relative to an elliptic shape showing the same width.

Occasionally, a high LWR is explained as beneficial against overheating under high temperature (Parkhurst and Loucks, 1972). However, leaf temperature depends on absolute width, not on LWR (Leigh et al., 2016). Nevertheless, LWR is climate-sensitive, although in a complex way. In CLAMP, LWR is used in the form of separate LWR classes, which were found to be particularly common in certain environments, in combination with other traits. For example, long and toothed leaves appear to be characteristic for riparian vegetation (Wolfe, 1993). With respect to spatial patterns within European vegetation, Traiser et al. (2005) observed that short and wide leaves are most common under northeastern continental conditions whereas lengthy and narrow leaves occur preferentially in Mediterranean regions, together with smaller leaf size. LWR is also affected by an acuminate tip: the more acute the tip, the longer the main axis of the leaf. For tropical Africa, Jacobs (1999) observed a positive correlation of LWR with precipitation, due to higher occurrence of acuminate tips under high humidity. Site-specific LWR relationships of differ-
ent floras with climate, which are driven largely by vegetation composition, appear therefore to be discontinuous and influenced differentially by various combinations of factors.

With respect to intraspecific responses, a positive correlation between LWR and temperature appears to be not uncommon. For example, Xu et al. (2012) reported a decrease of LWR under experimental warming in *Betula utilis*. A similar response was observed by Günthardt-Goerg et al. (2013) for experimental warming of various oak species and also for most species considered by McKee et al. (2019). In many cases, however, the response of LWR to warming was moderate and not significant.

**Trait-Variability and Palaeoclimate**

Despite the various caveats discussed above, trends in circularity, LWR and leaf size of *Platanus neptuni* and *Daphnogene cinnamomifolia* obtained in the present study appear to be quite consistent with site-specific CLAMP results (Appendix 2), at least for the Oligocene and early Miocene. For example, the data obtained in this study for circularity – and also leaf size – corroborate the notion of the Rupelian site Rauenberg as a quite warm locality, and the late Chattian site Witznitz-TC as warmer than the early Chattian Borna-Ost-Bockwitz-TC site, as was also discussed by Kovar-Eder (2016) and Moraweck et al. (2019). For the early Miocene, warmer conditions than for the Chattian sites are indicated by leaf size and circularity data of *P. neptuni* for the Linz localities, also in accordance with CLAMP. For the younger Eferding site (for which no CLAMP results are available), however, *D. cinnamomifolia* shows circularity values similar to the Chattian sites.

It should be noted that comparing stratigraphic trends, as observed in the present study, with CLAMP climate results does not represent a “test” of CLAMP. Firstly, this is not possible with just three taxa. Secondly, independent climate data would be necessary. What this comparison – illustrated in Appendix 2 – shows, however, is that the plastic changes of our taxa are largely consistent with available climate data for the Oligocene and Miocene, but in a species-specific manner, as expected.

Climate was probably quite variable during the Oligocene, with various minor cooling and warming events (Li et al., 2018). For Central Europe, there is evidence for a warmer Rupelian and cooling during the Chattian (Li et al., 2018). Evidence for a warming event at the Rupelian-Chattian boundary in the southern North Sea Basin was found by De Man and Van Simaeys (2004), based on marine
proxies, which also indicate a cooling during the later Chattian. Pollen based data and δ^{18}O values of fossil bivalve shells and shark teeth from Western Germany also indicate a decrease of terrestrial and sea-surface temperatures during the Chattian, followed by warming towards the Oligocene-Miocene transition (Walliser et al., 2016).

Palaeoclimate data for the Paleogene and Neogene are, however, heterogeneous and vary geographically as demonstrated, for example, by data of Teodoridis and Kvaček (2015) for Central Europe, which are based on CLAMP and CA and indicate, all in all, a possible warming after the EOT during the early Oligocene, followed by cooling towards the Oligocene-Miocene boundary. Site-specific climate can also be strongly affected by geographic position and local conditions, and distinct spatial heterogeneity is to be expected reflecting meso-climate. For instance, distance to shorelines will have an effect, due to the marine influence (Li et al., 2018). The region of Linz, for example, was influenced by the warm Paratethys (Kvaček et al., 2006) and Rauenberg was situated at the margin of the Upper Rhine graben, which was a marine connection between the Paratethys and the North Sea whereas the locations of the Leipzig Embayment were exposed to the influence of the cooler North Sea (Moraweck et al., 2019). In particular, the lower Oligocene Rauenberg flora contains some subtropical Eocene relics (Actinodaphne pseudogermari, Doliostrobus taxifloris) and thermophilous species (Ceratozamia floersheimensis) that either disappeared, or were never present, in coeval florals of the coastal North Sea lowlands of the Leipzig Embayment (Kovar-Eder et al., 2008; and Moraweck et al., 2019). Despite these limitations and caveats, however, the presented data for circularity, as well as for leaf size and LWR, are partially in quite good agreement with site-specific CLAMP data as well as with general proxy-indicated trends for Oligocene climate, such as a quite warm Rupelian, and cooler conditions in the Chattian.

**High Trait Variability for the Eocene**

The most conspicuous feature of the data set, however, is the strong variability of morphometric data for the Eocene, particularly shown by *Daphnogene cinnamomifolia*, but also indicated by the more limited Eocene data of *Platanus neptuni*. Statistically significant differences are most frequent for late Eocene material in all considered taxa. For example, in *D. cinnamomifolia*, which provided material for four late Eocene sites, leaf size peaked distinctly for two Priabonian data sets whereas leaf size for the two other localities is markedly lower (Figure 3F). In many cases of the collected data set, the late Eocene variability of the considered leaf traits largely covers the total range of variability over the entire considered time interval.

The values for leaf size-related data of *Platanus neptuni* at the Profen-Schwerzau site are conspicuously low. It is possible that this result is due to the quite limited number of available suitable leaves from this site. With respect to climate, MAT is in the range of the other Eocene sites, whereas precipitations is high (Appendix 2). High precipitation, however, causes leaching of nutrients (Givnish, 2002) which in turn may promote smaller leaf size due to lower nutrient availability. Furthermore, another conspicuous feature of the Profen-Schwerzau site is the very low PI, the lowest for all other sites and taxa (Figure 2). As shown by Balaguer et al. (2001), homogeneous light conditions, meaning more open landscapes, can lead to low plasticity. Furthermore, leaves acclimated to high irradiation may tend to smaller lamina size (Scoffoni et al., 2015). It is therefore possible, that specific local conditions at the Profen-Schwerzau site caused low leaf size and low intraspecific variability.

The late Eocene was a period of global cooling, which is also broadly reflected by plant proxies and vegetational composition. The turnover of vegetation in Central Europe from broadleaved evergreen forests towards mixed mesophytic forests started well before the EOT during the late Priabonian (Kvaček et al., 2014; Teodoridis and
Kvaček, 2015; Utescher et al., 2015; Kunzmann et al., 2016; Pound and Salzmann, 2017). Cooling predominantly increased temperature seasonality (Mosbrugger et al., 2005; Eldrett et al., 2009). The strong morphometric variability shown by the trait data of the considered species for the late Eocene material appears to be not consistent with available CLAMP temperature proxy data, which show mostly moderate variations (Appendix 2). Moreover, trends in lamina area, circularity and LWR data for the late Eocene data appear to be in contradiction to temperature trends indicated by CLAMP (Appendix 2). However, precipitation data indicated by CLAMP results – when available – show a quite high variation for the considered Eocene sites (Appendix 2). There is evidence that in addition to cooling the pattern and amount of precipitation changed in Central Europe, with precipitation becoming more seasonal as also indicated by the paleoclimate reconstruction results (Moraweck et al., 2015). This is also indicated by sediment data from northwestern central Europe, featuring changes in mineral composition as well as content of N-isotopes (Tramoy et al., 2016). These data provide evidence for a transition to drier conditions from the EOT onwards and also a distinct event, namely a dip towards cooler and/or drier conditions during the Priabonian (Tramoy et al., 2016). Possibly, also other soil conditions than drier conditions during the Priabonian (Tramoy et al., 2016). Possibly, also other soil conditions than drier conditions during this time, adding another confounding factor because many leaf traits are affected by nutrients.

From these considerations, it appears to be quite possible that the high morphometric variability shown by two of the three considered taxa for the late Eocene data indicate a general environmental instability during this period of major climate transition. Changing CO₂ values may have also possibly affected leaf traits indirectly, because stomatal conductance, and therefore water demand, is expected to be inversely correlated with atmospheric CO₂ as shown by various data and modeling results (Ainsworth and Rogers, 2007; Konrad et al., 2008; Leakey et al., 2009). In fact, elevated CO₂ was demonstrated to decrease the water demand of a mature forest (Bader et al., 2013), albeit the detailed ecophysiological ramifications of changing CO₂ on plants in natural habitats are complex (Donohue et al., 2017).

With respect to deposit type, the data do not allow for drawing definite conclusions on a possibly specific habitat signal. In Moraweck et al. (2019), leaf size-related data indicated a possible trend for larger leaves in marine deposits, likely caused by the accumulation of leaves of different habitats in these allochthonous assemblages and/or the marine influence on climate. This effect on morphometric leaf traits is much less distinct in the present data set. However, PI appears to tend to higher values in marine deposits (Figure 2). Also, Moraweck et al. (2019) found – for micromorphological traits (stomata, trichome densities) of Platanus neptuni – evidence for higher intraspecific variability in allochthonous assemblages. This may be explained by the heterogeneous composition of leaf material, originating from different sites from the hinterland with different local climate as well as possibly mixing leaves from different ecotypes. However, more data are required to better pinpoint this hypothetical effect.

CONCLUSIONS

All three considered fossil-species show various site-specific and significant differences, for leaf size-related traits, as well as for leaf shape-related traits. For the considered time interval, the data vary mostly within a taxon-specific range, and therefore within a taxon-specific morphospace. There is no indication for trends with respect to deposit type and therefore no specific deposit-habitat relationships. However, data from marine and therefore allochthonous deposits show the highest plasticity, probably due to the accumulation of heterogeneous plant material from different growing sites.

For the Oligocene and Miocene, the results are mostly consistent with palaeo-temperature data and here in particular for circularity data. Since circularity and length/width ratios are closely related for non-lobed taxa, as were considered in this study, the results confirm both traits as useful indicators for temperature. The high variability of various traits for the Eocene are, however, difficult to attribute to temperature. Rather, the considerable variability of Eocene trait data may be explained by environmental instability during the climate transition phase, such as changing precipitation patterns.

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APPENDIX 1.

All data used for the present study. Data for Appendix 1 and 2 is in zipped file for download at https://palaeo-electronica.org/content/2021/3285-taxon-specific-variability.

APPENDIX 2.

CLAMP results for mean annual temperature (MAT) and growing season precipitation (GSP), for the considered sites (site numbers see Table 1). Data for Appendix 1 and 2 is in zipped file for download at https://palaeo-electronica.org/content/2021/3285-taxon-specific-variability.