# Cryptic diversity and phylogeographic patterns of Mediodactylus species in the Eastern Mediterranean region 

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#### Abstract

Cryptic diversity poses a great obstacle in our attempts to assess the current biodiversity crisis and may hamper conservation efforts. The gekkonid genus Mediodactylus, a well-known case of hidden species and genetic diversity, has been taxonomically reclassified several times during the last decade. Focusing on the Mediterranean populations, a recent study within the $M$. kotschyi species complex using classic mtDNA/nuDNA markers suggested the existence of five distinct species, some being endemic and some possibly threatened, yet their relationships have not been fully resolved. Here, we generated genome-wide SNPs (using ddRADseq) and applied molecular species delimitation approaches and population genomic analyses to further disentangle these relationships. The most extensive nuclear dataset, so far, encompassing 2,360 loci and $\sim 699,000 \mathrm{bp}$ from across the genome of Mediodactylus gecko, enabled us to resolve previously obscure phylogenetic relationships among the five, recently elevated, Mediodactylus species and to support the hypothesis that the taxon includes several new, undescribed species. Population genomic analyses within each of the proposed species showed strong genetic structure and high levels of genetic differentiation among populations.


## 1. Introduction

Cryptic species constitute a major challenge in studies assessing biodiversity and a great obstacle in the global efforts to preserve species diversity. The term "cryptic species" is used to describe two or more distinct lineages that have been classified as a single nominal species due to their superficially indistinguishable morphology (Bickford et al., 2007). Although the concept has been known since the 18th century
(Winker, 2005), advances in DNA sequencing, including highthroughput sequencing, now allow for elucidating complex evolutionary histories, shedding light on complex speciation processes in nonmodel organisms and revealing a plethora of cryptic species in mammals (Herrera et al., 2022), reptiles (Engelbrecht et al., 2019), fishes (Guimarães et al., 2022), annelid worms (Bolotov et al., 2022), insects (Schär et al., 2022), mollusks (Sun et al., 2016), plants (Nitta \& Chambers, 2022), fungi (Wyrębek et al., 2021), and bacteria (Williamson

[^0]et al., 2022).
With current species extinction rates being up to 1,000 times higher (Pimm et al., 2014) than the background rate (that is, the pre-human extinction rate or the extinction rate that is not related with anthropogenic factors), the discovery of such unrecognized species is now more important than ever (Dirzo \& Raven, 2003) in order to reevaluate conservation actions and optimize conservation strategies to protect what remains. This is particularly important for the focal region of our study, the Mediterranean basin, one of the world's biodiversity hotspots (Myers et al., 2000). Being at the crossroad of three continents (Africa, Europe, Asia) and exhibiting a complex geological history that left an imprint on the biogeography of many taxa (Lymberakis \& Poulakakis, 2010; Poulakakis et al., 2015), the Mediterranean basin is also a "scientific research hotspot" due to its species richness, its high levels of endemism (Lymberakis \& Poulakakis, 2010; Tierno de Figueroa et al., 2013; Lymberakis et al., 2018), and its susceptibility to climate change (Vogiatzakis et al., 2016). The herpetofauna of the region counts 398 reptile species with $54 \%$ of them being endemic and distributed throughout the basin (Kerim \& Oğzukan, 2017) and 13 \% being threatened (i.e., categorized by the IUCN as vulnerable-VU, endangeredEN, or critically endangered-CR, (IUCN, 2008)). A fraction of these species has been discovered during the last 20-25 years (e.g. 79 reptile species have been added to the herpetofauna of the European region between 2000 and 2020 (Uetz et al., 2022)).

One of the most characteristic examples are the wall lizards of the genus Podarcis in southern Europe, here the initial number of species [17 in Harris and Arnold (1999)] has increased by over 50 \% (Poulakakis et al., 2005; Pinho et al., 2007; Carretero, 2008; Lymberakis et al., 2008; Larbes et al., 2009; Salvi et al., 2017; Psonis et al., 2018; Senczuk et al., 2019; Kiourtsoglou et al., 2021; Psonis et al., 2021) reaching the 26 species that are recognized today (Uetz et al., 2022). Similarly, recent studies have identified new species within various taxa that are distributed in the Mediterranean basin and were long considered as being single species or species complexes, including the blind snake Xerotyphlops (Kornilios et al., 2020a), the Roughtail Rock Agama Laudakia (Karameta et al., 2022), the skink lizard Ablepharus (Skourtanioti et al., 2016), and the green lizard Lacerta (Kornilios et al., 2020b). Many of these "newly described" species are endemics and/or have extremely narrow distribution ranges. For example, the wall lizard $P$. levendis is a steno endemic species only found on two islets (south of the Peloponnese) and is classified by the IUCN as VU, whereas L. cypriaca is endemic to Cyprus and has not been classified by the IUCN yet due to its recent elevation to the species level, but it may be considered as being threatened under criterion B (IUCN criteria for the Red List categories; Criterion B refers to the geographic range of a species).

Evidently, cryptic diversity, the "biodiversity wildcard" (Bickford et al., 2007), constitutes a major challenge in our efforts to devise conservation actions since the discovery of new species, especially endemic ones, substantially affects the conservation strategies as it changes the species richness indicators and the levels of endemism in a given region. Biodiversity parameters such as species richness and endemism are taken into account in the design/identification of protected areas and Key Biodiversity Areas (IUCN, 2016).

The Mediterranean thin-toed gecko (Mediodactylus kotschyi complex) was, until recently, one case of a "species complex" (Böhme et al., 2009). Its taxonomy was reevaluated (Kotsakiozi et al., 2018) based on nuDNA and mtDNA data, recognizing five distinct species within the complex (Fig. 1), some of them being endemic to geographically restricted areas; M. kotschyi (Steindachner, 1870) distributed in the mainland Balkans, most of the Aegean Islands and Italy, M. orientalis (Štěpánek, 1937) in Levant, Cyprus, southern Anatolia, and the south-eastern Aegean Islands, M. danilewskii (Strauch, 1887) in the Black Sea region and in south-west Anatolia, M. bartoni (Štěpánek, 1934) in Crete, and M. oertzeni (Boettger, 1888) occurring only in the southern Dodecanese Islands. This taxonomy was recently adopted by the 2020 update of the Species list of the European herpetofauna (Speybroeck et al., 2020).

Nevertheless, the inter- and intra-phylogenetic relationships of these species remain mostly uncertain.

While DNA-based species delimitation methods have proved to be useful, the identification of speciation events under incomplete lineage sorting (ILS) is challenging (Bamberger et al., 2021). Modern sequencing approaches [such as RADseq (Davey \& Blaxter, 2010), ddRADseq (Peterson et al., 2012), ezRAD (Toonen et al., 2013)] can generate sufficient data to address this challenge. Recent investigations in the lacertid genus Podarcis using genomic data revealed hidden patterns of genetic diversity and provided an improved resolution of their phylogenetic relationships (Garcia-Porta et al., 2019; Psonis et al., 2021; Yang et al., 2021), also suggesting the need for taxonomic revisions. Likewise, genome-wide SNPs have revealed a clearer picture of the phylogenetic relationships and provided a more stable taxonomy for eastern Mediterranean taxa including (a) the Aegean green lizards of the genus Lacerta, leading to the recognition of $L$. citrovittata and L. diplochondrodes (Kornilios et al., 2019, Kornilios et al., 2020b), (b) the Bufotes toads in the eastern Mediterranean (Dufresnes et al., 2019), and (c) the land snail Albinaria cretensis in the western part of the island of Crete (Bamberger et al., 2021).

In this study, we employed a ddRAD sequencing approach and analyzed genome-wide SNP data to elucidate the phylogenetic relationships among the eastern Mediterranean lineages of the genus Mediodactylus as defined in Kotsakiozi et al. (2018). Our objective was to re-evaluate the current taxonomy as well as assess the genomic diversity and the geographic structure of the populations using species-level genomic data.

## 2. Materials and methods

### 2.1. Samples, $D N A$ extraction, $d d R A D$-seq libraries preparation

In total, we used 94 specimens (Table S1) from 60 sampling localities (Fig. 1), covering the largest part of the distribution range of the five species (M. danilewskii, M. kotschyi, M. oertzeni, M. bartoni, M. orientalis; also see Table S1 for the number of individuals sampled per species) in the eastern Mediterranean and representing all major clades and subclades revealed in previous phylogenetic studies (Kasapidis et al., 2005; Kotsakiozi et al., 2018). Total genomic DNA was isolated from tail or tongue tissue of specimens that were preserved frozen $\left(-80{ }^{\circ} \mathrm{C}\right)$ or in ethanol. DNA was isolated using either the DNeasy Blood \& Tissue Extraction kit (Qiagen®, Hilden, Germany) according to the manufacturer's instructions, or an Ammonium Acetate based DNA extraction procedure (Bruford et al., 1998). The quality of the extracted DNA was evaluated using agarose gel electrophoresis (TAE, $1.5 \%$ gel) and quantification of the DNA extracts was performed using the Qubit® 2.0 Fluorometer (Invitrogen ${ }^{\circledR}$, Carlsbad, California, USA).

The double-digest restriction site-associated DNA (ddRAD) libraries were prepared following the protocol of Peterson et al. (2012). Briefly, for the ddRAD library preparation, $\sim 750$ ng of high-quality DNA was simultaneously double-digested using SbfI and MspI (New England BioLabs ${ }^{\circledR}$, Ipswich, MA, USA) restriction enzymes following the manufacturer's instructions. The individual barcoding was followed by the selection of fragments using the Blue Pippin electrophoresis platform (Sage Science, Beverly, MA, USA) under the range selection of 415-515 bp. Targeted fragments were amplified through 11 cycles of Polymerase Chain Reaction (PCR) using the Phusion® Polymerase kit (New England BioLabs®, Ipswich, MA, USA). Libraries were pooled and sequenced (paired-end sequencing, 150-bp reads long) on an Illumina Hi-Seq 2000 lane at the Yale Center for Genome Analysis (Yale University, New Haven, USA).

### 2.2. Sequence data processing

Raw Illumina reads were processed using ipyRAD v.0.9.77 (Eaton \& Overcast, 2020). Samples were demultiplexed using their unique


Fig. 1. The sampling locations of the studied specimens. The green shaded area indicates the distribution range of what was previously considered as the Mediodactylus kotschyi species complex according to the IUCN database. Numbers correspond to the sampling location codes provided in Table S1. The differently colored sampling locations indicate the most recently proposed species-level taxonomy as described in Kotsakiozi et al. (2018): M. kotschyi (red; 1-35), M. orientalis (yellow; 36-49), M. danilewskii (blue; 50-53), M. bartoni (light blue; 54-55), and M. oertzeni (purple; 56-60). (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)
sequence barcodes and Illumina indexes allowing no mismatches between the barcodes of the two reads (Illumina paired-end sequencing). Base calls with Phred quality scores below 20 (default setting; precision of the base call is $99 \%$ ) were converted into undetermined characters $(\mathrm{N})$ and reads including more than five (default setting) Ns were discarded. The minimum genotype depth was set to 6 (according to the ipyrad manual this is approximately the minimum depth at which a heterozygous base call can be distinguished from a sequencing error). The clustering threshold for the de novo assembly was set to 0.90 based on a preliminary analysis (not shown) of our data while following a similar reasoning used by Razkin et al. (2016) and Viricel et al. (2014), we also tested the clustering thresholds of 0.85 and 0.95 . The remaining parameters were left at their default settings, including the minimum number of individuals that have a given locus (set to 4). As a result, we got a sparse matrix, including loci for which at least four samples contain data. Thus, a high proportion of missing data was present in the assembled dataset. To assess the impact of missing data in getting a resolved phylogeny, for the final data assembly, we applied an extra filtering criterion (i.e., the min_taxa; as in https://github. com/ddarriba/ddrad-seq; see below), aiming to determine the minimum amount of data retaining sufficient phylogenetic information for a resolved phylogeny. This is described in detail in recent studies dealing with the effect of missing data on phylogenomic inference of lizard species (Psonis et al., 2018; Psonis et al., 2021). Thus, instead of discarding all loci with missing data above a particular threshold (as one would do by adjusting the min_samples_locus parameter in ipyrad), we retained loci that are phylogenetically informative for parts of the phylogeny with the aim to increase the potential to retain additional phylogenetic information for distinguishing among more divergent taxa at deeper splits in the tree (e.g. see Eaton et al. (2017)).

We generated four different datasets with distinct fractions of phylogenetically informative loci by varying the min_taxa threshold. In the first dataset, we set min_taxa: $=4$ (dataset: Med100) considering that this dataset contains $100 \%$ of the loci. Subsequently, we gradually decreased the amount of missing data by requiring more phylogenetically informative loci to be present [i.e., min_taxa: $=8$ (dataset: Med50), min_taxa $:=13$ (dataset: Med25), min_taxa $:=17$ (dataset: Med12) that correspond to about $50 \%, 25 \%$, and $12.5 \%$ of the loci of the initial Med100 dataset, respectively]. For each one of these datasets, we estimated the missing data per individual and per locus using the propTyped function of the adegenet package in R .

To evaluate these datasets with respect to the impact of missing data and justify our choice of the most stable dataset for comprehensive and final analyses, prior to the phylogenomic analyses, we used Pythia (Haag et al., 2022). Pythia is an open source software tool (https://github. com/tschuelia/PyPythia), that predicts a priori the expected behavior or difficulty of phylogenetic tree searches. We predicted this difficulty for each of the four datasets. Given that a Maximum Likelihood analysis, especially on a large genomic dataset, is time and resource intensive, it is helpful to predict a priori the "potential" of a given dataset to either converge to topologically similar tree topologies or to result in multiple statistically indistinguishable yet topologically highly distinct trees. In other words, Pythia predicts and quantifies, on a scale ranging between 0.0 (easy dataset) and 1.0 (extremely difficult), the difficulty of analyzing a given dataset. As such, it increases user awareness and allows to devise an effective as well as appropriate analysis strategy (e.g., increase the number of independent tree searches to construct a reliable tree on a "difficult" dataset). Although Pythia predicted the dataset with the least missing data (Med12; score 0.07; see Results) as being least difficult, the scores provided for the other three datasets were low (easy-to-analyze datasets) as well (0.09-0.16; see Results). Therefore, we also performed i) preliminary DAPC (see Section "Population Genomics Analyses" below) and ii) Maximum Likelihood analyses (for settings see Section "Phylogenomic Analyses"), on all four datasets. Then, we used the --rfdist option to compute the topological Robinson-Foulds (RF) distance (Robinson \& Foulds, 1981) among 50 ML trees, in a preliminary
investigation on how the amount of missing data (See Results Section) affects the results.

### 2.3. Phylogenomic analyses

For the dataset that Pythia suggested (Dataset Med12 including the 94 samples and the full sequences with a length of $698,737 \mathrm{bp}$; see below) as having the best potential for a resolved phylogeny, we used ModelTest-NG (https://github.com/ddarriba/modeltest; (Darriba et al., 2019), to predict the best model of evolution for the phylogenetic analyses. We performed a Maximum Likelihood (ML) tree inference using RAxML-NG (v.1.0.3; (Kozlov et al., 2019)) under the GTR + gamma model, with 50 random starting trees using 25 random and 25 parsimony-based starting trees (the default value for this step is 20 tree searches, but we increased this number to 50 to explore the tree space more thoroughly). To check the bootstrap convergence of the best scoring tree in each analysis we used the -bsconvergence option and the bootstrap support (BS) was also calculated and mapped onto the bestscoring ML tree of the selected dataset. We also performed an ML analysis using only the unlinked SNPs (one SNP per locus, the dataset was assembled using the $R$ scripts available at https://github. com/ddarriba/ddrad-seq) of the selected dataset (Med12) using the Lewis (Lewis 2001) ascertainment bias correction. The command lines used for the ML analysis using RAxML-NG are provided in the Supplementary Material (Code for analyses). A Bayesian Inference (BI) analysis was performed for the selected (Dataset Med12 see below) dataset using MrBayes v.3.2.7 (Ronquist et al., 2012) and under the GTR + gamma model. The MCMC analysis ran for $1,000,000$ generations using two independent runs with four chains each. The result was saved every 1,000 generations and for the "burn in" we discarded the first $25 \%$ of samples. Apparent convergence of the BI analysis was evaluated using the Estimated Sample Size (ESS $>200$ ) and the Potential Scale Reduction Factor (PSRF $=1.0$ ). The produced trees were visualized using FigTree v.1.4.4 (http://tree.bio.ed.ac.uk/software/figtree/).

To test if the uneven representation of species and relevant missing data (see Results Section; Tables 2 and S2) affect our phylogenomic analyses, we performed an additional ML analysis on a pruned version of the Med12 dataset. The distributional pattern of missing data in our dataset is due to the overrepresentation of M. kotschyi (Table S1; ~60 \% of the samples) with respect to the remaining species (see also Section 3.4. Species Delimitation). Thus, we pruned the dataset down to 22 samples used in order to keep between 4 and 6 samples per species (except M. bartoni for which only 2 samples are available). The samples were selected such as to have similar proportions of missing data (Table S2).

In order to account for incomplete lineage sorting (ILS) that can induce gene trees/species trees incongruences that in turn might heavily impact phylogenetic reconstructions we also performed a coalescent based phylogenetic analysis with SVDquartets (Chifman and Kubatko, 2014) as implemented in *PAUP (Cummings, 2004) using the multi locus data of the 94 sample dataset (Med12). SVDquartets infers the species tree directly from the site patterns and therefore bypasses the impact of gene tree estimation error. The analysis was executed i) considering the two best supported 8 - and 12 -species delimitation schemes and ii) based on the current taxonomy considering the five species. Runs were performed using exhaustive Quartet sampling with 200,000 random quartets and 1,000 bootstrap replicates.

The trees inferred by all phylogenetic inference methods were unrooted. Initially, we attempted to root the tree, using the Mediterranean house gecko (Hemidactylus turcicus) as outgroup. However, due to the high amount of missing data, the $H$. turcicus sequences were excluded from the final dataset. To determine the most probable root of the tree, we used the RootDigger tool (Bettisworth \& Stamatakis, 2021) using as input the ML tree. RootDigger can indicate the most likely root location on a given unrooted tree and infers a confidence value for the possible root placement. We kept the parameters as default and the
exhaustive mode which evaluates the likelihood of placing the root into every branch of the tree, and as such it allows us to quantify root placement uncertainty.

### 2.4. Population genomic analyses

The population structure within each species was evaluated using the Bayesian clustering method implemented in STRUCTURE v.2.3.4 (Pritchard et al., 2000) via the STRUCTURE_THREADER pipeline v.1.3.10 (Pina-Martins et al., 2017). We used STRUCTURE to identify genetic clusters and assign individuals to these clusters without prior information about the sampling location. The analysis was performed on the Med12 dataset (based on the Pythia score and the RF distances) and was conducted on a per species basis (as defined in Kotsakiozi et al. (2018) and currently adopted by the 2020 update of the Species list of the European herpetofauna), though only for those species where more than six samples were available (M. kotchyi, M. orientalis, M. oertzeni). To comply with the assumption of independence across loci, we subsampled our dataset by selecting one SNP per locus using respective R scripts (https://github.com/ddarriba/ddrad-seq). This filtered dataset was also used in all population genomic analyses (see below) and from now on, we will refer to it as Med12_1snp dataset. For each analysis the most likely allocation of samples to clusters (K), was determined by conducting 10 independent runs for each K ranging from 1 to 10. Each run assumed an admixture model and independent allele frequencies and used a burn-in period of 100,000 and 500,000 generations. The best K was selected based on the deltaK method of Evanno et al. (2005) using STRUCTURE_THREADER (Earl \& vonHoldt, 2012). Results were summarized and plotted with CLUMPAK that accounts for label switching and multimodality (Kopelman et al., 2015).

To complement the Bayesian analysis, we also performed a Principal Component Analysis (PCA) with the R package LEA (Frichot \& François, 2015) and a Discriminant Analysis of Principal Components (DAPC) of ADEGENET R package, using the Med12_1snp dataset. We used the find. clusters option of the ADEGENET R package (Jombart et al., 2010) in order for individuals to be assigned to DAPC-defined clusters, without $a$ priori defining samples to populations/groups. The number of DAPCclusters is chosen based on the lowest BIC value. DAPC transforms the raw data using a PCA and then a DA is applied on the retained principal components to provide an efficient description of the genetic clusters using a few synthetic variables (discriminant functions) that are linear combinations of the original variables (raw data) (Jombart et al., 2010). Thus, the among-group variance is maximized while the within-group variance is minimized.

Same Med12_1snp dataset was then used to estimate the FST distance and perform AMOVA analyses. Pairwise genetic differentiation (FST) between groups of populations and their statistical support (p-value: 0.05) were calculated in Arlequin v3.5.2.2 (Excoffier \& Lischer, 2010), using 16,000 permutations (according to the manual that guarantees to have less than $1 \%$ difference with the exact probability in $99 \%$ of the cases). The partitioning of the genomic variation among and within populations was evaluated through a hierarchical Analysis of MOlecular VAriance (AMOVA) (Excoffier et al., 1992), as implemented in Arlequin, using 16,000 permutations. Details on the grouping for the AMOVA analyses are provided in Table S3.

### 2.5. Species delimitation analysis

Acknowledging that species delimitation can be challenging and that different approaches may yield conflicting results, we conducted species delimitation using two methods; a) the BFD* method (Leaché et al., 2014) and b) the multi-rate PTP (mPTP) (Kapli et al., 2017).

Species delimitation with the BFD* method was performed on a subset of the Med12_1snp dataset (selected dataset; see Results Section) using SNAPP (Bryant et al., 2012) package in BEAST2 v.2.7.5 (Bouckaert et al., 2019). This was deemed necessary since BFD* is computationally
demanding, and thus we reduced our dataset based on the tree topology inferred via ML and BI to contain fewer representatives from every major clade or subclade (named Med12_snapp dataset; see Table S1 for the samples used in this reduced dataset). However, given the substantially more $M$. kotschyi samples used compared to the other species (see Fig. 1 and Table S1), this resulted in an overrepresentation of the M. kotschyi haplotypes. This overrepresentation might be an issue that should be taken into account in a species delimitation analysis [for details see (Magoga et al., 2021)] since the higher the number of sampled haplotypes, the higher the probability to find intermediate haplotypes among closely related species becomes. The case of M. orientalis is analogous, though less evident. Thus, to test if this unbalanced Med12_snapp dataset affects our results, we performed one more BFD* analysis (on the dataset named Med12_snapp2) by randomly subsampling the Med12_snapp dataset, in order for each of the species to be equally represented by $4-8$ samples (except $M$. bartoni that had only two samples). SNP data was converted to binary format with phrynomics R package (https://github.com/bbanbury/phrynomics/). BFD* uses a Yule prior with a parameter lambda $(\lambda)$ representing the speciation rate. We estimated the $\lambda$ value using the pyule script (https://github. com/joaks1/pyule). The script required the tree height (estimated based on the tree produced by the analysis of the concatenated sequences of the most stable dataset: Med12; see Results; Phylogenomic Analyses) and the number of tips/species as input. The number of tips/ species varied from four to twelve depending on the species model scheme (see Table 1 for details in the scenarios), thus resulting in different $\lambda$ values (from 40.1 considering four species to 77.9 considering twelve species). Mutation rates $u$ and $v$ were set to one and were not sampled, while intraspecific variance was set to $0.1(10 \%, \alpha=1, \beta=$ 10 , Rateprior $=$ gamma) and coalescence rate was sampled with a starting value of 10 , following the settings used in similar studies for lizard species (e.g. Psonis et al. (2018)). The BFD* analysis was run with a chain length of 100,000 generations, alpha $=0.3,50 \%$ burn-in percentage and 48 steps. The analyses were executed in BEAST using a chain length of $1,000,000$ generations and samples were stored every 10,000 generations. Apparent convergence for each delimitation scheme analysis as well as species tree estimation was assessed using Tracer and ESS values (ESS > 200).

Specimens were assigned to the following alternative species delimitations (i) Model 1 (RunA), the four groups revealed by

Table 1
BFD* analysis results for Mediodactylus species delimitation models. Clades coding refers to Fig. 3. Bayes Factor (BF) delimitation was used for model selection and was estimated based on the marginal likelihood estimate (MLE) value for each model. Positive BF values indicate support for the alternative model, and negative BF values indicate support for the null model (the model with the highest MLE). The most highly supported scheme is shown in bold.

| Model [partition of clades/ <br> subclades] | Species | MLE | Rank | BF |
| :--- | :--- | :--- | :--- | :--- |
| RunH [A1/A2/A3/B1-Muğ1a/B1- <br> Cyprus/B1-Israel/B1-Adana <br> /B2/C/D/E1/ E2] - tree | 12 | -115.97 | 1 | - |
| topology-mPTP delimitation <br> RunG [A1/A2/A3/B1/B2/C/D/E] <br> $\quad$ - tree topology | 8 | -368.53 | 2 | -158.88 |
| RunF [A1/A2/A3/B/C/D/E] - tree <br> topology | 7 | -457.96 | 3 | -683.98 |
| RunE [A/B1/B2/C/D/E] - tree <br> topology <br> RunD [A1A2/A3/B/C/D/E] - tree <br> topology <br> RunC [A/B/C/D/E] - current <br> taxonomy <br> RunB [A1/A2/A3/BCD/E] - DAPC <br> groups | 6 | 6 | -504.88 | 4 |

preliminary PCA on the entire 94 sample dataset, (ii) Model 2 (RunB) the five groups revealed by DAPC analyses on the entire 94 sample dataset, (iii) Model 3 (RunC), the five currently recognized species model, (iv) Model 4-8 (RunD-H), the groups revealed by the phylogenetic, DAPC and STRUCTURE analyses, in which the species number ranged from four to twelve (Table 1). More specifically, Model 4 (RunD), six species model with two species within M. kotschyi, Model 5 (RunE), six species model with two species within $M$. orientalis, Model 6 (RunF), seven species model with three species within M. kotschyi, Model 7 (RunG) eight species model with three species within M. kotschyi, and two species within M. orientalis, and Model 8 (RunH) twelve species model with three species within M. kotschyi, five species within M. orientalis and two species within M. oertzeni. Following Leaché et al. (2014), Bayes factor Delimitation (BFD*) was used to select among alternative delimitations and estimated as follows: $\mathrm{BF}=2 \times$ (MLE1-MLE0) where MLE0 was the marginal likelihood estimate value of the best model (Table 1) and MLE1 was the marginal likelihood estimate value for each alternative model evaluated against model 0 . The strength of support from BF comparisons of competing models can be evaluated using the framework of (Kass and Raftery, 1995). The BF scale is as follows: $0<$ $\mathrm{BF}<2$ is not worth more than a bare mention, $2<\mathrm{BF}<6$ is positive evidence, $6<\mathrm{BF}<10$ is strong support, and $\mathrm{BF}>10$ is decisive.

The second species delimitation approach we employed, mPTP (Kapli et al., 2017), is an improved version PTP and does not require the user to define any analysis parameters (i.e. similarity thresholds, cutoffs, etc). The method uses a Markov chain Monte Carlo (MCMC) sampling approach and computes support values for each delimitation of the input tree. Those values can be used to assess the confidence of the inferred ML delimitation scheme. For the mPTP analysis we used the concatenated sequence data of the Med12 dataset and the respective ML tree which we uploaded to the mPTP web server (https://mcmc-mptp.h-its.org/mcm $\mathrm{c} /$ ).

## 3. Results

## 3.1. ddRADseq data metrics

After quality filtering, the sequencing of the ddRAD libraries resulted in $1,046,505$ reads on average per sample (Table S2). The number of loci per sample after applying the clustering threshold, the average depth of clusters per individual, and the percentage of complete genotypes per individual are presented in Table S2. The filtered ipyrad assembly included 32,964 loci, each being present in at least four samples (Min$\operatorname{Cov}=4$, paralogs removed) with an average of 3,554 loci per sample (Table S2). The SNPs matrix produced by ipyrad included 419,529 variable sites ( $84.6 \%$ missing) with a total of 30,517 unlinked SNPs.

The application of the extra filter to the ipyrad dataset resulted in four datasets (Med100, Med50, Med25, Med12), where the number of loci, the length of sequences in the assembly, and metrics regarding the percentage of missing data are presented in Table 2. For the selected by Pythia dataset (Med12 see below), the fraction of missing genotypes per sample ranged from $37 \%$ to $91 \%$ (Table S2). The proportion of missing data per locus (see also Table 2) ranged from $0 \%$ (i.e. some loci and specifically 86 out of the 2360 loci, were present in all 94 samples) to 83 \% (i.e. 3 out of the 2360 loci have missing data in $83 \%$ of the individuals).

### 3.2. Phylogenomic analyses

Pythia suggested the Med12 dataset while the scores for all four datasets (Med100, Med50, Med25, and Med12) were also low i.e. 0.16, $0.09,0.15$, and 0.07 respectively. This dataset was also suggested by estimating the RF distances among all pairs of 50 inferred ML trees of each dataset. Therefore, this dataset was used for subsequent ML, BI, and the SVDquartets analyses. ML analysis converged after 400 trees (cut-off threshold 0.01 ) and resulted in the robustly supported tree (average BS

Table 2
Number of loci and length of sequences (bp) in the assembly for the four filtered datasets (min_taxa $=4,8,13,17$ ). Estimates for the missing data per dataset are also provided and specifically the percentage of missing data for the total matrix (\% of missing data), the average percentage of missing data per individual (\% average per indv), the range of missing genotypes (from the 94 -sample dataset) per locus (Range miss per locus), and the number of loci that are missing in more than $50 \%$ and $75 \%$ of the samples ( $>50 \%$ and $75 \%$ ) in each dataset.

| dataset | Med100 <br> (min_taxa 4) | Med50 <br> (min_taxa 8) | Med25 <br> (min_taxa <br> 13) | Med12 <br> (min_taxa <br> 17) |
| :---: | :---: | :---: | :---: | :---: |
| \% of loci | 100 \% | 50 \% | 25 \% | 12.5 \% |
| Number of loci | 18,300 | 8,664 | 4,182 | 2,360 |
| Sequence length (bp) | 5,342,266 | 2,547,210 | 1,234,877 | 698,737 |
| $\begin{aligned} & \text { \% of missing } \\ & \text { data }^{1} \end{aligned}$ | 86.8 \% | 78.7 \% | 69.5 \% | 61.7 \% |
| $\begin{aligned} & \text { \% of missing } \\ & \text { data }^{2} \end{aligned}$ | 84 \% | 76,3\% | 68 \% | 61.5 \% |
| \% average per indv ${ }^{1}$ | 86.6 \% | 78.6 \% | 69.7 \% | 61.7 \% |
| Range miss per locus ${ }^{1}$ | 0 \%-98 \% | 0 \%-95 \% | 0 \%-90 \% | 0 \%-84 \% |
| $>50 \%$ and | 12,884 and | 5,792 and | 2,603 and | 1,761 and |
| $75 \%^{1}$ | 11,463 | 4,393 | 1,394 | 439 |

${ }^{1}$ for the SNP dataset where one SNP per locus was retained and used in the population genomics analyses.
${ }^{2}$ for the complete sequence dataset used in the phylogenomics analyses.
on the tree equals to 92.2) presented in Fig. 2. BI analysis resulted in a tree with high BS Posterior Probabilities (PP; 0.96-1.00) and with identical topology (PP values are also presented in Fig. 2) to the one from ML. The phylogenomic inference confirmed the presence of five major clades within the eastern Mediterranean Mediodactylus taxa, each with high statistical support $[\mathrm{PP}=1.00, \mathrm{BS}=100]$, which correspond to the five currently recognized species. The SVDquartets analysis (Fig. 3) resulted in a tree with the same topology as the ML/BI trees presented in Fig. 2. Importantly, the species tree inferred with SVDquartets showed twelve highly supported clades that are geographically separated i.e. species occupy non overlapping regions, as shown in Fig. 3. The ML analysis on the SNPs matrix (not shown) also robustly supported (BS values 94-100) the presence (and the grouping of samples within each one) of the twelve clades (see Fig. 3). Finally, the tree topology remained unaltered for the ML analysis on the pruned dataset with 22 samples.

The rooted tree produced by RootDigger analysis placed, with high probability (lwr $=0.99$ ), M. danilewskii (Fig. 2), a species that ranges from Crimea to the coastline of Türkiye, and to the East Aegean islands (Fig. 1; blue), as being the most likely root of the tree. The most densely sampled clade, the one of $M$. kotschyi, can be robustly subdivided into three subclades; one hosts samples from continental Greece and the north/central Aegean Islands (called A1), another one includes the Kythira/Antikythira Islands samples (A2), and the third one comprises the Cyclades and the island of Kos that geographically belongs to the east Aegean Islands (A3). M. oertzeni which is distributed in the southeast Aegean Islands (Fig. 1; violet) seems to be a sister clade of M. kotschyi and closely related to the Crete's clade, M. bartoni. Last, M. orientalis (Fig. 2), which is further subdivided into two subclades; one including samples from western Türkiye (i.e. Aydin) and the east Aegean Islands (called B2) and one including samples from southern Türkiye (i.e. Adana, Gaziantep), Cyprus, and Israel (called B1).

### 3.3. Population genomics analyses

Genetic structure: The Evanno method (Evanno et al., 2005) on the population STRUCTURE analysis for M. kotschyi (Fig. 2, Fig. S3) supported the presence of two clusters $(\mathrm{K}=2 ; \mathrm{Q}$ values $>0.95)$, which


Fig. 2. Maximum Likelihood (ML) tree reconstructed using ddRAD data. Bayesian Inference (BI) analysis resulted in an identical topology, bootstrap (BS) support values and Posterior Probabilities (PP) from the ML and the BI analyses, respectively, are shown on the branch nodes of the tree. Individual codes follow those in Table S1 with the first two digits representing the map codes of Fig. 1. Letters and numbers on the nodes are used to label the respective clade/subclade of the tree and are consistent using the coding used for the BFD* grouping schemes in Table 1. The vertical STRUCTURE bar plots on the right, indicate the groups of populations identified by the respective analysis on each clade/subclade of the tree. The spots on the emended maps indicate the geographic distribution of the STRUCTURE defined groups.


Fig. 3. Bootstrap $50 \%$ majority-rule consensus tree from SVDQuartets analysis for twelve lineages/species as they were delimited by BFD* and mPTP analyses and their respective distribution on the maps.
correspond to the A1/A2 and A3 clades of the phylogenetic tree that contains a split within this clade forming two monophyletic lineages; A1/A2 and A3 (Fig. 3). Hierarchical STRUCTURE analysis then showed the separation of A1 from A2 (STRUCTURE on the A1/A2 cluster; $\mathrm{K}=2$ ) and then clear geographic differentiation within each subclade (Fig. 2, Fig. S3) -that were also supported by BFD* and mPTP as possibly different species (see Results section-Species delimitation). Specifically, the three population clusters supported by DeltaK for A1, coincide with the split observed within this subclade (Fig. 2) separating the islands from continental Greece and the north Aegean Islands as well as from the Peloponnese (Fig. 2). The DeltaK method resulted in similar conclusions for subclades A2 and A3 as in both cases $K=2$ is returned as the most likely choice. In both cases the clustering (A2 = Kythira and Antikythira Islands and satellite islets; A3 = north and south part of the Cyclades) coincide with the splits observed in the tree of Fig. 2. For M. oertzeni (Fig. 2, Fig. S4), Delta K method supported two clusters of populations, while for M. orientalis (Fig. 2, Fig. S4) K = 6 was the best supported value according to the deltaK method, albeit five major clusters were plotted by CLUMPAK (Fig. S4; 1-Cyprus, 2-Adana/Gaziantep, 3-Israel, 4-Aydin/ East Aegean Islands, $5-\mathrm{Mug}$ la) since the 6 th cluster ( Q value of the 6 th cluster in major cluster equals 0.0004 ) appears only in the minor clustering (in 3 out of the 10 CLUMPAK runs) scheme. In both cases the results of STRUCTURE analyses supported the geographic differentiation and are in agreement with the tree topology of Fig. 2.

Regardless of the filtering used (retaining $12.5 \%$ to $100 \%$ of the loci), PCA (Fig. S1.A) suggested the differentiation of $M$. kotschyi from all the remaining ones, while DAPC suggested the differentiation of
M. kotschyi and M. oertzeni (Fig. S1.B) from the remaining ones. Based on this finding we proceeded to the next two DAPC analyses using the Med12 dataset (as indicated by Pythia) and filtered as to keep one SNP per locus. For M. kotschyi, the DAPC analysis (Fig. 4A) supported the presence of eight DAPC-groups that are in agreement to both ML/BI and coalescent trees in Figs. 2, 3). In particular, we found a clear distinction according to the first axis, between the samples that originated from the north/central Aegean Islands and continental Greece (A1) from the remaining samples. Based on the second axis of DAPC, the samples from the Kythira/Antikythira Islands (A2) are differentiated from those from the southern Cyclades Islands (A3). The DAPC-groups defined within clade A1 (Fig. 4A; groups 2 to 5; Peloponnese, Kythnos Isl., continental Greece, and central Aegean Islands, respectively) largely coincide with the distinct clusters defined by the hierarchical STRUCTURE analysis within clade A1 (Fig. 2). The DAPC on the other group of species, indicated a clear distinction between species as M. bartoni, M. oertzeni, and $M$. danilewskii which form distinct groups. Interestingly, M. orientalis showed substantial differentiation (Fig. 4B) that is also similar to the STRUCTURE clustering for this species (Fig. 2, Fig. S4) forming five groups; (i) Adana-Israel, (ii) Cyprus, (iii) Muğla (Türkiye), and (iv) Ikaria-Fournoi Islands (east Aegean), and (v) one sample from the Muğla clusters within the danilewskii-group.

The results of the AMOVA analysis are presented in Table S3. The vast majority of the genetic variation ( $66 \%-91.7 \%$ ) was observed among groups. More specifically, when we considered six to twelve groups (grouping as in Schemes D, F, G, H of the BFD* analysis; See Table 1) the variation among groups exceeded $90 \%$ compared to a


Fig. 4. Discriminant Analysis of Principal Components (DAPC) for Mediodactylus populations that belong (A) to M. kotschyi and (B) to other Mediodactylus lineages of the eastern Mediterranean. Individuals are represented as dots with the different colors representing the DAPC-groups defined. A bar plot of eigenvalues for the discriminant analysis (DA eigenvalues) is displayed in each inset. The plots are made using the first two DAs in both cases.
variation of $68 \%$ among groups that the current taxonomy scheme (presented in RunC of Table 1) attained.

Genetic differentiation: Pairwise FST estimates between the major clades of the tree (Fig. 2; A, B, C, D, and E; current taxonomy) receive high values $(F S T>0.68)$. Also high values $(F S T>0.75)$ were received among the three clades of $M$. kotschyi that coincide with delimited species (see below). Regarding clade A1, high FST values (0.46-0.57) were estimated between groups of populations (i.e. north Cyclades/ north Aegean-continental Greece/Peloponnese). A lower level of differentiation ( $F S T=0.36$ ) was recorded between the two clusters of south Cyclades Islands (Clade A3) and a higher level of differentiation $(F S T=0.63)$ was observed between the Kythira/Antikythira Islands (Clade A2). The two subclades of Clade B (Figs. 2, 3; B1/B2) showed a moderate differentiation compared to the rest level of differentiation (Fst $=0.34$ ), whereas high differentiation $(F S T=0.81)$ was observed between the two subclades of Clade E (Figs. 2, 3; E1/E2).

### 3.4. Species delimitation

The Marginal Likelihood Estimates (MLE) that were obtained from the first BFD* analysis, which was based on the PCA, DAPC, STRUCTURE and ML/BI tree topology (dataset; Med12_snapp), are presented in Table 1 and the results of the second BFD* analysis (dataset; Med12_snapp2) aiming to avoid overrepresentation of the sample-rich clades (i. e., including 4-8 samples per clade) are presented in Table S4. Both analyses supported the twelve lineages scheme (Table 1; RunH) as the delimitation of choice ( BF values $>10$; decisive) coinciding with well supported lineages in ML/BI and coalescent trees (Figs. 2, 3). The mPTP also supported the presence of twelve delimited species. More specifically, both analyses supported the two recently recognized species (M. danilewski, M. bartoni) and supported additional delimited species within M. kotschyi (subclades A1, A2, A3 as being distinct species), M. orientalis, and $M$. oertzeni clades. Within the $M$. orientalis clade, BFD* and mPTP supported the delimitation of five species (1-Cyprus, 2-Israel, 3Adana, 4-Muğla, 5-Ikaria-Fournoi-Aydin) and within M. oertzeni clade supported the delimitation of two species (Karpathos-Rhodes and SymiTilos).

## 4. Discussion

During the last decade, the feasibility to use thousands of genomewide DNA markers in non-model organisms opened a new era in phylogenomics, revolutionized the field and revealed complex evolutionary processes and biogeographic patterns. In this study, using an extensive nuclear dataset including thousands of loci across the genome of the five recently recognized Mediodactylus species of the eastern Mediterranean not only we were able to confirm their monophyly, but also reveal additional hidden species diversity in the study area. Our analyses produced a clearer picture of the evolutionary relationships and intraspecific population structure and revealed that three species (M. kotschyi, M. orientalis, and M. oertzeni) comprise species complexes. For M. kotschyi, the presence of three species is robustly supported by our results. For M. orientalis and M. oertzeni our data support the presence of five and two species within each complex respectively, however, these findings should be interpreted with caution given the high proportion of missing data for these two species. Last, the twelve delimited species seem to have non overlapped distributional ranges and that the paleogeography of the region played an important role on shaping their distributions.

## Species delimitation and phylogenetic relationships

The five recognized Mediodactylus species (M. kotschyi, M. orientalis, M. danilewskii, M. bartoni, and M. oertzeni) of the Eastern Mediterranean region form well-supported, monophyletic clades, confirming the morphological grouping of Beutler (1981) and the recent raising of those groups to species level (Kotsakiozi et al., 2018). More specifically, the kotschyi, bartoni and oertzeni morphological groups proposed by Beutler (1981) represented exactly these taxa, while the danilewskii group was split into two species; M. danilewskii and M. orientalis. The most differentiated species is $M$. kotschyi with a relatively broad geographic range and high levels of genetic differentiation among populations. The species tree produced by SVDquartets revealed that the three highly supported lineages within M. kotschyi - which were also supported by both species delimitation methods as being different species- comprise sister taxa with A3 from south-eastern Cyclades being the most differentiated one. Mediodactylus danilewskii, and M. orientalis seem to be more closely
related to each other compared with the other species as they cluster close to each other in DAPC (Fig. 4). However, it is important to note that the conclusions about these species should be interpreted with caution, as these species, particularly the relatively widespread $M$. danilewskii, are undersampled.

The species delimitation analyses supported the scheme of twelve putative species. Specifically, mPTP and BFD* analyses indicated M. danilewskii and M. bartoni clades as distinct species while they supported the presence of three species within M. kotschyi, the five-species scheme within $M$. orientalis and the presence of two species within M. oertzeni. Note that the species delimitation supported by BFD* and mPTP for $M$. orientalis and $M$. oertzeni completely coincide with the DAPC and STRUCTURE results for these two species. AMOVA analysis further supported the pattern indicated by BFD* as the percentages of variation among groups were maximized ( $>90 \%$ ) when we considered the species delimitation schemes that were best supported by BFD*. This enhanced the validity of this specific grouping of lineages.

We emphasize that the high percentage of missing data for a number of samples, which is anticipated since our dataset includes several distinct species, and the fact that a couple of species are undersampled considering their distribution range, did not allow us to draw strong conclusions regarding a possible taxonomic revision. We tested the effect of missing data on the analyses by producing four datasets (Med100, Med50, Med25, Med12) containing different percentages of missing data (from $61 \%$ to $86 \%$; Table 2). Among the four datasets, and as expected, the most stable dataset was the one (Med12) with the lowest proportion of missing data. However, Pythia predicted low scores for all four datasets and identical (or almost identical) results were obtained for the four datasets during preliminary analyses (tree topology and population clustering). These observations supported the idea that the percentage of missing data, although it was -on average- relatively high, does not affect the main results. This can be attributed to the fact that the filters applied here, aim to retain phylogenetic informativeness and preserve the phylogenetic signal in the data. The inclusion of more missing data among more divergent taxa increased the probability of encompassing more phylogenetic information for deeper cladogenetic events in a tree (Eaton et al., 2017). Similar findings regarding the effect of missing data on phylogenomics have been observed in other studies (Takahashi et al., 2014; Wang et al., 2017; Psonis et al., 2018; Psonis et al., 2021). Nonetheless, we do have strong evidence that more species complexes exist within the taxon. For example, one of the species that appeared to be a species complex with possibly five species is M. orientalis. However, this species is undersampled and exhibited a high percentage of missing data. This indicates that a denser sampling strategy is needed which will result in a more complete genomic dataset for this species before strong conclusions about taxonomic revisions can be drawn. The other species complexes revealed were $M$. oertzeni, a species with a restricted geographic distribution and M. kotschyi. Although these species cannot be considered as being undersampled, complementary studies (e.g. ecology, traditional taxonomy etc) are needed prior to issuing taxonomic recommendations. However, it is important to note that the higher number of loci used here, allowed us to unravel hidden diversity that remained undetected before. In particular, two out of the five species were confirmed (M. danilewskii, M. bartoni) and the presence of three other species complexes is suggested; M. kotschyi with three, M. orientalis with five, and M. oertzeni with two robustly supported distinct groups of lineages (putative species), respectively.

## Phylogeography

For M. kotschyi, the three distinct groups of lineages (subclades) supported by SVDquartets results are also geographically distinct (see Fig. 3) with the first (A1) being present in mainland Greece and the north/central Aegean islands, the second (A2) in the Kythira island group, and the third (A3) in the central and southern Cyclades, with high FST values among them. Based on the subspecies taxonomy, the first subclade includes different morphological subspecies [e.g. M. kotschyi
skopjensis (Karaman, 1965) and M. k. kotschyi (Steindachner, 1870] from the A3 lineage [e.g. A3 samples had been assigned to M. k. concolor (Bedriaga, 1881, 1882)]. The absence of a clear intraspecific structure and fully resolved differentiation within $M$. kotschyi in the study of Kotsakiozi et al. (2018) can potentially be attributed to the use of few loci in contrast to the genome wide information of a high number of loci of the present study. Here, the higher resolution that the SNPs data offer, allowed for a more fine-grained species delimitation and detection of three distinct groups of lineages within M. kotschyi, two of which are located solely on islands. This further supports the theory that islands (as here the Mediterranean islands) harbor hidden diversity (Pérez-Delgado et al., 2022).

Focusing on the geographic distribution of M. kotschyi subclades, there is a north-to-south and an east-to-west differentiation. Within M. kotschyi, the split of the Cyclades into north-western (in subclade A1) and south-eastern islands (subclade A3), has also been observed in other animal species, such as the Euscorpius scorpions (Parmakelis et al., 2006), vipers (Vipera ammodytes complex; Thanou et al. (2023)), and partially in Podarcis lizards (Poulakakis et al., 2005; Yang et al., 2021) as well as in Mesobuthus scorpions (Parmakelis et al., 2006), reflecting the known geological separation of the Cyclades ( $\sim 3.5$ Mya) (Popov et al., 2004). In subclade A1, there are three distinct subgroups (Figs. 2, 4) exhibiting clear geographical differentiation: the Peloponnese, the northwestern Cyclades Islands/Skyros Island, and mainland Greece. The inclusion of the north Aegean Islands (Lemnos Island and Thassopoula islet) within the mainland populations is likely due to their recent geological separation from the nearby mainland (Popov et al., 2004). Subclades A2 and A3 exclusively consist of island populations. Subclade A2 consists of individuals originated from a biogeographical distinct island group at the southwest edge of the Aegean archipelago, which includes Kythira Island (to the north), Antikythira Island (to the south), and the Pori and Lagouvardos islets (in between). Phylogenetically, subclade A2 is closely related to subclade A1, which includes the Peloponnese and shares geographical proximity with the Kythira island group. This biogeographical pattern is observed in several animal taxa, such as Podarcis (Spilani et al., 2019) and Ablepharus (Skourtanioti et al., 2016).

According to palaeogeographic data, the larger island of Kythira to the north submerged during the Pliocene (Meulenkamp, 1985), whereas there is no evidence of similar tectonic movements affecting Pori, Lagouvardos, and Antikythira (a larger islet south of Pori and Lagouvardos) during that period. The presence of M. kotschyi on all the islands within this group suggests that either the group was colonized from the Peloponnese following the re-emergence of Kythira or that the populations of Pori, Lagouvardos, and Antikythira islets remained unaffected by the Pliocene tectonic rearrangements, giving rise to the population of Kythira following its re-emergence. These findings align with the divergence times inferred in Kotsakiozi et al. (2018), supporting the divergence of the Kythira island group from the mainland at the end of the early Pliocene ( 3.9 Mya ), while the differentiation within this group occurred during the Pleistocene.

The subclade A3 consists of individuals that originated from the central and south Cyclades Islands. Of particular note, the island of Kos that biogeographically belongs to the east Aegean Islands clustered within A3. The Aegean Sea constitutes a major contemporary barrier to biotic exchange between mainland Greece and Türkiye. The palaeogeographic evolution of the Aegean region has been described in detail in several studies (Parmakelis et al., 2006; Papadopoulou et al., 2010; Poulakakis et al., 2015; Kornilios et al., 2019), starting from the united landmass (Ägäis) of the middle Miocene to the formation of the Aegean Barrier (AB) in the late Miocene (10-9 Mya) [for more details see Kornilios et al. (2019)], causing the separation of the west Aegean (Cyclades Islands) from the east Aegean Islands. This pattern is imprinted in the biogeography and phylogeny of animal species (especially in those with limited overseas dispersal abilities). However, there are several cases of animal species, characterized as 'naughty' in Poulakakis et al. (2015),
that have passed the Aegean Barrier (e.g., Ablepharus kitaibelii, Podarcis erhardii, Pelophylax bedriagae, Trachelipus aegaeus, Albinaria brevicollis, Dichomma dardanum, Zonites rhodius). So, the presence of Kos (east Aegean) in the subclade of Cyclades can be either one more case of 'naughty' animal or a human-aided dispersal, which is not uncommon for Mediodactylus (Koynova et al., 2017; Mares \& Novarini, 2020; Urošević et al., 2021).

On the other hand, M. orientalis, with a broad distribution area (Fig. 1), is divided into two robustly supported clades (see Fig. 2. B1; Türkiye, Cyprus, and Israel and B2; western Türkiye and East Aegean Islands). The first subclade showed a clear geographic differentiation as it is divided into four lineages (Fig. 3) exhibiting an east-west differentiation. The first lineage is from Israel, the second is from southeast Türkiye, the third from southwestern Türkiye and the fourth from Cyprus, which can be explained by the geomorphology of the area (e.g., the Taurus Mountains, Anatolian Diagonal, Nur Mountains) and the isolation of Cyprus. The Anatolian Diagonal is a line of mountain ranges that run from the south of Gümüşhane - Bayburt in the north, southwest across Türkiye to the Taurus Mountains (Mutun, 2010). It is thought to consist a significant geographic barrier shaping the distribution of various species across Türkiye and dividing lineage distribution into east and west (Ciplak et al., 1993; Rokas et al., 2003; Sengor et al., 2003; Mutun, 2010; Bilgin, 2011). The uplift of the Nur Mountains (during late Pliocene) seems to explain the isolation of the populations distributed at the southern-east edge of the taxon's distribution. Of particular interest is the region of southwestern Türkiye (Muğla region in our case), in which two different species of Mediodatylus are present (M. orientalis and M. danilewskii). This area is extremely rich in biodiversity, with the presence of distinct phylogenetic lineages, even at species level, in particular for several reptile species, such as Ophiomorus kardesi (Kornilios et al., 2018), Laudakia stellio (Karameta et al., 2022), Xerotyphlops vermicularis (Kornilios, 2017), and Blanus strauchi (Sindaco et al., 2014).

Considering the lineage from Cyprus, it seems that Cyprus is more closely related to south Türkiye. Cyprus has been isolated for at least 5.3 Ma from the surrounding continental regions, with which it has probably never been connected, making it one of the very few and by far the largest, oceanic islands of the Mediterranean Sea (Dimitriou et al., 2022). Kotsakiozi et al (2018) estimated that the isolation of the Cyprus lineage occurred in the late Miocene ( $\sim 6 \mathrm{Mya}$ ). This time corresponds to the Messinian Salinity Crisis ( $\sim 6-5.3$ Mya; (Krijgsman et al., 1999)), when the island was connected with, or being closer to Anatolia either through a land bridge or via a series of intermediate islets used as stepping-stones (Poulakakis et al., 2013).

The second subclade of $M$. orientalis corresponds to the area of east Aegean islands and western Türkiye, which can be attributed to the paleogeographic history of this region where the east Aegean islands were connected to Türkiye even during the late Pleistocene (Perissoratis \& Conispoliatis, 2003; Lykousis, 2009; Sakellariou \& Galanidou, 2017). It is worth noting that the five groups of populations within $M$. orientalis clade indicated by STRUCTURE analysis and coinciding with the subclades and lineages of the phylogenetic tree were also supported as delimited species by mPTP and BFD* analyses. Interestingly, these five lineages (see Fig. 2) also correspond to distinct subspecies; the lineage from Israel to M. orientalis orientalis (Štěpánek, 1937), the lineage of Cyprus to M. orientalis fitzingeri (Štěpánek, 1937) while the lineage from Adana-Gaziantep has been suggested to belong to M. orientalis bolkarensis (Rösler, 1994). However, a finer-scale sampling strategy along the Middle East coastline might help to disentangle the phylogenetic relationships within the taxon and provide insights into its phylogeographic history. In any case, we stress the need for additional studies focusing on the ecology, the morphology, and the biology of the taxa under study to fully conclude on the suggested species status.

Two other clades that appear in the tree (M. bartoni and M. oertzeni), are island species with restricted geographic distribution (southeast Aegean for M. oertzeni and Crete and surrounding islets for M. bartoni). The presence of only two individuals for $M$. bartoni did not permit us to
assess the intraspecific diversity of this species in more detail. For M. oertzeni on the other hand, there was a clear differentiation between the islands of Rhodes and Karpathos from Tilos and Symi islands. Populations of these two groups had previously been described to belong to two distinct subspecies [oertzeni: Rhodes-Karpathos and beutleri: Symi; See Valakos et al. (2008)] that completely coincide with the two delimited species supported by our analysis. The close phylogenetic affinity of Mediodactylus geckos from Rhodes and Karpathos islands is a common pattern in animal species [e.g. water frogs of the genus Pelophylax (Lymberakis et al., 2007) and ground beetles of the genus Dendarus (Trichas et al., 2020)]. Karpathos, which was an island during the Miocene, was joined with Rhodes and Anatolia in the Early Pliocene (Daams \& Van de Weerd, 1980) and it was permanently isolated during the Late Pliocene (Böger \& Dermitzakis, 1987). Taking into account the estimated time of divergence of Karpathos and Rhodes by Kotsakiozi et al. (2018) in the Middle Pleistocene ( $\sim 1$ Mya), the distribution of M. oertzeni on Karpathos Isl. is the result of the dispersal of an ancestral form of $M$. oertzeni from Rhodes Isl. to Karpathos Isl., when Karpathos was already an island.

Mediodactylus danilewskii was estimated as the most probable root of the tree (Fig. 2). The species is distributed in a broad geographic area, expanding from Crimea to south Türkiye. Unfortunately, our small sample size did not allow us to investigate the genetic structure of its populations. However, given the concordance between our findings and those of Kotsakiozi et al. (2018) and by taking into account the presence of this species in Bulgaria and its subsequent introduction to different areas of Bulgaria (Koynova et al., 2020) and along the Turkish coasts of the Black Sea (Bülbül et al., 2023), we can hypothesize that this species covers a much broader area than the one sampled here, as samples from north Greece, north Türkiye, and Bulgaria [included in Kotsakiozi et al. (2018), but not in the present study] cluster within it. Thus, given the substantial morphological variation that this species exhibits within its range (Ajtić 2014; Pulev et al., 2014), a finer sampling strategy and a subsequent population genetic/omic analysis within M. danilewskii will shed more light on its evolutionary history.

## 5. Conclusion

Genomic data, phylogenomic analyses and current species delimitation methods are powerful tools for the study of cryptic diversity (Bickford et al., 2007; Chattopadhyay et al., 2016; Tang et al., 2022). These tools enabled us to reveal the relationships among Mediodactylus species at almost the entire distribution range of Mediodactylus in the eastern Mediterranean region, unveiling hidden diversity. More specifically, genomic data confirmed the monophyly of the recent raising of M. kotschyi, M. orientalis, M. danilewskii, M. bartoni, and M. oertzeni lineages to species level, revealing, however, that three of them are species complexes that require further investigation. Our results suggest that there are possibly twelve Mediodactylus potential species with nonoverlapping distributional ranges in the Eastern Mediterranean region. This is because M. kotschyi, M. orientalis, and M. oertzeni appear to form species complexes with three, five, and two species within each complex, respectively. However, we emphasize the need for additional studies before a potential systematic revision, including a larger number of sampled localities for at least two of these species as well as the inclusion of other types of evidence, such as, for instance, morphological studies. Some of the newly suggested species are island endemics (e.g., M. bartoni endemic to Crete and satellite islets, M. oertzeni E1 or E2 lineage endemic to southeast Aegean Islands) and some of them may be classified as being threatened in upcoming IUCN evaluations. Given the rate of species discovery since the adoption of the phylogenetic species concept, the distribution and the number of hotspots around the globe (Peterson \& Navarro-Siguenza, 1999) might still change substantially. Unraveling cryptic diversity contributes to addressing several of the shortfalls that Hortal et al. (2015) identified as biodiversity knowledge gaps. These shortfalls (e.g., Linnean Shortfall; knowledge gaps in
taxonomy, Wallacean Shortfall; in species distribution, Prestonian Shortfall; in abundance and population dynamics) severely affect our efforts to preserve biodiversity, which is critical for the ecosystems and human societies (Díaz et al., 2018). The use of genomic data and current species delimitation methods serve as a first step to unravel cryptic diversity, even for taxa that display complex evolutionary relationships.

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## CRediT authorship contribution statement

Panayiota Kotsakiozi: Writing - original draft, Visualization, Investigation, Formal analysis. Aglaia Antoniou: Writing - review \& editing, Resources, Investigation, Data curation. Nikolaos Psonis: Writing - review \& editing, Resources. Kostas Sagonas: Writing - review \& editing, Resources. Emmanouela Karameta: Writing - review \& editing, Resources. Çetin Ilgaz: Writing - review \& editing, Resources. Yusuf Kumlutaş: Writing - review \& editing, Resources. Aziz Avcı: Writing - review \& editing, Resources. Daniel Jablonski: Writing review \& editing, Resources. Diego Darriba: Writing - review \& editing, Methodology, Data curation. Alexandros Stamatakis: Writing - review \& editing, Methodology, Data curation. Petros Lymberakis: Writing review \& editing, Resources. Nikos Poulakakis: Writing - review \& editing, Supervision, Resources, Project administration, Funding acquisition, Conceptualization.

## Data availability

Code used for analyses is included in the Supp Material. Genomic data produced have been deposited in SRA NCBI. The Radseq data used in the analyses are available on the NCBI SRA in demultiplexed form, under BioProject number PRJNA1051519 and BioSample accessions SAMN38777126- SAMN38777219.

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## Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi. org/10.1016/j.ympev.2024.108091.

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## Supplementary Material

Supplementary material for the manuscript titled:
Cryptic diversity and phylogeographic patterns of Mediodactylus species in the Eastern Mediterranean region

This document contains:

- Tables S1-S4
- $\quad$ Figures S1, S2, S3, S4, S5
- $\quad$ Code for the analyses


## Tables

Table S1. Mediodactylus samples used in this study, including samples that were also used in Kotsakiozi et al. (2018) study and newly added samples in the dataset that are indicated by bold characters. Map codes are as in Figure 1 and sample codes are as in Figure 2. NHMC: Natural History Museum of Crete.


| 11_Lagouvardos(1634) | Lagouvardos isl., South of Peloponnese [11] | 80.3.85.1634 |
| :---: | :---: | :---: |
| 12_Pori(1635) | Pori isl., South of Peloponnese [12] | 80.3.85.1635 |
| 12_Pori(1636) | Pori isl., South of Peloponnese [12] | 80.3.85.1636 |
| 13_Thasopoula(1637)\# | Thasopoula isl., North Aegean [13] | 80.3.85.1637 |
| 13_Thasopoula(1638) | Thasopoula isl., North Aegean [13] | 80.3.85.1638 |
| 13_Thasopoula(1639) | Thasopoula isl., North Aegean [13] | 80.3.85.1639 |
| 14_Milos(1641) | Milos isl., Cyclades [14] | 80.3.85.1641 |
| 14_Milos(1642) | Milos isl., Cyclades [14] | 80.3.85.1642 |
| 15_Milos(1646) | Milos isl., Cyclades [15] | 80.3.85.1646 |
| 15_Milos(1647)\#2 | Milos isl., Cyclades [15] | 80.3.85.1647 |
| 16_Polyaigos(1649) | Polyaigos isl., Cyclades [16] | 80.3.85.1649 |
| 16_Polyaigos(1650) | Polyaigos isl., Cyclades [16] | 80.3.85.1650 |
| 17_Kimolos(1653) | Kimolos isl., Cyclades [17] | 80.3.85.1653 |
| 17_Kimolos(1654)\#2 | Kimolos isl., Cyclades [17] | 80.3.85.1654 |
| 17_Kimolos(1655) | Kimolos isl., Cyclades [17] | 80.3.85.1655 |
| 18_Sikinos(1662) | Sikinos isl., Cyclades [18] | 80.3.85.1662 |
| 18_Sikinos(1663)\# | Sikinos isl., Cyclades [18] | 80.3.85.1663 |
| 19_Ismarida(1665)\# | Ismarida lake, Thrace [19] | 80.3.85.1665 |
| 20_Loudias(1666) | Loudias river, Thessaloniki [20] | 80.3.85.1666 |
| 21_Thira(1667)\# | Thira isl., Cyclades [21] | 80.3.85.1667 |
| 21_Thira(1668) | Thira isl., Cyclades [21] | 80.3.85.1668 |
| 22_Skyros(1670)\#2 | Skyros isl., Aegean [22] | 80.3.85.1670 |
| 23_Skyros(1671) | Skyros isl., Aegean [23] | 80.3.85.1671 |
| 24_Skyros(1673) | Skyros isl., Aegean [24] | 80.3.85.1673 |
| 25_Naxos(1696)\# | Naxos isl., Cyclades [25] | 80.3.85.1696 |


|  | 26_Kythira(1707) | Kythira isl., South of Peloponnese [26] | 80.3.85.1707 |
| :---: | :---: | :---: | :---: |
|  | 26_Kythira(1708)\#2 | Kythira isl., South of Peloponnese [26] | 80.3.85.1708 |
|  | 26_Kythira(1709) | Kythira isl., South of Peloponnese [26] | 80.3.85.1709 |
|  | 27_Lemnos(1716) | Lemnos isl., North East Aegean [27] | 80.3.85.1716 |
|  | 28_Peloponnese(1715) | Arkadia, Peloponnese [28] | 80.3.85.1715 |
|  | 29_Peloponnese(1718)\# | Arkadia, Peloponnese [29] | 80.3.85.1718 |
|  | 30_Peloponnese(1808)\#2 | Geraki, Peloponnese [30] | 80.3.85.1808 |
|  | 31_Peloponnese(1814) | Sikionas, Peloponnese [31] | 80.3.85.1814 |
|  | 32_Peloponnese(1820) | Gargaliani, Peloponnese [32] | 80.3.85.1820 |
|  | 33_Kythnos(1858)\# | Kythnos isl., Cyclades [33] | 80.3.85.1858 |
|  | 34_Folegandros(1868) | Folegandros isl., Cyclades [34] | 80.3.85.1868 |
|  | 35_los(1888) | Ios isl., Cyclades [35] | 80.3.85.1888 |
| M. orientalis | 36_Cyprus(1967)\#2 | Cyprus [36] | 80.3.85.1967 |
|  | 36_Cyprus(1889) | Cyprus [36] | 80.3.85.1889 |
|  | 36_Cyprus(1890) | Cyprus [36] | 80.3.85.1890 |
|  | 36_Cyprus(1891) | Cyprus [36] | 80.3.85.1891 |
|  | 37_Adana(1387) | Feke, Adana, Türkiye [37] | 80.3.85.1387 |
|  | 37_Adana(1723)\#2 | Feke, Adana, Türkiye [37] | 80.3.85.1723 |
|  | 38_Mugla(1528) | Kötekli, Muğla, Türkiye [38] | 80.3.85.1528 |
|  | 38_Mugla(1529) | Kötekli, Muğla, Türkiye [38] | 80.3.85.1529 |
|  | 39_Fournoi(1710) | Fournoi isl., SouthEast Aegean [39] | 80.3.85.1710 |
|  | 39_Fournoi(1711) | Fournoi isl., SouthEast Aegean [39] | 80.3.85.1711 |
|  | 40_Ikaria(1712)\#2 | Ikaria isl., SouthEast Aegean [40] | 80.3.85.1712 |
|  | 41_Ikaria(1713) | Ikaria isl., SouthEast Aegean [41] | 80.3.85.1713 |
|  | 41_Ikaria(1714) | Ikaria isl., SouthEast Aegean [41] | 80.3.85.1714 |


|  | 42_Adana(1721)\#2 | Kedikli Kozan, Adana, Türkiye [42] | 80.3.85.1721 |
| :---: | :---: | :---: | :---: |
|  | 43_Mugla(1722)\#2 | Muğla University, Muğla, Türkiye [43] | 80.3.85.1722 |
|  | 44_Aydin(1753)\#2 | Dam-Çine, Aydin, Türkiye [44] | 80.3.85.1753 |
|  | 45_Gaziantep(1758)\#2 | Zincirli-Islahiye, Gaziantep, Türkiye [45] | 80.3.85.1758 |
|  | 46_Adana(1763) | Kutkulagi-Ceyhan, Adana, Türkiye [46] | 80.3.85.1763 |
|  | 47_Mugla(1764) | Muğla, Muğla, Türkiye [47] | 80.3.85.1764 |
|  | 48_Israel(1824)\#2 | Malkishua, Israel [48] | 80.3.85.1824 |
|  | 49_Israel(1830) | Modiin, Israel [49] | 80.3.85.1830 |
| M. danilewskii | 50_Antalya(1389)\#2 | Gazipasa, Antalya, Türkiye [50] | 80.3.85.1389 |
|  | 51_Kastelorizo(1717) | Kastelorizo, South East Aegean [51] | 80.3.85.1717 |
|  | 52_Mugla(1759)\#2 | Kapikargin-Dalaman, Muğla, Türkiye [52] | 80.3.85.1759 |
|  | 53_Crimea(1772)\#2 | Theodosiyan, Crimea [53] | 80.3.85.1772 |
| M. bartoni | 54_Chrysi(1542)\#2 | Chrysi isl., SE of Crete [54] | 80.3.85.1541 |
|  | 55_Mikronisi(1704)\#2 | Mikronisi isl., South of Crete [55] | 80.3.85.1704 |
| M. oertzeni | 56_Symi(1556)\# | Symi isl., SouthEast Aegean [56] | 80.3.85.1556 |
|  | 56_Symi(1557) | Symi isl., SouthEast Aegean [56] | 80.3.85.1557 |
|  | 56_Symi(1558) | Symi isl., SouthEast Aegean [56] | 80.3.85.1558 |
|  | 57_Rodos(1559)\#2 | Rhodes isl., SouthEast Aegean [57] | 80.3.85.1559 |
|  | 57_Rodos(1560) | Rhodes isl., SouthEast Aegean [57] | 80.3.85.1560 |
|  | 58_Karpathos(1578) | Karpathos isl., SouthEast Aegean [58] | 80.3.85.1578 |
|  | 59_Karpathos(1620) | Karpathos isl., SouthEast Aegean [59] | 80.3.85.1620 |
|  | 59_Karpathos(1621)\#2 | Karpathos isl., SouthEast Aegean [59] | 80.3.85.1621 |
|  | 60_Tilos(1615) | Tilos isl., SouthEast Aegean [60] | 80.3.85.1615 |
|  | 60_Tilos(1616)\#2 | Tilos isl., SouthEast Aegean [60] | 80.3.85.1616 |

[^1]2: Samples also used in the Med12_snapp2 dataset for the SNAPP analysis. Samples were randomly selected from the Med12_ snapp dataset in an attempt to have a more balanced dataset in terms of species representation (each species was represented by 3-8 samples).

Table S2. Marker discovery and ddRADseq data metrics for all the Mediodactylus samples used in this study. The number of raw reads (raw_reads) after Illumina sequencing, the mean number of loci after applying the clustering threshold equal to 0.90 (clusters), the average depth of clusters (avg_depth) the number of loci in the final filtered ipyrad assembly (loci) and the percentage of complete genotypes in the final selected dataset (Med12) after applying the min_taxa filtering on a per sample basis are presented. Sample codes are as in Table S1.

| Sample | raw_reads | clusters | avg_depth | loci | Med12_co <br> mplete ${ }^{\mathbf{1}}$ | Med12_co <br> mplete $^{2}$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 01_Gyaros( <br> 1530) | 1134974 | 144503 | 81.39 | 3259 | 49.2 | 47.3 |
| 01_Gyaros( <br> 1532) | 1009758 | 100667 | 61.26 | 4059 | 57.1 | 54.9 |
| 02_Serifos( <br> 1544) | 828316 | 104915 | 51.23 | 4256 | 59.4 | 56.9 |
| 02_Serifos( <br> 1545) | 749049 | 75601 | 60.69 | 3425 | 52.4 | 50.2 |
| 03_Andros <br> (1551) | 1838469 | 210334 | 83.02 | 5019 | 61.2 | 59.2 |
| 03_Andros <br> (1552) | 792419 | 90331 | 55.48 | 3845 | 53.6 | 51.8 |
| 04_Paros(1 <br> 567) | 1170918 | 133890 | 68.24 | 4503 | 58.3 | 54.4 |
| 05_Astypal <br> aia(1573) | 1310127 | 124829 | 95.77 | 3352 | 48.7 | 45.5 |
| 05_Astypal <br> aia(1574) | 862984 | 87697 | 67.82 | 3465 | 50.4 | 47.3 |
| 06_Kos(15 <br> 76) | 1180819 | 101033 | 81.5 | 3837 | 53.1 | 49.7 |
| 06_Kos(15 <br> 77) | 1055829 | 155349 | 77.17 | 3564 | 51.4 | 48.5 |
| 07_Amorg <br> os(1609) <br> 07_Amorg <br> os(1610) | 1302763 | 146540 | 104.95 | 3098 | 45.5 | 42.4 |


| 07_Amorg <br> os(1611) | 1516583 | 162593 | 93.47 | 4178 | 58.1 | 54.9 |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- |
| 08_Trichon <br> ida(1623) | 650822 | 78996 | 53.99 | 2634 | 45.0 | 43.8 |
| 09_Syros(1 <br> 626) | 1206941 | 152069 | 73.55 | 3954 | 55.5 | 52.8 |
| 09_Syros(1 <br> 627) | 877343 | 105137 | 57.09 | 3938 | 49.2 | 47.3 |
| 10_Antikit <br> hira(1630) | 1117785 | 148188 | 67.54 | 4121 | 55.6 | 53.1 |
| 10_Antikit <br> hira(1632) | 1056053 | 134804 | 73.11 | 3392 | 48.1 | 45.7 |
| 11_Lagouv <br> ardos(1633 <br> ) | 1130114 | 110685 | 69.58 | 4561 | 57.8 | 54.4 |
| 11_Lagouv <br> ardos(1634 <br> ) | 921165 | 119277 | 68.4 | 3829 | 52.6 | 50.5 |
| 12_Pori(16 <br> 35) | 443972 | 48968 | 44.14 | 2767 | 57.8 | 37.5 |
| 12_Pori(16 <br> 36) | 1652716 | 198852 | 61.7 | 5755 | 58.8 | 56.5 |
| 13_Thasop <br> oula(1637) <br> 13_Thasop <br> oula(1638) <br> 13_Thasop <br> oula(1639) | 1032742 | 1408213 | 141671 | 65481 | 56.06 | 4815 |


| 15_Milos(1 <br> 647) | 1192775 | 142226 | 74.08 | 4339 | 59.2 | 55.4 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 16_Polyaig <br> os(1649) | 914692 | 114417 | 56.05 | 4584 | 53.8 | 50.6 |
| 16_Polyaig <br> os(1650) | 1003672 | 91543 | 73.01 | 3655 | 53.4 | 50.0 |
| 17_Kimolo <br> s(1653) | 497616 | 60094 | 45.45 | 3241 | 47.8 | 37.5 |
| 17_Kimolo <br> s(1654) | 771376 | 77768 | 61.47 | 3468 | 51.4 | 47.9 |
| 17_Kimolo <br> s(1655) | 606791 | 75910 | 36.53 | 4555 | 55.3 | 51.2 |
| 18_Sikinos( <br> 1662) | 1095197 | 105342 | 74.94 | 4024 | 56.2 | 52.9 |
| 18_Sikinos( <br> 1663) | 890040 | 128253 | 71.65 | 3276 | 47.0 | 43.4 |
| 19_Ismarid <br> a(1665) | 663337 | 62416 | 50.54 | 3231 | 47.9 | 45.8 |
| 20_Loudias <br> (1666) | 655276 | 64949 | 60.79 | 2832 | 42.7 | 39.9 |
| 21_Thira(1 <br> 667) | 1019002 | 114019 | 51.86 | 5232 | 60.9 | 58.2 |
| 21_Thira(1 <br> 668) | 776457 | 92410 | 52.66 | 4007 | 56.9 | 54.2 |
| 22_Skyros( | 1421306 | 111580 | 88.9 | 3821 | 57.1 | 54.1 |
| 1670) <br> 23_Skyros( <br> 1671) <br> 24_Kythira <br> (1707) | 886853 | 92200 | 68.17 | 3184 | 48.3 | 46.4 |
| 25_Nkyros( <br> 1673) | 1506351 | 150407 | 83.6 | 4296 | 60.0 | 57.6 |


| $\begin{gathered} \text { 26_Kythira } \\ (1708) \end{gathered}$ | 718855 | 66815 | 50.87 | 3753 | 52.7 | 50.9 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| $\begin{gathered} \text { 26_Kythira } \\ (1709) \end{gathered}$ | 806999 | 101530 | 51.24 | 4158 | 55.9 | 53.6 |
| $\underset{(1716)}{27}$ | 632674 | 95220 | 45.91 | 3289 | 47.1 | 44.6 |
| 28_Pelopo nnese(171 5) | 2290383 | 245862 | 105.04 | 4403 | 54.7 | 51.7 |
| 29_Pelopo nnese(171 <br> 8) | 1102900 | 104516 | 64.9 | 3966 | 57.8 | 54.5 |
| 30_Pelopo nnese(180 <br> 8) | 1332512 | 85474 | 67.86 | 5722 | 61.2 | 59.4 |
| 31_Pelopo nnese(181 <br> 4) | 2411785 | 182668 | 111.67 | 6014 | 62.9 | 60.4 |
| 32_Pelopo nnese(182 $0)$ | 4343002 | 391421 | 160.89 | 6299 | 60.9 | 59.2 |
| $\begin{aligned} & \text { 33_Kythno } \\ & \text { s(1858) } \end{aligned}$ | 95792 | 21491 | 14.06 | 1437 | 22.4 | 22.0 |
| 34_Folega ndros(1868 ) | 190449 | 29853 | 23.32 | 2475 | 31.8 | 31.0 |
| $\begin{gathered} 35 \_ \text {los(188 } \\ 8) \end{gathered}$ | 381718 | 31048 | 35.2 | 3634 | 44.8 | 43.3 |
| Average for species M. kotschyi | 1109879 | 117553.2 | 67.40 | 4040.11 | 53.55 | 50.5 |
| $\begin{gathered} \text { 36_Cyprus( } \\ \text { 1967) } \end{gathered}$ | 1446819 | 180150 | 90.31 | 2551 | 12.5 | 16.1 |
| $\begin{gathered} \text { 36_Cyprus( } \\ \text { 1889) } \end{gathered}$ | 742856 | 90062 | 55.97 | 2615 | 14.1 | 18.2 |


| $\begin{gathered} \text { 36_Cyprus( } \\ \text { 1890) } \end{gathered}$ | 1492420 | 168313 | 50.39 | 4586 | 31.1 | 34.3 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| $\begin{gathered} \text { 36_Cyprus( } \\ \text { 1891) } \end{gathered}$ | 979773 | 126975 | 71.38 | 2537 | 14.4 | 18.4 |
| 37_Adana( 1387) | 463927 | 59704 | 37.47 | 2079 | 13.6 | 17.8 |
| 37_Adana( 1723) | 263488 | 37515 | 27.72 | 1624 | 10.7 | 14.4 |
| $\begin{gathered} \text { 38_Mugla( } \\ \text { 1528) } \end{gathered}$ | 1422327 | 209587 | 98.39 | 3064 | 13.8 | 18.1 |
| $\begin{gathered} \text { 38_Mugla( } \\ \text { 1529) } \end{gathered}$ | 492609 | 64817 | 56.72 | 2509 | 10.0 | 13.5 |
| $\begin{aligned} & \text { 39_Fournoi } \\ & \text { (1710) } \end{aligned}$ | 1108486 | 124333 | 69.26 | 3473 | 17.0 | 22.6 |
| $\begin{aligned} & \text { 39_Fournoi } \\ & \text { (1711) } \end{aligned}$ | 2479092 | 231876 | 114.21 | 3882 | 17.3 | 22.4 |
| $\begin{gathered} \text { 40_Ikaria(1 } \\ 712) \end{gathered}$ | 625557 | 115109 | 43.42 | 3093 | 13.6 | 17.6 |
| $\begin{gathered} \text { 41_Ikaria(1 } \\ \text { 713) } \end{gathered}$ | 456763 | 69144 | 43.36 | 2663 | 12.5 | 16.5 |
| $\begin{gathered} \text { 41_Ikaria(1 } \\ 714) \end{gathered}$ | 1694687 | 204035 | 81.21 | 3752 | 16.7 | 21.7 |
| $\begin{gathered} \text { 42_Adana( } \\ \text { 1721) } \end{gathered}$ | 541594 | 85656 | 40.42 | 1978 | 14.4 | 18.6 |
| $\begin{gathered} \text { 43_Mugla( } \\ \text { 1722) } \end{gathered}$ | 835393 | 106926 | 73.55 | 2816 | 12.3 | 16.3 |
| $\begin{gathered} \text { 44_Aydin(1 } \\ 753) \end{gathered}$ | 1195509 | 140945 | 45.56 | 4157 | 18.7 | 24. 4 |
| $\begin{gathered} \text { 45_Gaziant } \\ \text { ep(1758) } \end{gathered}$ | 782427 | 114215 | 61.46 | 1877 | 12.8 | 16.9 |
| $\begin{gathered} \text { 46_Adana( } \\ \text { 1763) } \end{gathered}$ | 927972 | 122158 | 63.83 | 2071 | 13.8 | 18.2 |
| $\begin{gathered} \text { 47_Mugla( } \\ \text { 1764) } \end{gathered}$ | 294947 | 56525 | 30.45 | 2552 | 10.4 | 13.6 |


| $\begin{gathered} \text { 48_Israel(1 } \\ 824) \end{gathered}$ | 1106859 | 67318 | 70.43 | 1869 | 13.0 | 17.9 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| $\begin{gathered} \text { 49_Israel(1 } \\ 830) \end{gathered}$ | 414477 | 57131 | 28.52 | 1783 | 12.8 | 17.1 |
| Average for species M. orientalis | 941332.5 | 115833 | 59.71571 | 2739.571 | 14.55 | 18.78 |
| 50_Antalya (1389) | 754196 | 104525 | 50.52 | 1408 | 13.6 | 17.5 |
| 51_Kastelo rizo(1717) | 408014 | 107881 | 21.78 | 1338 | 12.2 | 15.7 |
| $\begin{aligned} & \text { 52_Mugla( } \\ & \text { 1759) } \end{aligned}$ | 291647 | 48754 | 28.49 | 960 | 09.2 | 11.7 |
| $\begin{gathered} \text { 53_Crimea } \\ (1772) \end{gathered}$ | 571390 | 71510 | 36.76 | 1387 | 11.3 | 14.9 |
| Average for species M. danilewskii | 506311.8 | 83168 | 34.3875 | 1273.25 | 11.57 | 14.95 |
| $\begin{aligned} & \text { 54_Chrysi( } \\ & \text { 1542) } \end{aligned}$ | 899234 | 107814 | 49.25 | 1122 | 14.3 | 18.2 |
| 55_Mikroni si(1704) | 784569 | 84700 | 55.41 | 1007 | 12.9 | 16.1 |
| Average for species M. bartoni | 841901.5 | 96257 | 52.33 | 1064.5 | 13.6 | 17.15 |
| $\begin{gathered} \text { 56_Symi(1 } \\ \text { 556) } \end{gathered}$ | 685926 | 97382 | 55.89 | 3284 | 14.0 | 18.0 |
| $\begin{aligned} & \text { 56_Symi(1 } \\ & \text { 557) } \end{aligned}$ | 1140652 | 118825 | 71.46 | 3974 | 19.4 | 24.7 |
| $\begin{gathered} \text { 56_Symi(1 } \\ 558) \end{gathered}$ | 1850738 | 287744 | 100.27 | 3798 | 16.2 | 20.3 |
| $\begin{aligned} & \text { 57_Rodos( } \\ & \text { 1559) } \end{aligned}$ | 1150528 | 163357 | 57.68 | 4284 | 20.6 | 26.4 |


| $\begin{aligned} & \text { 57_Rodos( } \\ & \text { 1560) } \end{aligned}$ | 1453036 | 163166 | 77.32 | 4205 | 20.6 | 26.3 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 58_Karpat hos(1578) | 872843 | 130102 | 59.59 | 3329 | 17.7 | 22.3 |
| 59_Karpat hos(1620) | 775559 | 102432 | 43.45 | 3831 | 20.5 | 25.9 |
| 59_Karpat hos(1621) | 950093 | 120032 | 52.33 | 4060 | 21.9 | 27.8 |
| $\begin{aligned} & 60 \text { _Tilos(16 } \\ & \text { 15) } \end{aligned}$ | 1079491 | 164478 | 73.22 | 3649 | 17.1 | 21.8 |
| $\begin{gathered} 60 \text { _Tilos(16 } \\ \text { 16) } \end{gathered}$ | 1672509 | 215170 | 64.82 | 4671 | 20.2 | 25.9 |
| Average for species M. oertzeni | 1163138 | 156268.8 | 65.603 | 3908.5 | 18.8 | 23.9 |
| 1 Dataset Med12 after filtered using the code in https://github.com/ddarriba/ddrad-seq and one SNP per locus was kept |  |  |  |  |  |  |

Table S3. Analyses of molecular variance (AMOVA). Populations are divided into groups as in species delimitation analysis shown in Table 1.

| Groups [clades/subclades of the phylogenetic tree] | Source of variation | df | Percentage of variation (\%) |
| :---: | :---: | :---: | :---: |
| 5 groups [A / B / C / D / E] | Among groups | 4 | 68.1 |
| as current taxonomy | Within groups | 14 | 26.8 |
|  | Within populations | 75 | 5.1 |
| 4 groups [A1 / A2 / A3 / BCDE] | Among groups | 3 | 86.0 |
| as PCA grouping | Within groups | 15 | 8.5 |
|  | Within populations | 75 | 5.5 |
| 5 groups [A1 / A2 / A3 / BCD / E] | Among groups | 4 | 85.0 |
| as DAPC grouping | Within groups | 14 | 9.1 |
|  | Within populations | 75 | 5.9 |
| 6 groups [A / B1 / B2 / C / D / E] | Among groups | 5 | 65.5 |
| as tree topology | Within groups | 13 | 29.2 |
|  | Within populations | 75 | 5.2 |
| 6 groups [A1A2 / A3 / B / C / D / E] | Among groups | 5 | 91.7 |
| as tree topology | Within groups | 13 | 2.6 |
|  | Within populations | 75 | 5.7 |
| 7 groups [A1 / A2 / A3 / B / C / D / E] | Among groups | 6 | 90.9 |
| as tree topology | Within groups | 12 | 3.1 |
|  | Within populations | 75 | 6.0 |


| 8 groups [A1 / A2 / A3 / B1 / B2 / C / D / E] | Among groups | 7 | 90.8 |
| :---: | :--- | :---: | :---: |
| as tree topology | Within groups | 11 | 3.1 |
|  | Within populations | 75 | 6.1 |
| 12 groups [A1 / A2 / A3 / B1a / B1b / B1c / | Among groups | 11 | 90.1 |
| B1d / B2 / C / D / E1 / E2] |  |  |  |
| as tree topology | Within groups | 7 | 3.5 |
|  | Within populations | 75 | 6.3 |

[^2]Table S4. BDF* results for Mediodactylus species delimitation models, performed using 3-8 samples per major clade in order to account for the overrepresentation of Clade A haplotypes. Clades' coding refers to Fig. 3. Bayes Factor (BF) delimitation was used as a model selection tool and it was estimated based on the marginal likelihood estimate (MLE) value for each model. Positive BF values indicate support for the alternative model, and negative BF values indicate support for model 0 (here the highly supported model). Best supported scheme is shown in bold.

| Model [partition of <br> clades/ subclades] | Species | MLE | Rank | BF |
| :---: | :---: | :---: | :---: | :---: |
| RunH [A1 / A2 / A3 / B1- <br> Adana/ B1-Mugla/ B1-Cyprus/ B1- <br> Isreal / B2 / C / D / E]-tree topology <br> and mPTP delimited scheme | $\mathbf{1 2}$ | -109.9 | $\mathbf{1}$ | - |
| RunG [A1 / A2 / A3 / B1 / B2 / C / D / <br> E]-tree topology | 8 | -294.49 | 2 | -369.18 |
| RunF [A1 / A2 / A3 / B / C / D / E]-tree <br> topology | 7 | -361.51 | 3 | -503.22 |
| RunE [A / B1 / B2 / C / D / E]-tree <br> topology | 6 | -366.52 | 4 | -513.36 |
| RunD [A1A2 / A3 / B / C / D / E]-tree <br> topology | 6 | -397.11 | 5 | -574.54 |
| RunC [A / B / C / D / E] ; current <br> taxonomy | 5 | -538.29 | 6 | -856.90 |
| RunA [A1 / A2 / A3 / BCDE] - PCA <br> groups | 4 | -2369.84 | 8 | -4520.00 |
| RunB A1 / A2 / A3 / BCD/ E]- DAPC <br> groups | 5 | -1122.35 | 7 | -2025.02 |
| R |  |  |  |  |

Figures


Figure S1. (A) Principal components analysis (PCA) as implemented and plotted in LEA package, presenting the projection of all individual geckos on the first two PCs and (B) DAPC as implemented in adegenet on all the Mediodactylus populations.


Figure S2. Maximum Likelihood (ML) tree reconstructed using ddRAD data. Bootstrap (BS) support values from the ML analysis are shown on the branch nodes of the tree. Individual codes follow those in Table S1 with the first two digits representing the map codes of Figure 1.


Figure S3. (A) Structure bar plot ( $K=2$, best supported number of clusters based on the Evanno method) for all Mediodactylus populations belonging to M. kotschyi. (B) Structure bar plots for each one of the Subclades A1, A2, A3 ( $K=2$ or 3 was the best grouping). Individuals are represented as vertical bars along the plot. The Y axis represents the percentage of each individual ( Q value) assigned to a cluster; the height of each color represents the probability of assignment to a genetic cluster. The black vertical lines indicate clade/subclade limits. (C) Sampling locations for the groups of populations as indicated by structure analyses. Colors correspond to the structure bar plots shown in panel B. The dot and rectangular points correspond to the blue and the orange cluster in panel A respectively.


Figure S4. (A) Structure analysis for M. orientalis and M. oertzeni. Individuals are represented as vertical bars along the plot. The Y axis represents the percentage of each individual ( Q value) assigned to a cluster; the height of each color represents the probability of assignment to a genetic cluster. The black vertical lines indicate subclade limits. (B) Sampling locations for the groups of populations as indicated by structure analyses for $M$. orientalis ( $\boldsymbol{L}^{(1)}$ and . oertzeni ( $\bullet$ ). The differentially colored points correspond to the clusters within each species as shown in the bar plots above.


Figure S5. Bootstrap 50\% majority-rule consensus tree from SVDQuartets analysis for eight lineages/species (the second best delimited scheme indicated by BFD*) and five species (as it is the current taxonomy) and their respective distribution on the maps.

## Code used for the analyses

## For the RaxML phylogenetic tree inference

raxml-ng --parse --msa Mediodactylus.phy --model GTR+G \#Compress and store MSA in the binary format and Estimate memory requirements and optimal number of CPUs
raxml-ng --check --msa Mediodactylus.phy --model GTR+G \#check the MSA
raxml-ng --msa *.rba --model GTR+G --threads 44 --seed 2617 --prefix MLTreedata --tree pars\{25\}, rand\{25\} \#Increase the number of starting trees to 50 (the default is 20) to explore the tree space more thoroughly.
raxml-ng --rfdist --tree MLtree --prefix RF \#check if the 50 starting trees were enough for the analysis
raxml-ng --bootstrap --msa *.rba --model GTR+G --prefix boot --seed 2988 --threads 44 --bs-trees 300 \#Bootstrap analysis. Increase the number of replicates to 300 (default is 50 replicates)
raxml-ng --bsconverge --bs-trees boot. raxml. bootstraps --prefix convergence --seed 1234 --threads 4 --bs-cutoff 0.01
raxml-ng --support --tree MLTree. raxml. bestTree --bs-trees boot.raxml.bootstraps --prefix MLwithBoot --threads 44 \#map the bootstrap values onto the best-scoring/best-known ML tree

For the RaxML phylogenetic tree inference using only unlinked SNPs data matrix we used --model GTR+ASC_LEWIS for the Lewis correction, and we run the analysis according to the above mentioned commands.


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[^1]:    \#: Samples used in the Med12_snapp dataset for the SNAPP analysis were selected based on the tree topology and structure barplots.

[^2]:    Abbreviations: df: degrees of freedom.

