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Net diversification rates of the woody plant genus *Petalidium* (Acanthaceae) are highest in the ancient and arid Namib Desert

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At present, tropical arid biomes house less woody plant species diversity than tropical moist biomes, which could be due to lower rates of evolutionary diversification in the recent or distant past. Here, we study the evolutionary diversification of Petalidium (Acanthaceae), a genus of 36 species of woody shrubs found in the Namib Desert of southwest Africa, and surrounding areas. We generated a new, nearly fully sampled and temporally calibrated phylogeny for Petalidium using RADseq SNP data and secondary calibrations. We then investigated variation in net diversification rate across the phylogeny, the ancestral climatic niche of lineages and the link between the two. We find that arid climatic conditions are linked with increased rates of net species diversification in the genus. Despite its great age, the Namib Desert clearly hosts young plant radiations. This apparent contradiction can be explained by a scenario of high evolutionary turnover, in this case potentially caused by alternating hyper-arid and relatively mesic phases. Hyper-arid phases could result in high plant mortality and extinction of species, leading to ecological opportunity and diversification during mesic phases. Taken together, our results contribute to a growing body of literature that shows evidence for elevated rates of plant diversification in the Quaternary in arid biomes across the globe.

KEYWORDS

Namib Desert, arid biomes, Acanthaceae, plant diversification, RAD sequencing

Introduction

Within the tropics, the diversity of plant species is generally linked to water availability, with more moist biomes housing higher taxonomic diversity, particularly for woody plants (O'Brien et al., 2000; Daru et al., 2016; Esquivel-Muelbert et al., 2017). Within individual woody plant clades this pattern is also evident, with families and genera tending to be more species-rich in moist biomes compared to more arid biomes (Currie, 1991; Pennington et al., 2006). Assuming no systematic difference in the age of clades, these discrepancies could suggest that diversification rates are lower in arid versus moist biomes.

In contrast, scientists since Stebbins (1952) have argued that arid environments promote plant speciation (Axelrod, 1972; Böhnert et al., 2022). Arid biomes may house a greater diversity of edaphic conditions that can facilitate plant speciation (Stebbins, 1952; Axelrod, 1972; Buira et al., 2021), and the patchy distribution of species in arid biomes may be conducive to intermediate levels of gene flow among populations that can promote adaptive diversity and differentiation (Wright, 1940; Stebbins, 1952).

Present-day arid biomes have a dynamic climatological past, and increasing aridity since the Miocene across the globe has been cited as a causal factor in diversification of arid-affiliated plant lineages (Wu et al., 2018; Böhnert et al., 2022; Čalasan et al., 2022; Hua et al., 2022). Global aridification continued to increase throughout the Pliocene and Pleistocene climate cycles, with multiple examples of Quaternary diversification in arid-affiliated plant lineages (Kadereit and Abbott, 2021). There is ample evidence that arid biomes are areas of active and recent plant speciation. However, most studied examples pertain to herbaceous and succulent plant lineages (e.g., Klak et al., 2004; Arakaki et al., 2011; Guerrero et al., 2013; Böhnert et al., 2019; Lavor et al., 2019; Merklinger et al., 2021). Macroevolutionary studies of woody plant lineages, which show the most marked differences in diversity between arid and moist biomes, are scarce (although see Simon et al., 2009; Singhal et al., 2021). We do not yet have a firm idea of the relationship between arid climatic conditions and diversification in woody plant lineages.

Sub-Saharan Africa as a whole is dry in comparison to other regions of the tropics, such as the Americas and southeast Asia. Yet, arid regions of sub-Saharan Africa have received limited macroevolutionary study compared to regions like the Atacama Desert (Guerrero et al., 2013; Böhnert et al., 2019; Merklinger et al., 2021). The Namib Desert, in southwest Africa, is one of the oldest deserts on Earth, originating 55-80 Ma (Ward et al., 1983). It houses ancient iconic plant lineages such the morphologically bizarre gymnosperm Welwitschia as (Welwitschiaceae) and the similarly bizarre angiosperm lineage Tiganophyton (Tiganophytaceae, Swanepoel et al., 2020). These speciespoor lineages are notable for occurring on long, isolated evolutionary branches, which have been termed "depauperons" (Donoghue and Sanderson, 2015). In contrast, other lineages emblematic of the Namib Desert have achieved markedly high taxonomic diversity, such as Euphorbia (Euphorbiaceae), Indigofera (Fabaceae), Commiphora (Burseraceae) and several genera of Acanthaceae (Craven, 2009; Darbyshire et al., 2020). Most of these clades are diverse across tropical Africa, and the species present in the hyper-arid region of southwest Africa are derived from multiple lineages.

One genus of Acanthaceae, *Petalidium*, stands out from other diverse genera in the region in that its diversity is centred in the Namib Desert and surrounding areas, and is hypothesized to derive from a single colonisation event (Tripp et al., 2017). *Petalidium* is furthermore a lineage of woody, non-succulent plants, a habit usually associated with higher diversity in moist regions. In the present study, we aim to test whether the arid conditions in the Namib Desert may have helped to spur diversification in the genus *Petalidium*, as proposed by Tripp et al. (2017). If supported, *Petalidium* would represent a notable counter-example to the assumption that tropical arid biomes inhibit woody plant diversification.

Petalidium is a genus of 36 accepted species that occur almost entirely in Namibia and Angola, with the exception of one species sister to the African taxa, which occurs in India and Nepal in a more moist climate and two species restricted to northeastern South Africa and adjacent areas in Zimbabwe (Tripp et al., 2017). *Petalidium* is the fifth most diverse genus in the arid Kaokoveld Region of northwestern Namibia and southwestern Angola (Craven, 2009) and is ecologically successful, being the dominant shrub cover in major portions of this region and also forming large populations of co-occurring species in many areas (all authors, pers. obs.). The total taxonomic diversity of *Petalidium* is likely higher than currently known as new species of *Petalidium* continue to be found after fieldwork in poorly studied regions (Swanepoel, 2020; Swanepoel and Manzitto-Tripp, 2022). African *Petalidium* appear to be undergoing an evolutionary radiation, with estimates for the crown age of the 35 African species ranging from 1.4 Ma to 3.6 Ma (Tripp et al., 2017). Species of *Petalidium* can occur in extremely arid areas where mean annual precipitation (MAP) is less than 50 mm per year, but other species of African *Petalidium* occur in more moist areas where MAP exceeds 700 mm. We know that at present the majority of species of *Petalidium* occur in more arid regions (Tripp et al., 2017), but the climatic history of the clade has not been studied. We do not know whether it originated in more moist or arid conditions. If it originated in more arid conditions, then its greater diversity in arid regions may simply be a result of a longer history of occupancy.

In order to examine the link between climatic conditions and diversification in *Petalidium*, we undertook the following steps: (i) we generated a new phylogeny for the genus using RAD sequencing, improving the species sampling from previous efforts by 50% (Tripp et al., 2017); (ii) we assessed variation in net diversification rate across the phylogeny; (iii) we reconstructed ancestral climatic niches across the phylogeny; and (iv) we tested for a link between net diversification and estimated ancestral climatic niche. Our overall aim was to test whether *Petalidium* shows higher species richness in arid climatic conditions relative to moist ones because of higher rates of net diversification in arid conditions or because of a longer history of occupancy of arid climatic environments, with equivalent or even lower net diversification in arid conditions.

Materials and methods

Ecological analyses

We compiled 1,349 occurrence records of *Petalidium* from herbarium collections. We manually checked this database, verified specimens' identifications and removed those that could not be confidently identified or geo-referenced. We then extracted values for the 19 Worldclim bioclimatic variables (Fick and Hijmans, 2017) for remaining occurrences. We computed species means for all variables and performed a principal component analysis (PCA) on mean environmental preference using the princomp function (on the correlation matrix) in the base package of the R Statistical Environment (R Core Team, 2013).

Field methods

Populations of *Petalidium* were sampled for herbarium and silica-gel vouchers during four expeditions to Namibia, one to South Africa and one to Angola, by E.A.T and K.G.D. from 2010 to 2017. Specimens are housed at the following herbaria: RSA-POM, COLO, WIND, LUBA, K, and E. All specimens included in this phylogenetic work were fertile, and we are confident in the identifications.

DNA extraction and sequencing

For 35 samples, we extracted DNA from silica-dried leaf tissue using a standard CTAB protocol (Doyle and Doyle, 1987). DNA was then sequenced using a ddRADseq approach (see Tripp et al., 2017 for protocols including custom barcode design, which was modified from Parchman et al., 2012). We used *EcoR1* and *Mse1* for double digestion followed by size selection of 250–500 bp fragments to reduce the overall genomic material that was sequenced. Libraries were pooled and then submitted to the University of Colorado's BioFrontiers Sequencing Facility for quality control, cleanup, and 1×100 sequencing on an Illumina NextSeq.

Bioinformatics

All bioinformatic work was conducted on the University of Colorado's SUMMIT supercomputer, which comprised a total of 450 compute nodes, each containing 24 cores, for a total of 10,800 available cores (each with 200 GB of RAM). We used FastQC (Andrews, 2017) to assess overall quality of our sequencing. Subsequently, the data were demultiplexed using ea. fastq-multx v.1.03, which is distributed as part of the ea-utils package (Aronesty, 2011). Then, adapters were removed and reads were cleaned using cutadapt v.1.4.2 (Martin, 2011). The resulting fastq files were assessed for total quantity of reads containing the CAATT cutsite in Unix, and any samples with fewer than 100,000 reads with the cutsite were omitted from further consideration. We used ipyrad (Eaton and Overcast, 2020) to trim low-quality bases, filter reads and then assemble the data into loci by implementing reference-guided methods, leveraging a fully assembled, although not yet published nuclear genome for Petalidium crispum (Manzitto-Tripp et al., unpub. data). The following parameters were implemented in ipyrad: max_low_ qual_bases = 10, mindepth_majrule = 2; cust_threshold = 0.85; max_Ns_ consens=5; max_Hs_consens=25; min_samples_locus=25; max_ Indels_locus=100; and max_shared_Hs_locus=0.5. Paralogous loci were excluded from consideration using the max_shared_Hs_locus=2 flag. The final RADseq phylogenomic dataset is available on GenBank under the Sequence Read Archive Study SUB12441993.

Phylogenetic reconstruction and divergence time estimation

We used the matrix of SNPs to infer phylogenetic relationships under the GTR model of substitution using maximum likelihood (ML) in RaxML (Stamatakis, 2014) with 100 bootstrap replicates. We included SNPs that were missing data for up to 90% of the accessions in the analysis. We explored lower thresholds of missing data, but found that the 90% level gave the strongest bootstrap support values (as in Tripp et al., 2017). To avoid bias in branch length estimation due to the absence of constant sites, we applied the ascertainment bias correction to likelihood calculations (Leaché et al., 2015). We also ran a phylogenetic analysis in IQ-TREE multicore version 2.0.3 (Nguyen et al., 2015) under the GTR model of substitution and the ascertainment bias correction, with 1,000 ultrafast bootstrap replicates (Hoang et al., 2018). Missing data in our matrix of SNPs precluded the use of SNAPP, a Bayesian method implemented in BEAST2 that can simultaneously infer tree topology and node ages (Bouckaert et al., 2014). Instead, we dated the ML phylogenetic tree to absolute time using penalized-likelihood, implemented in treePL (Smith and O'Meara, 2012). We set a secondary calibration using an interval of 1.3-5 Ma corresponding to the 95%HPD of the crown age of Petalidium estimated by Tripp et al. (2017). We used crossvalidation to estimate the best value of the smoothing parameter. There were two species (*P. ohopohense* and *P. lanatum*) for which sequence data were available from earlier RADseq runs, but with insufficient sequence data to place in the phylogeny in our complete phylogenetic analysis. Preliminary analyses with different sets of data allowed us to develop a robust idea of whom their closest relative(s) were. These two species were grafted into the final phylogeny mid-way along the branch subtending their closest relative(s) in R using the function bind.tip from the package phytools 1.2.0 (Revell, 2012).

Estimation of ancestral character states and diversification rate

We mapped the logarithm of precipitation of driest quarter along the phylogeny and conducted ancestral state estimation using the function contmap of the package phytools v1.2.0 (Revell, 2012), which in turn relies on the approach of Felsenstein (1985). Of the readily available bioclimatic variables, this one best mechanistically quantifies water availability to plants when it is most scarce, across the driest 3 months of the year. Precipitation of the driest month may be influenced by lags in depletion of water from preceding months, while precipitation in wetter periods may not accurately measure stress from low water availability.

We estimated the net diversification rate of *Petalidium* in a bayesian framework using the software BAMM v.2.5.0. (Rabosky, 2014). BAMM was run with a global sampling fraction of 35/36 and for 10 million generations. Convergence of the mcmc chains was checked in R using the coda package v0.19.4 (Plummer et al., 2008) and plots were made with the BAMMtools package v2.1.10 (Rabosky et al., 2014).

We tested for an association between aridity and net diversification rate using the Quantitative State Speciation and Extinction model (QuaSSE, FitzJohn, 2010), implemented in the R package diversitree v0.9.16 (FitzJohn, 2012). We ran QuaSSE on the first principal component of the climate space occupied by species, which broadly represents a gradient in water availability, as well as on the logarithm of precipitation of the driest quarter. We compared a null model of trait-independent diversification vs. models where the diversification rate has a linear, sigmoidal or hump-shaped response to the environmental variable, with and without drift included. Because trait-dependent diversification models have been shown to have an elevated type-I error rate (Rabosky and Goldberg, 2015), we conducted parametric bootstrapping to evaluate the significance of our results. First, we inferred the root state of the genus for the two variables using the function fastANC from the phytools package v1.2.0 (Revell, 2012). We then used the estimated root state and the estimate of the diffusion parameter from the null model in QuaSSE to perform 100 simulations of Brownian-motion trait evolution along the phylogeny, using the function sim.char from the geiger R package v2.0.10 (Harmon et al., 2007; Pennell et al., 2014). Finally, these 100 simulated trait histories were subjected to a QuaSSE model comparison as with the observed empirical data set. The log-likelihood of the best model minus the log likelihood of the null model (hereafter ΔLL) was recorded for each simulated dataset and compared to the observed Δ LL. The proportion of simulated Δ LL values that were greater than the observed ΔLL value was then calculated to assess the robustness of QuaSSE results. If less than 0.05 of the simulated values are greater than the observed value, then the observed value is considered to be significantly different from that expected by chance in this 1-tailed test.

Results

Ecological analyses

After cleaning the occurrence data, we obtained 982 occurrence records for 35 species. The first two principal components (PC1 and PC2) from a principal component analysis of the mean climatic variables for each species explained 48.1 and 27.8% of the variation in the climate data respectively, with subsequent axes explaining little variation (Figure 1). PC1 was related primarily to amounts of

precipitation (total annual precipitation and total precipitation for different quarters and months), while PC2 was related primarily with mean temperature variables (annual, quarterly and monthly).

Sequencing and bioinformatics

After sequencing and bioinformatic processing, we obtained 83,547 SNPs. Prior to phylogenetic inference with an ascertainment bias correction, only true SNPs (i.e., sites with two bases present across all individuals) were retained, while all sites with one base or an IUPAC



FIGURE 1

Position of species in a principal component analysis of the mean value for each species for 19 bioclimatic variables (from www.worldclim.org). The first two components are shown, which explain 48.1 and 27.8% of the variation in the data, respectively. Species from the "diverse and arid clade" which shows a faster rate of net diversification (see Figure 4) are denoted in red, and all other species are denoted in blue. The length of the arrow for each climatic variable is related to the strength of its correlation with the first two principal components. Climatic variables are abbreviated: Mean_Diurnal_Range=mean diurnal temperature range; T_Ann_Range=temperature annual range; T_Season=temperature seasonality; Max_T_ Warmest_M=maximum temperature of the warmest month; P_Driest_M=total precipitation of the driest month; P_Driest_Q=total precipitation of the coldest quarter; Mean_T_Warmest_Q=mean temperature of the warmest quarter; Ann_P=total annual precipitation; P_Wettest_Q=mean temperature of the wettest quarter; P_Wettest_M=mean temperature of the wettest quarter; Mean_T_Coldest_Q=mean temperature of the coldest quarter; Mean_T=Coldest_M=mean temperature of the coldest month; P_Season=precipitation seasonality; and lsothermality. We interpret the first principal component as a precipitation-dominated axis, with positive values being hotter and negative values being colder. We interpret the second principal component as a precipitation-dominated axis, with positive values being drier and negative values being wetter.

character, which are treated as invariants by many softwares including RaxML, were removed, resulting in a final matrix of 34,017 SNPs for 35 species.

Phylogenetics, ancestral state reconstruction and diversification analyses

RaxML (Figure 2) and IQ-TREE (Supplementary Figure S1) analyses yielded highly similar phylogenetic trees with similar support values. Deep nodes in the ML phylogenetic tree were fully resolved, but bootstrap support values were low within one clade. That clade holds the majority of *Petalidium* species, yet has the shortest branch lengths (Figure 2). Our temporally calibrated phylogeny had an estimated crown age for African *Petalidium* of 4.33 Ma. The clade with short branch lengths and most *Petalidium* species had an estimated crown age of 1.63 Ma (Figure 3).

Ancestral state reconstruction of precipitation of the driest quarter suggests that *Petalidium* originated in more moist conditions and adapted to high levels of aridity in its recent evolutionary history (Figure 4). There are two lineages found within arid areas: one comprises two species (*P. luteo-album* and *P. giessii*). The other is the clade mentioned above with short branch lengths and comprising the majority of species in the genus. The first branch in this clade belongs to *P. rupestre*, a local endemic in the arid Namib Desert of southwest Angola, while the second branch belongs to *P. aromaticum*, a species occurring in more moist conditions along the Limpopo River on the border of South Africa and Zimbabwe. The remaining species in this clade occur in the arid Namib Desert and Kaokoveld. We hereafter refer to this clade as the *diverse arid clade*.

In the BAMM analysis, the best model was one with one shift in diversification rate (posterior probability=0.6), which is placed at the base of the *diverse arid clade* (Figure 4). This clade shows a higher net diversification rate than the rest of the phylogeny. When visualising model-averaged variation in net diversification across the phylogeny, we find that estimated net diversification rate ranged from 0.6 to 1.5 species per myrs.

With our QuaSSE analysis we found that diversification rate in *Petalidium* was best modelled as linearly related to PC1 (the "precipitation" component), with positive values (reflecting lower levels of precipitation, and presumably water availability) being related to higher net-diversification. Net diversification was negatively related to precipitation of the driest quarter, with highest net diversification in the most dry conditions. However, our simulations show that these relationships are not significantly different from what would be expected by chance. Indeed, the simulated Δ LL was greater than observed Δ LL 8% of the time for PC1 (p = 0.08) and 44% of the time for precipitation of driest quarter (p = 0.44). Overall, these results indicate that while there is a link between water availability and net diversification rate, with higher net diversification in arid regions, this link is not significant in a statistical sense.

Discussion

In this study, we found that the higher species richness of *Petalidium* in arid areas relative to moist areas may indeed be linked to higher rates of net diversification in arid areas, in contrast to the alternative hypothesis of a longer history of occupancy in arid areas. Our analyses suggest that the genus as a whole originated in more moist conditions





ca 5 Ma and that two lineages subsequently shifted to occurring in more arid conditions ca 1.6 Ma (Figure 4). One lineage in particular, the *diverse arid clade*, showed elevated rates of net diversification, and this clade occupies the most arid areas within the overall range of the genus. Yet, the singular nature of the *diverse arid clade* prevents us from making strong causal links between occupation of arid areas and net diversification rate within the genus *Petalidium*. However, combining the results here for *Petalidium* with published results for other clades suggests there indeed may be a causal link (Fisher et al., 2015; Darbyshire et al., 2020).

An evolutionary pattern of higher diversification rates in arid areas during the Pleistocene has been found in other lineages of Acanthaceae in the Namib Desert and surrounding regions (Fisher et al., 2015; Darbyshire et al., 2020). Similarly, the majority of the species-rich lineages in the neighbouring Succulent Karoo arid biome are younger than 10 Ma (Klak et al., 2004; Verboom et al., 2009; García-Aloy et al., 2017). Thus, *Petalidium* represents an additional data point, notably from a woody plant clade, supporting a growing body of evidence for rapid Quaternary diversifications of plants in the Namib Desert and surrounding regions. It is key to note though that the Namib Desert also houses incredibly old evolutionary lineages that are found nowhere else on earth, such as *Welwitschia*, which diverged from *Gnetum* in the Cretaceous (Hou et al., 2015) and *Tiganophyton*, an early diverging member of the Brassicaceae which originated in the Eocene (Hendriks et al., 2022).

How then do we explain the co-occurrence of recently radiated lineages and ancient depauperons (Donoghue and Sanderson, 2015) in the same space? We hypothesize that the answer lies in exceptional evolutionary turnover. Recent studies (Koenen et al., 2015; Pennington et al., 2015) have suggested that high diversity in tropical rain forests can be explained by the persistence of numerous evolutionary lineages through episodic periods of extinction, which then provide ample 'seed lineages' from which radiations can arise. If episodes of extinction in arid biomes are more severe, then fewer lineages would



FIGURE 4

Left: Ancestral state reconstruction of precipitation of driest quarter for the genus *Petalidium* (Revell, 2012). Most species of *Petalidium* occur in arid conditions where they receive no precipitation for at least 3 months during the driest quarter, yet the analysis suggests that the ancestor and early lineages of *Petalidium* (i.e., older nodes) occurred in more moist conditions. Thus, occupation of arid conditions appears to be a derived state. Right: Variation in net rate of diversification across *Petalidium* (Rabosky, 2014). This analysis suggests that one clade of *Petalidium*, that which encompasses the majority of species, is diversifying at a faster rate than the other clades in the phylogeny (with red, orange, and yellow branches). This clade also tends to occur in more arid conditions, and is referred to as the *diverse arid clade* in the main text.

persist throughout these episodes and fewer would radiate afterwards. Overall, plant richness would be lower in arid compared to moist biomes (as observed), but there nonetheless could still be multiple lineages that show recent diversification.

There is extensive evidence that the Namib desert experienced climatic fluctuations during the Quaternary with alternating arid and mesic phases, or pluvial cycles (Scott et al., 1991, 2022; Lancaster, 2002; Dupont, 2006; Gil-Romera et al., 2006; Brook et al., 2007; Grodek et al., 2013; Greenbaum et al., 2014; Lim et al., 2016). These oscillating mesic-arid phases could have provided opportunities for isolation, but also opportunities for previously isolated populations to exchange genes and produce novel genetic combinations, with both processes contributing to diversification, as suggested by Stebbins (1952). Furthermore, high levels of inter-annual variability in rainfall and extended droughts may also drive vegetation die back (Crouchet et al., 2019; Wright et al., 2023), leaving a "clean slate" available for colonisation during mesic periods. In fact, in our most recent fieldwork in the Namib Desert and surrounding areas, we observed extensive dieback of plant populations compared to our first observations in the early 2010s, presumably as a result of widespread drought during the last decade.

Our molecular dataset of thousands of SNPs was sufficient to resolve the deepest nodes in the phylogeny of *Petalidium*, but not relationships within the *diverse arid clade*, which shows low bootstrap support values (Figure 2, Supplementary Figure S1). Such low support was not expected given that RADseq has shown sufficient power to resolve fine-scale phylogenetic relationships in *Barleria*, a closely related genus of Acanthaceae (Comito et al., 2022). This poor resolution in the shallow part of our phylogeny could be explained by hybridisation or incomplete lineage sorting, both of which may cause conflict among genes for relationships. Although field observations (all authors, pers. obs.) document the existence of putative hybrids in *Petalidium*, i.e., individuals with morphology intermediate between two co-occurring species, a preliminary test for hybridisation with our SNP data did not suggest any evidence for hybridisation (Supplementary Figure S2). Given the young age (1.63 Ma) and the high species richness of the *diverse arid clade*, incomplete lineage sorting could be a plausible explanation for the lack of support, but new genetic data, from full genomes or numerous unlinked nuclear loci (c.f. Morais et al., 2019), will be necessary to fully address this issue in *Petalidium*.

Future phylogeny construction based on unlinked nuclear loci, rather than SNP data, may also allow for the implementation of better algorithms for temporally calibrating phylogenies (e.g., Mirarab et al., 2014). We relied on penalized likelihood to calibrate our ML phylogeny, a method that is known to bias node ages towards older time (Carruthers and Scotland, 2022). Nevertheless, our estimates for the ages of younger clades in the phylogeny, such as the *diverse arid clade* are similar to Tripp et al. (2017), suggesting that future improvements to the phylogeny will be unlikely to invalidate our finding that the majority of diversification in *Petalidium*, particularly in the *diverse arid clade*, occurred during the Quaternary.

Our findings show a clear correlation between adaptation to extreme aridity and increased net diversification in *Petalidium*, albeit not a statistically significant one. We have suggested that alternating mesic and arid phases in a present-day arid region have driven this pattern, but there are alternative explanations. The genus exhibits a remarkable diversity of floral forms, associated with different pollinators, including open cup-shaped flowers (a generalist pollination syndrome), large tubular red flowers (bird pollinated), and small maroon flowers (pollinated by long-tongued flies), and the *diverse arid clade* has the greatest floral diversity in *Petalidium*. Interactions with pollinators is a long-hypothesized driver of angiosperm diversity (Van der Niet et al., 2014; Tripp et al., 2021), and changes in floral form linked to switching of pollinators and subsequent reproductive isolation may have promoted speciation in *Petalidium*. Pollinators do seem scarce in this arid region at present, but they may have been less so during past mesic phases. Clearly, further investigation is needed to unravel the potentially interacting factors that promoted such a recent and fast diversification in one of the driest environments on Earth.

Data availability statement

The datasets presented in this study can be found in online repositories. The name of the repository and accession number can be found at: NCBI; PRJNA930345.

Author contributions

OL, EM-T, and KGD conceived the study. EM-T and KGD collected the samples in the field that underlie the phylogeny. EM-T executed laboratory analyses. OL and EM-T conducted bioinformatic and phylogenetic analyses. WS contributed key insights into identification of specimens and phylogenetic results. OL and KGD conducted downstream macroevolutionary analyses and wrote the first draft of the article. All authors contributed to the article and approved the submitted version.

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Supplementary material

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