Lizards of a different stripe: phylogenetics of the *Pedioplanis undata* species complex (Squamata, Lacertidae), with the description of two new species

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Abstract

The lacertid genus *Pedioplanis* is a moderately speciose group of small-bodied, cryptically-colored lizards found in arid habitats throughout southern Africa. Previous phylogenetic work on *Pedioplanis* has determined its placement within the broader context of the Lacertidae, but interspecific relations within the genus remain unsettled, particularly within the *P. undata* species complex, a group largely endemic to Namibia. We greatly expanded taxon sampling for members of the *P. undata* complex and other *Pedioplanis*, and generated molecular sequence data from 1,937 bp of mtDNA (ND2 and cyt b) and 2,015 bp of nDNA (KIF24, PRLR, RAG-1) which were combined with sequences from GenBank resulting in a final dataset of 455 individuals. Both maximum likelihood and Bayesian analyses recover similar phylogenetic results and reveal the polyphyly of *P. undata* and *P. inornata* as presently construed. We confirm that *P. husabensis* is sister to the group comprising the *P. undata* complex plus the Angolan sister species *P. huntleyi* + *P. haackei* and demonstrate that *P. benguelensis* lies outside of this clade in its entirety. The complex itself comprises six species including *P. undata*, *P. inornata*, *P. rubens*, *P. gaerdesi* and two previously undescribed entities. Based on divergence date estimates, the *P. undata* species complex began diversifying in the late Miocene (5.3 ± 1.6 MYA) with the most recent cladogenetic events dating to the Pliocene (2.6 ± 1.0 MYA), making this assemblage relatively young compared to the genus *Pedioplanis* as a whole, the origin of which dates back to the mid-Miocene (13.5 ± 1.8 MYA). Using an integrative approach, we here describe *Pedioplanis branchi* sp. nov. and *Pedioplanis mayeri* sp. nov. representing northern populations previously assigned to *P. inornata* and *P. undata*, respectively. These entities were first flagged as possible new species by Berger-Dell’mour and Mayer over thirty years ago but were never formally described. The new species are supported chiefly by differences in coloration and by unique amino acid substitutions. We provide comprehensive maps depicting historical records based on museum specimens plus new records from this study for all members of the *P. undata* complex and *P. husabensis*. We suggest that climatic oscillations of the Upper Miocene and Pliocene-Pleistocene era in concert with the formation of biogeographic barriers have led to population isolation, gene flow restrictions and ultimately cladogenesis in the *P. undata* complex.

Key Words

Biogeography, molecular phylogeny, phylogenetics, southern Africa, species description, taxonomy
Introduction

The Lacertidae is a large family of Old World lizards with a center of diversity in southern Africa, particularly in xerothermic habitats (Arnold et al. 2007; Mayer and Pavlicev 2007; Garcia-Porta et al. 2019). Overall, this family has a conservative general morphology and the degree of homoplasy and convergence in external features is very high, especially in predominantly arid-zone lineages (Borsuk-Bialynicka et al. 1999; Hipsley and Müller 2017). With 13 currently recognized species, *Pedioplanis* Fitzinger, 1843 is the most diverse genus of lacertid lizards in southern Africa (Conradie et al. 2012), with taxa occupying semi-arid to highly xeric habitats, including coastal fynbos, Nama and succulent Karoo, grassland, savannah, and hard-packed gravel plains and deserts (Branch 1998; Makokha et al. 2007). Nevertheless, previous studies have detected additional, novel lineages among *Pedioplanis* species that await resolution and formal description (Mayer and Berger-Dell´mour 1987; Makokha et al. 2007; Conradie et al. 2012). Members of the *Pedioplanis undata* species complex are autochthonous to Namibia and South Africa and have proven particularly taxonomically challenging (Mayer and Berger-Dell´mour 1987; Makokha et al. 2007). All are similar in size (adult SVL 40–56 mm) and overall body form, but exhibit differences in dorsal patterning and coloration, arrangement of scales in the lower eyelid, and geographic distribution (Mayer and Berger-Dell´mour 1987; Branch 1998; Makokha et al. 2007).

*Pedioplanis undata* was originally described as a single widespread species, *Lacerta undata* Smith, 1838, and one year later transferred to the genus *Eremias* Fitzinger, 1834 by Duméril and Bibron (1839). Three non-nomotypical subspecies, *Eremias undata rubens*, *E. u. gaerdesi* and *E. u. inornata* (the last originally described as full species, *Eremias inornata* Roux, 1907) were subsequently recognized based on morphological differences (Boulenger 1921; Mertens 1954, 1955). Mayer and Berger-Dell´mour (1987) used allozyme data to delineate a total of seven geographically coherent forms including northern and southern varieties of *P. u. undata* and *P. u. inornata*. They provisionally elevated *P. gaerdesi* and *P. rubens* to full species, and later used the same methods to describe *P. husabensis*, which together with *P. u. undata* and *P. u. inornata* resulted in five putative members of the complex (Berger-Dell´mour and Mayer 1989). Morphologically-based phylogenetic analyses by Arnold (1991) recovered the same five named taxa at the species level, although their results suggested that the *P. undata* complex was rendered paraphyletic by *P. namaquensis* and *P. benguelensis*.

Multiple recent molecular phylogenies have confirmed the monophyly of the *P. undata* species complex (including *P. husabensis*, as sister to all remaining species; Makokha et al. 2007; Conradie et al. 2012), however, some interspecific relationships remain unresolved. Maximum likelihood analyses place *P. rubens* in a clade with *P. undata* that is sister to all remaining species within the complex (Makokha et al. 2007), whereas Bayesian analyses suggest that the former is more deeply nested within the complex (Makokha et al. 2007; Conradie et al. 2012). The monophyly of *P. inornata* has also been an ongoing subject of debate. Makokha et al. (2007) identified two moderately divergent (ND2 = 7–8%; 16S rRNA = 3–4%; RAG-1 = 1–2%) lineages within the species, a west-central Namibian clade and a southern Namibian/Northern Cape (South Africa) clade. *Pedioplanis gaerdesi* was recovered as the sister to the former, rendering *P. inornata* paraphyletic (Makokha et al. 2007). These results corroborate Mayer and Berger-Dell´mour’s (1987) report of two morphologically and geographically distinct forms of *P. inornata*, however, no formal taxonomic changes were made due to limited geographic sampling. Finally, the status of *P. undata* has also previously come into question, as Mayer and Berger-Dell´mour (1987) identified northern and southern forms of the species, yet the most recent phylogenetic analyses detected no such divergent clades given their limited sampling (Makokha et al. 2007; Conradie et al. 2012).

We present the most comprehensive phylogenetic assessment yet of the *P. undata* species complex in order to resolve interspecific relationships among described species, investigate the status of previously recognized unnamed lineages, and elucidate the geography of species diversification within the complex. We compiled a new and extended multi-locus dataset and reconstructed the phylogeny of the genus *Pedioplanis* as a whole with a greatly increased sample set for each member of the *P. undata* species complex throughout their ranges, particularly in Namibia.

Materials and methods

Field sampling

The majority of specimens included in this study were collected by the authors, under permitted fieldwork in Namibia, South Africa and Botswana. Specimens were euthanized, fixed in 10% neutral buffered formalin, and stored in 70–75% ethanol (EtOH). Prior to fixation, tail, thigh muscle or liver tissue was preserved in 95% EtOH for subsequent DNA sequencing. Specimens have been deposited in the Museum of Comparative Zoology at Harvard University (MCZ), the California Academy of Sciences (CAS), the National Museum of Namibia in Windhoek (NMMNW), and the Museum für Naturkunde in Berlin (MfN, ZMB). All locality data were recorded by means of handheld GPS devices in decimal degrees (DD) using the WGS 84 reference coordinate system.

Molecular phylogeny

Genomic DNA was extracted using the Qiagen DNAeasy Kit. PCR amplification was performed on an Eppendorf Mastercycler gradient thermocycler using primer pairs
obtained from published sources (see Suppl. material 4: Table S1 for primer pairs). Molecular sequence data were generated for two mitochondrial genes, NADH dehydrogenase subunit 2 (ND2) and cytochrome b (cyt b), and for three nuclear genes, recombination activation gene 1 (RAG-1), prolactin receptor (PRLR), and kinesin family member 24 (KIF24). Nuclear genes were sequenced for only a subsample of individuals representing unique mitochondrial haplotypes (KIF24: n = 229; PRLR: n = 214; RAG-1: n = 244). PCR products were visualized using 1.5% agarose gels and then purified with a homemade magnetic bead solution (Rohland and Reich 2012). Cycle Sequencing was performed using the BigDye Terminator v3.1 Cycle Sequencing Kit (Applied Biosystems, Foster City, CA, USA) and purified with an additional magnetic bead cleanup before being sequenced on an ABI 3730xl DNA analyzer. Sequences were assembled and edited in Geneious v9.1.8; initial sequence alignments were made using MUSCLE v3.8.31 (Edgar 2004) before adjusting by eye. Sequence alignments were used to calculate average uncorrected pairwise genetic distances (p-distance) using the distance matrices generated in Geneious v9.1.8.

Sequences were generated for 428 individuals, including all 13 currently recognized species of Pedioplanis and eight outgroup taxa (Nucras aurantiacus, N. holubi, N. tessellata, Heliobolus lugubris, Ichnotropis capensis, Meroles knoxii, M. reticulatus, M. suborbitalis). Based on recent molecular phylogenies (Mayer and Pavlicev 2007; Engleder et al. 2013) the sister pair Nucras + Heliobolus were used to root the tree. Within the P. undata species complex, the number of individuals for which sequences were generated are as follows: 18 P. husabensis (14 localities, see Kirchhof et al. 2017), 71 P. undata (28 localities), three P. rubens (2 localities), 110 P. inornata (58 localities), 48 P. gaerdesi (37 localities). Additional Pedioplanis sequences from the genes ND2 (n = 45) and RAG-1 (n = 40) were downloaded from GenBank, providing thorough representation of each species across its known geographic range (see Suppl. material 1 for complete list of tissue specimen and GenBank sampling).

Phylogenetic analyses using Maximum likelihood (ML) and Bayesian inference (BI) were performed on the CIPRES Science Gateway v3.3 (Miller et al. 2010). Separate analyses were performed on individual genes and on the concatenated sets of mitochondrial and nuclear genes separately. Best-fitting models of evolution were identified using PartitionFinder v2.1.1 (Lanfear et al. 2016) in which partitioning schemes were selected using the Bayesian Information Criterion (BIC) score for each analysis. All ML partitions were run under the GTR+Γ model of evolution using RAxML v8.1.24 (Stamatakis 2014) for 1,000 rapid nonparametric bootstrap replicates; BI partitions were run under the models listed in Suppl. material 5: Table S2. MrBayes v3.2.6 (Ronquist and Huelsenbeck 2003) was used to perform all Bayesian phylogenetic analyses and run for 50,000,000 generations, sampling every 10,000 generations; convergence of the Markov chains was assessed by eye using Tracer v1.7 (Rambaut et al. 2018) and the initial 25% of trees were discarded as burn-in.

Morphological data and analysis

Morphological comparisons among members of the P. undata complex were made following initial phylogenetic analysis. A diversity of mensural and meristic measures was recorded for 120 specimens including the type series of the new species described herein, and additional specimens representative of major clades revealed by the phylogenetic analyses. In order to obtain adequate sampling to capture morphological variation in each clade, we added non-genotyped adult individuals (≥ 40 mm snout–vent length) of P. undata sensu stricto (n = 6) collected from the Oanob Dam (near Rehoboth) population, and of P. rubens (n = 9) from the Klein Waterberg population, to the morphological analyses (see Appendix 1 for a complete list of specimens used for the morphological analyses). Morphological data included the following measurements in mm (for bilateral features only right side values were recorded, unless damaged): Snout–vent length (SVL) – snout tip to the anterior edge of the cloaca; body length (BL) – anterior edge of cloaca to posterior edge of collar; collar-snout length (CSL) – posterior edge of collar to snout tip; head length (HL) – posterior edge of the occipital to snout tip; maximum head width (HW); lower jaw length (JL) – anterior edge of jaw bone to tip of lower jaw; length of the fourth finger (FiL) and fourth toe (ToL); inter-limb length (IL) – length between axillary and inguinal regions; hind limb (tibia) length (HiL); forearm length (FoL) – elbow to wrist; tail length (TaL) – posterior edge of cloaca to tail tip. All measurements were collected using Mitutoyo digital calipers (to 0.1 mm). Comparative data for Angolan taxa were obtained from Conradie et al. (2012).

Features of head and body scalation recorded were: prefrontal contact (PF); number of small granules in front of supraoculars touching frontal and prefrontal (G); number of rows of granules between supraoculars and supraciliary (RG); number of supraciliaries (Su); interparietal-occipital contact (IO); number of infralabials (IF); number of gular scales in a straight line between the chin symphysis and median collar plate (Gu); number of subdigital lamellae on fourth toe (SuL); number of femoral pores on right leg (left if right leg was damaged or pores obscured) (Fe). Features of dorsal color and pattern were also recorded. All recorded morphological data for the type specimens are reported in the species descriptions below and listed in Tables 1 and 2; summaries of the meristic and mensural character data for members of the P. undata species complex can be found in Suppl. materials 2, 3 (Suppl. material 2: mensural character data for P. undata species complex voucher specimens; Suppl. material 3: meristic character data for P. undata species complex voucher specimens).
X-rays and Micro-computed (micro-CT) scans of specimens in the type series designated in this study were prepared at the California Academy of Sciences and Museum für Naturkunde, respectively, to obtain presacral vertebrae counts (PV). All x-ray images were taken using a Faxitron Cabinet X-ray System Model 43855C (Faxitron, Tucson, AZ, USA) at 45 kV for 60 s. Micro-CT scans were obtained using a Phoenix nanotom X-ray/tube at 85 kV and 120 or 150 μA, generating 1440 projections with 750 ms per scan, with differing kV-settings depending on the respective specimen size. Effective voxel size, i.e., resolution in three-dimensional space, ranged from 12.8 to 14.3 μm. The cone beam reconstruction was performed using the datos|x-reconstruction software (GE Sensing & Inspection Technologies GMBH phoeni(x-ray datos|x 2.2) and the data were visualized in VG Studio Max 3.1. Micro-CT scans have been deposited on the MN data repository (https://doi.naturkundemuseum.berlin/10.7479/4jg6-ae46) and are available on the MorphoSource website (http://morphosource.org). X-ray images are also available on MorphoSource (See Appendix 1 for DOIs).

Due to limited sample sizes we ran our morphological analyses on a dataset combining data for adult males and females of each species; all statistical analyses were performed in R (R Core Team 2019). Meristic variables were tested for normal distribution using the Shapiro-Wilk normality test as part of the R package (Wickham 2016). We refrained from additional allometric effects were taken into account when mensural characters were compared. We built bivariate linear models using the raw measurements of each character in relation to the respective individual’s SVL and compared the slopes of the regression lines visually using the ggplot2 package (Wickham et al. 2021) and for homogeneity of variance with Levene’s test. We excluded the categorical variables prefrontal contact (PF) and interparietal-occipital contact (IO) and number of rows of granules between supraoculars and supraciliaries (RG) from this analysis. Allometric effects were taken into account when mensural characters were compared. We built bivariate linear models using the raw measurements of each character in relation to the respective individual’s SVL and compared the slopes of the regression lines visually using the ggplot2 package (Wickham et al. 2021) and for homogeneity of variance with Levene’s test. We excluded the categorical variables prefrontal contact (PF) and interparietal-occipital contact (IO) and number of rows of granules between supraoculars and supraciliaries (RG) from this analysis. Allometric effects were taken into account when mensural characters were compared. We built bivariate linear models using the raw measurements of each character in relation to the respective individual’s SVL and compared the slopes of the regression lines visually using the ggplot2 package (Wickham et al. 2021) and for homogeneity of variance with Levene’s test. We excluded the categorical variables prefrontal contact (PF) and interparietal-occipital contact (IO) and number of rows of granules between supraoculars and supraciliaries (RG) from this analysis.

Results

Phylogenetic analysis

Final alignments of the in-group Pedioplanis taxa were as follows: ND2: 905 base pairs (429 variable, 379 parsimony informative); cyt b: 1,032 base pairs (444 variable, 363 parsimony informative); KIF24: 484 base pairs (72 variable, 50 parsimony informative); PRLR: 544 base pairs (73 variable, 46 parsimony informative); RAG-1: 987 base pairs (124 variable, 74 parsimony informative).

There were no conflicts in the topologies between the BI and ML analyses and both retrieved generally high nodal support (bootstraps – BS > 70%; posterior probabilities – PP > 0.95) throughout their respective trees. Separate analyses among the nuclear genes revealed no conflicts, though individually they provided little resolution and were thus concatenated into a single alignment. Within the concatenated nuclear tree (not shown) P. benguelensis, P. husabensis and P. rubens were found to be monophyletic with strong support (P. benguelensis: PP = 1.0, BS = 100%; P. husabensis: PP = 1.0, BS = 100%; P. rubens: PP = 0.85, BS = 100%). However, all supra-specific clades within the P. undata complex were weakly supported by our nuclear data, and relationships among them remained largely unresolved.

Separate analyses were performed on the individual and concatenated mitochondrial genes ND2 and cyt b. Each analysis resulted in a well-resolved tree and no taxonomically relevant topological conflicts were revealed. Our concatenated mitochondrial analyses (Fig. 1) recovered a strongly supported monophyletic Pedioplanis as sister to Meroles + Ichnotropis (PP = 1.0, BS = 98%). Within Pedioplanis, P. burchelli + P. laticeps are sister to all remaining congeners (PP = 1.0, BS = 100%). Pedioplanis breviceps and P. lineoeccellata are recovered as sister taxa (PP = 1.0, BS = 100%) and are, in turn, sister to P. namaquensis plus all remaining Pedioplanis (PP = 1.0, BS = 100%). Pedioplanis benguelensis, P. husabensis and (P. huntleyi + P. haackei) are successively more closely related to the P. undata complex (PP = 1.0, BS = 99–100%). Within the clade containing the four recognized species of the P. undata species complex (P. undata, P. gaerdesi, P. inornata, P. rubens) we recovered a total of six highly divergent lineages (Figs 1, 2). In all analyses P. undata appears polyphyletic, comprising two geographically distinct lineages, one from northern Namibia (P. undata “North” sensu Mayer and Berger-Dell’mour 1987) and another that occurs in central Namibia from the Khomas Highlands, south into parts of the northern Hardap Region and west into the Erongo Region (P. undata “South” sensu Mayer and Berger-Dell’mour 1987). The two forms were not recovered as sister taxa and are highly divergent from one another (average uncorrected p-distance + SD: cyt b: 12% + 0.9; ND2: 14% + 4.1). The southern lineage is deeply nested within the complex, whereas P. undata “North” is sister to all other P. undata complex lineages (PP = 1.0, BS = 100%). Pedioplanis rubens was recovered nested within the P. undata species complex and sister to the remaining clades (PP = 1.0, BS = 99%). Individuals that superficially resemble P. inornata or P. gaerdesi from central-west Namibia in the Erongo region form a clade, (P. inornata “Central” sensu Makokha et al. 2007) and are sister to P. gaerdesi sensu stricto (PP = 1.0, BS = 100%), collected largely from north of the Ugab River. A second P. inornata clade (P. inornata “South”) was represented by specimens from as far north as the Kuiseb River and south into the Northern Cape Province of South Africa. These P. inornata lineages are moderately divergent

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from each other (average uncorrected p-distance + SD): cyt b: 7.7% + 0.5; ND2: 8.8% + 4.3). A sister relationship between *P. undata* “South” and *P. inornata* “South” is recovered in all analyses (PP = 0.99, BS = 91%), although without support in the case of nuclear data. A sampling map for sequenced tissues of *P. husabensis* and all clades recovered within the *P. undata* species complex is depicted in Fig. 1, and within these clades, major geographic structuring is depicted in Fig. 2a, b.

**Morphological analysis**

The final morphological dataset (adults only) contained 13 *P. undata* “North” (7 males, 6 females), 20 *P. inornata* “Central” (15 males, 5 females), 22 *P. inornata* “South” (15 males, 7 females), 21 *P. rubens* (4 males, 6 females), 21 *P. gaerdesi* (8 males, 13 females), and 11 *P. undata* “South” (2 males, 9 females). We ran Shapiro-Wilk normality tests on all meristic characters and found that only Gu data was normally distributed (W = 0.9838, p = 0.2782) and met the assumption of homogeneity of variance (Levene’s test, F = 1.8392, p = 0.113), whereas all other characters were not normally distributed: G (W = 0.95389, p = 0.0018), Su (W = 0.56256, p < 0.001), IF (W = 0.74376, p < 0.001), SuL (W = 0.95704, p = 0.0044), Fe (W = 0.93945, p < 0.001) (see Suppl. material 6: Table S3 for detailed results).

Using ANOVA or Kruskal-Wallis rank sum tests (and subsequently applied Wilcoxon rank sum test) we found significant differences only in the number of subdigital lamellae on the fourth toe (SuL; Kruskal-Wallis chi-squared = 11.603, p = 0.0407) and the number of femoral pores on the leg (Fe; Kruskal-Wallis chi-squared = 24.992, p < 0.001) between the species of the *P. undata* complex. Since none of these differences contribute meaningfully to the diagnosis of the new taxa we present the data in the Suppl. material 7: Table S4 for reference, but refrain from overinterpreting them in this present study.

For the difficult to distinguish species pairs *P. undata* “North” and *P. undata* “South”, as well as *P. inornata* “Central” and *P. inornata* “South”, we assessed mensural characters in an allometric context by comparing their linear regressions (see Suppl. material 8: Fig. S1, Suppl. material 9: Fig. S2 for linear regression comparisons of mensural characters). However, in our dataset, the average SVL of adult *P. inornata* (“Central”) and *P. inornata* (“South”) were significantly different, with *P. inornata* (“Central”) being significantly smaller (Wilcoxon rank sum test, W = 122, p = 0.01401), preventing a meaningful comparison of all other size-related body measurements by this method. While the similar-sized *P. undata* (“North”) and *P. undata* (“South”) (Wilcoxon rank sum test, W = 78, p = 0.728), might differ with regard to forelimb length (FoL) and lower jaw length (JL) (see Suppl. material 8: Fig. S1), this may be due to other factors (e.g., sampling bias, potential sexual size dimorphism), but limited sample sizes prevented the use of multivariate analyses for clarification.
Systematics

Our phylogenetic results support the existence of two highly divergent, unnamed entities within the \textit{Pedioplanis undata} species complex, each previously recognized as a putatively distinct, but unnamed lineage of a recognized taxon (\textit{P. undata} and \textit{P. inornata}). The results of the phylogenetic analyses further indicate that these lineages exhibit species-level genetic differentiation and, on this basis, we elevate the \textit{P. undata} “North” clade and the \textit{P. inornata} “Central” clade to full species, which we describe here using the general lineage concept of species (de Queiroz 1998).

\textit{Pedioplanis undata} “South”/\textit{P. undata} “North”

Mayer and Böhme (2000) examined the type material of \textit{Lacerta undata} Smith, 1838 and noted that the rediscovered syntype specimens in the National Museums of Scotland collection were, in fact, representatives of the taxon long associated with the name \textit{Pedioplanis lineoocellata pulchella} (Gray, 1845) and now, based on the findings of Edwards (2013), considered as \textit{P. lineoocellata} (Duméril & Bibron, 1839). They proposed to conserve the current usage of names within \textit{Pedioplanis} by calling on the ICZN in Case 3085 to set aside the original type designation and recognize the designation of a neotype for \textit{L. undata}. Opinion 1992 of the ICZN (Anonymous 2002) accepted this solution and thus the neotype was fixed as Naturhistorisches Museum Wien (NMW) 31886 from “near Windhoek, Namibia.” This links the name \textit{P. undata} to the \textit{P. undata} “South” clade. The assignment of the name \textit{Pedioplanis undata} to the more restricted, southern clade is notable in that the predominant concept of the species based on previous studies of \textit{P. undata}...
is largely affiliated with populations and representative specimens from the more broadly distributed northern clade (Branch 1998; Makokha et al. 2007; Conradie et al. 2012). In these cases the presence of bold dorsal stripes was considered diagnostic for the species, however based on our results this character is now largely applicable to the unnamed northern clade, and is exhibited inconsistently and with variation among true \textit{P. undata}. No other nomina are currently within the synonymy of \textit{Pedioplanis undata} and none are available for application to the \textit{P. undata} “North” clade, which we describe as:

\textit{Pedioplanis mayeri} sp. nov.

http://zoobank.org/1EDAC87A-1018-4A70-82BF-B9A0040D0E86
Figs 3–5

**Holotype (Fig. 3).** Adult ♀; MCZ-R193179 (field number MCZ-A28704); Namibia, Erongo Region, Omaruru District, at Farm Omandumba, ca. 1 km W of house (-21.49786, 15.62630, 1238 m above sea level (a.s.l.)); collected 15 June 2014 by Jackie L. Childers, Aaron M. Bauer, William R. Branch, Matthew Heinicke and Johan Marais. Holotype and part of type series to be transferred to the National Museum of Namibia.

**Paratypes.** n = 8 (all specimens are adults unless otherwise noted); three ♀: CAS 214643, MCZ-R185872, ZMB 89349; five ♂: CAS 214645, MCZ-R185870, MCZ-R190213, MCZ-R193125, ZMB 89350 collected from various localities in the Kunene Region of northern Namibia • CAS 214643, 59 km W of Kamanjab (-19.64871, 14.33984, 1163 m a.s.l.); collected 1 June 2000 by Aaron M. Bauer • CAS 214645; same collection data as for proceeding • MCZ-R185870; 35 km S of Epupa Falls on Okangwati Rd. (-17.26083, 13.21194, 1035 m a.s.l.); collected 14 August 2007 by Aaron M. Bauer, Johan Marais, Ross A. Sadlier, and Stuart V. Nielsen • MCZ-R185872; N Okangwati on Epupa Falls Rd. (-17.40361, 13.15861, 1140 m a.s.l.); collected 27 November 2011 by Aaron M. Bauer • MCZ-R193125; collected at the type locality on 13 June 2014 • ZMB 89349; subadult ♀; along the C35 road, ca. 2 km N of Fransfontein (-20.19976, 15.01453, 1097 m a.s.l.); collected 27 November 2011 by Aaron M. Bauer • MCZ-R193125; collected at the type locality on 13 June 2014 • ZMB 89350; along the C43 road, 29 km N of Palmwag (-19.62736, 13.87391, 1124 m a.s.l.); collected 15 August 2014 by Sebastian Kirchhof • ZMB 89350; along the C43 road, 29 km N of Palmwag (-19.62736, 13.87391, 1124 m a.s.l.); collected 15 August 2014 by Sebastian Kirchhof. Elevation data for the holotype and paratypes was obtained using the GPS or, if not available, from Google Earth (earth.google.com) using georeferenced GPS coordinates from the collecting localities.

**Diagnosis.** Distinguished from \textit{P. lineoocellata}, \textit{P. lateiceps} and \textit{P. burchelli} by having 10 longitudinal ventral scale rows (vs. 12 or more). It is distinct from \textit{P. gaerdesi}, \textit{P. benguelensis}, \textit{P. husabensis}, \textit{P. namaquensis} and \textit{P. breviceps} in possessing a semi-transparent lower eyelid with a brille formed by 2–4 enlarged scales (brille formed by a single scale in \textit{P. benguelensis} and \textit{P. gaerdesi}, lower eyelid with eight opaque scales in \textit{P. husabensis} and...
opaque and scaly in *P. breviceps* and *P. namaquensis*). Dorsal patterning is characterized by the presence of five, usually bold dark brown to black, straight-ridged dorsal stripes, distinguishing it from *P. rubens* (dorsum and tail uniform red-brown to brick red, lacking conspicuous markings with only a hint of a slightly brighter dorso-lateral line on each side), *P. inornata*, *P. gaerdesi* and *P. branchi* sp. nov. (dorsum may be light to dark gray becoming gradually more reddish towards the tail, possessing dark and/or light beige-yellowish speckling but lacking distinct longitudinal elements), *P. haackei* (only three dark dorsal stripes), and *P. undata* (dorsal striping bold or not, may be reduced with pale longitudinal elements or even a single middorsal stripe restricted to the nape).

It is further distinguished from *P. haackei* by typically having a smaller number of granules anterior to the first supraocular (9–16 vs. 12–32), from *P. huntleyi* in having a larger number of granules anterior to the first supraocular (9–16 vs. 7–13), and from *P. undata* in possessing a greater maximum number of granular scales anterior to the supraoculars (9–16 in *P. mayeri* sp. nov. versus 8–13 in *P. undata*), and a greater maximum number of femoral pores on a single leg (12–16 in *P. mayeri* sp. nov. versus 11–14 in *P. undata*) (see Suppl. material 7: Table S4 and Suppl. material 3). *Pedioplanis mayeri* is similar in most characteristics to *P. huntleyi*, which is restricted to Angola, but the latter species is typified by the restriction of the five dark dorsal stripes to the anterior half of the trunk, a condition found in only some *P. mayeri*. No single morphological character unambiguously distinguishes the allopatric species *P. mayeri*, *P. undata* and *P. huntleyi*. We therefore provide several diagnostic characters based on the mitochondrial gene ND2 in order to supplement the morphological data. *Pedioplanis mayeri* may be distinguished from all other *Pedioplanis* species in being characterized by having the amino acid histidine instead of a tyrosine at base pair 61 due to a codon change at that position. It also has the amino acid phenylalanine at base pair 223, instead of a threonine (*P. benguelensis*, *P. breviceps*, *P. haackei*, *P. huntleyi*, *P. husabensis*, *P. laticeps*, *P. lineooccellata*, *P. namaquensis*), an alanine (*P. burchelli*), or a serine (*P. branchi* sp. nov., *P. gaerdesi*, *P. inornata*, *P. rubens*, *P. undata*) due to a codon change at that position. Finally, it has the amino acid threonine due to a codon change at base pair 568, instead of a leucine (*P. rubens*), a valine (*P. haackei*, *P. laticeps*, *P. undata*), or an isoleucine (*P. benguelensis*, *P. branchi* sp. nov., *P. breviceps*, *P. burchelli*, *P. gaerdesi*, *P. husabensis*, *P. namaquensis*), an alanine (*P. burchelli*), or a serine (*P. branchi* sp. nov., *P. gaerdesi*, *P. inornata*, *P. rubens*, *P. undata*) due to a codon change at that position.
obliquely rectangular in shape, and strongly keeled; 24 presacral vertebrae.

**Color in alcohol.** Dorsum of crown gray-tan with small speckles of varying size on head shields; dorsum of snout tinged with reddish-brown. Lateral surface of head cream-colored with minute gray speckles on loreal scales, supralabials and infralabials. Granules bordering lower eyelid window white. Color and pattern of temporal region confluent and consistent with those on trunk. Trunk dorsum with three bold black stripes each bordered on either side by whitish-cream stripes that extend to the sacrum, becoming indistinct on the tail; middorsal black stripe divided anteriorly by a whitish–cream stripe that extends from the occiput to the level of the forelimb insertion, narrowing from a width of five granular dorsal scales to two from anterior to posterior. Below the lateralmost of these pale stripes, which originates on the temporal, lies a thick darkly pigmented line to which is fused the coalescence of a series of 9R/8L irregularly sized ocelli with pale gray-blue centers comprising 6–14 granules, each surrounded by a brown to black ring. This in turn is bordered ventrally by a diffuse white line which begins at the corner of the mouth, passes through the ear and continues to the groin. Below this line the flanks transition to the immaculate pearly white of the venter. Dorsal surfaces of forelimbs mottled brown with margins of enlarged scales pale; dorsal surfaces of hind limbs medium brown, enlarged scales on preaxial surface similar to those of forelimb, postaxial surfaces bearing scattered cream-colored spots, each 1–3 granular scales in extent; palms and soles gray with scattered diffuse pinkish to orange markings. Tail light brown, densely speckled with darker markings which continue the bold lateral black stripes of the body dorsum as a pair of narrow dark brown stripes running along the medial–most keels of the tail for at least three quarters of its length.

**Variation within the type series.** Measurements for the type series are summarized in Table 1 and mensural (n = 13) and meristic (n = 17) character data based on measurements of additional museum vouchers can be found in Suppl. material 2, 3, respectively. Paratypes have similar scalation to the holotype except as follows: in MCZ-R185872 the prefrontals do not contact and are instead divided by a granule; in MCZ-R193125 the interparietal and occipital are separated by a granule; in all paratype specimens the occipital is not fused with the parietals; the number of granules in the group preceding the anterior supraocular varies from nine to 15; in CAS 214645 and ZMB 89349 there is one row of granules dividing the anterior supraocular from the supraciliaries (usually two); in CAS 214645 and MCZ-R185870 there are 7R and 7L infralabials, respectively (usually six); granular scales in a straight line between the symphysis of chin shields and the median collar plate varies from 27 to 32; midbody scale rows vary from 57 to 64; subdigital lamellae under the fourth toe range from 25 to 28; unilateral

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**Figure 3.** *Pedioplanis mayeri* sp. nov. adult female holotype (MCZ-R193179) from Farm Omandumba in the Erongo Region of Namibia. (a.) Dorsal and (b.) ventral views of the whole body; scale bars indicate 10 mm; (c.) dorsal, (d.) ventral) and (e.) lateral views of the head; scale bar indicates 2.0 mm.
Table 1. Morphological data for *Pedioplanis mayeri* sp. nov. type series. CAS = California Academy of Sciences, MCZ = Museum of Comparative Zoology, ZMB = Museum für Naturkunde. Abbreviations for character values are as follows: C = In contact; S = Separated by granule; SF = Separated by frontal and frontonasal; INC = Tail incomplete. See Materials and Methods for character abbreviations and descriptions; Midbody scale rows (MB) were not recorded for all individuals.

| Type Status | Holotype | Paratype | Paratype | Paratype | Paratype | Paratype | Paratype | Paratype | Paratype |
|-------------|----------|----------|----------|----------|----------|----------|----------|----------|----------|
| Museum No.  | MCZ-R 193179 | CAS 214643 | MCZ-R 185872 | ZMB 89349 | CAS 214645 | MCZ-R 185870 | MCZ-R 190213 | MCZ-R 193125 | ZMB 89350 |
| Sex         | Female | Female | Female | Female | Male | Male | Male | Male | Male |
| SVL (mm)    | 45.5 | 40.7 | 47.0 | 43.3 | 44.0 | 51.4 | 49.2 | 43.5 | 121.5 |
| Tal. (mm)   | 116.7 | NA (INC) | 108.5 | 105.4 | 107.3 | 124.9 | 131.8 | 96.4 | 121.5 |
| PF          | C     | S      | C      | S      | C      | C      | C      | C      | C      |
| G           | 16    | 9      | 11     | 15     | 14     | 10     | 11     | 10     | 13     |
| RG          | 2     | 2      | 2      | 1      | 1      | 2      | 2      | 2      | 2      |
| Su          | 6     | 6      | 6      | 6      | 6      | 6      | 6      | 6      | 6      |
| IO          | C     | C      | C      | C      | C      | C      | C      | S      | C      |
| IF          | 6     | 6      | 6      | 6      | 6      | 6      | 6      | 6      | 6      |
| Gu          | 28    | 28     | 29     | 28     | 31     | 27     | 29     | 31     | 29     |
| SUL         | 27    | 25     | 25     | 26     | 27     | 28     | 27     | 26     | 26     |
| Fe          | 15    | 13     | 14     | 13     | 14     | 14     | 14     | 14     | 14     |
| PV          | 24    | 25     | 24     | 25     | 24     | 24     | 24     | 25     | 24     |
| MB          | –     | –      | 61     | –      | 57     | 64     | 64     | 58     | –      |

**femoral pores number 13 to 14 (right side only); and pre-sacral vertebrae number either 24 or 25.**

**Coloration in life (Fig. 4).** Dorsal coloration and pattern varied considerably across examined specimens. Dorsum of head gray-tan, sometimes possessing dark speckles of varying size; dorsum usually bearing three bold black, medium-brown or gray stripes bordered on each side by white stripes; dorsal stripes often terminate at the tail base or begin to fade into a gray and reddish hue midway towards the tail base; dorsal stripes may be absent entirely, with individuals possessing a uniformly gray dorsum, sometimes with dark speckling. All individuals examined possess yellow-centered ocelli or yellow spots along the flanks, usually bordered ventrally by a white stripe and dorsally by a black stripe (or with the dark rims of the ocelli confluent with such a stripe); lateral spots or ocelli vary in size from ca. 0.5 to 1.0 mm in diameter, in some cases smaller spots may merge with the white ventral stripe; anterior surface of the thighs and outermost ventral plates sometimes tinged or speckled with orange; limbs and feet more or less uniformly grayish (front), reddish-brown to brown, or speckled with light or black dots or blotches that may form bars; tail usually bright brick or orange-red, in some cases dark speckling may be present on the dorsal surface of the tail.

**Coloration in alcohol.** Pattern as described above though dark stripes may fade to dark gray, yellow lateral spots may appear cream white or gray-blue, and red coloration usually present on the dorsum of the midbody and tail may become a faded orange-pink or may fade to a faint, golden-brown hue.

**Distribution.** *Pedioplanis mayeri* sp. nov. is endemic to northern Namibia and occurs from south of the Kunene River and east of the Namib Desert along the eastern side of the escarpment, thence throughout the eastern Kunene Region, entering the northeastern parts of the Erongo Region and east through the Otjozondjupa Region, reaching at least as far east as Oshikango (TM17028) in the north, Gobabis (Omaheke Region) in the south-east, and Nauchas in the south. It does not enter the Kalahari dune fields, and is possibly absent from the Khomas Hochland, where it is replaced by *P. undata* (see Fig. 6 for map of locality records).

**Etymology.** The specific epithet is a patronym formed in the genitive singular honoring our friend and colleague, the Austrian lacertid specialist Werner Mayer (1943–2015), who first recognized the distinctiveness of his namesake species and whose contributions to the study of *Pedioplanis* have been seminal.

**Natural history.** *Pedioplanis mayeri* is widespread in northern Namibia and inhabits Tropical & Subtropical Grasslands, Savannas & Shrublands and Deserts & Xeric Shrublands biomes (following the classification of the WWF; Olson et al. 2001), but it is absent from true desert. Within the occupied biomes, *P. mayeri* inhabits Namibian savanna woodlands, Kalahari xeric savanna, Kalahari Acacia-Baikiaea woodlands and Angolan Mopane woodlands (terminology follows Olson et al. 2001). Here, the species is active on sandy clay soils and more compact, hard soils, rocky flats and slopes, in dense grassy patches as well as in more open areas interspersed with shrubs (see Fig. 5 for images of characteristic habitats). In rocky areas and stony slopes individuals were found sleeping under stones. The majority of the species’ range comprises Namibia’s arid regions with a median annual precipitation of 279 mm (61–559 mm) and a median annual temperature average of 21.1 °C (18–23.5 °C) (all climate data taken from Fick and Hijmans 2017).
females with up to four eggs were found in December
and February, and small juveniles (SVL = 27 mm) ap-
peared in mid-February (Kirchhof et al. 2014). Sym-
patric ground-living diurnal lizards include Trachylepis
damarana, Meroles squamulosus, Heliobolus lugu-
bris, Gerrhosaurus flavigularis, and Agama anchietae,
among others.

Conservation. The species occurs across a wide area
of Namibia in which potential threats from agriculture
and mining are scattered and localized. Local populations
occur in protected areas within Etosha National Park and
Waterberg Plateau Park as well as in many local conserv-
vancies. Applying IUCN criteria, we consider P. mayeri
to be Least Concern.
Figure 6. Topographic maps showing locality records for all the species of the Pedioplanis undata species complex and *P. husabensis*: (a.) *P. husabensis* (b.) *P. mayeri* sp. nov. (c.) *P. rubens* (d.) *P. branchi* sp. nov. (e.) *P. gaerdesi* (f.) *P. undata* (g.) *P. inornata*. Black dots represent museum vouchers from various museum collections, those with a white center were sequenced and included in the phylogenetic analyses. Museum voucher information was collected from the California Academy of Sciences San Francisco/USA, Carnegie Museum of Natural History Pittsburgh/USA, Ditsong National Museum Pretoria/South Africa, Iziko South African Museum Cape Town/South Africa, Museum für Naturkunde Berlin/Germany, Museum of Comparative Zoology Harvard University/USA, Museum of Vertebrate Zoology Berkeley/USA, Natural History Museum of Los Angeles County/USA, Bayworld Museum Port Elizabeth/South Africa, National Museum of Namibia Windhoek/Namibia, Zoological Research Museum Alexander Koenig Bonn/Germany, in parts via the Global Biodiversity Information Facility (GBIF; https://www.gbif.org/).
**Pedioplanis inornata** “South” / *P. inornata “Central”*

The type series of *Eremias inornata* Roux, 1907 comprises eight specimens collected from “Orange-Fluss, Kl.-Namaqualand” [the Orange River, Little Namaqualand, Northern Cape, Province, South Africa]. Syntype ZMA11049 (Zoological Museum Amsterdam) was subsequently designated as the lectotype (Daan and Hillenius 1966). The ZMA collections have subsequently been incorporated into those of the Naturalis Biodiversity Center in Leiden (RMNH). The geographic origin of the specimens and the relatively detailed description and accompanying plate, clearly tie the name-bearing lectotype of *Pedioplanis inornata* to the *P. inornata “South”* clade. No other nominas are currently within the synonymy of *Pedioplanis inornata* and none are available for application to the *P. inornata “Central”* clade, which is here described as:

*Pedioplanis branchi* sp. nov.

http://zoobank.org/7FDB1C1FF-BEB4-464A-88C5-6CDD19ED50F2

Figs 7–9

**Holotype (Fig. 7).** Adult ♂; CAS 214788 (field number AMB 6551); Namibia, S of Karibib at Junction of Road D1914 and Road D1952, Karibib District, Erongo Region (-22.27038, 15.57471, 1080 m a.s.l.); collected 8 June 2000 by Aaron M. Bauer. Holotype and part of type series to be transferred to the National Museum of Namibia.

**Paratypes.** n = 7 (adults); (two ♂: ZMB 89310, ZMB 89311; five ♂: CAS 214790, CAS 214792, CAS 214794, ZMB 89305, ZMB 89316) collected from various localities located in the Erongo Region of northwestern Namibia; CAS 214790: S of Karibib at junction of Rd. D1914 and Rd. D1952 (-22.27281, 15.57815, 1075 m a.s.l.); collected 8 June 2000 by Aaron M. Bauer; CAS 214792; same collection data as for proceeding; CAS 214794; same collection data as for proceeding; ZMB 89305; 22 km SW of Uis along C35 (-21.42419, 14.76271, 884 m a.s.l.); 10 August 2014; ZMB 89316; 22 km SW of Uis along C35 (-21.34513, 14.75676, 884 m a.s.l.); 23 January 2013 • ZMB 89310; Farm Friedhelm Sack (-22.52699, 15.54487, 709 m a.s.l.); 14 October 2014 • ZMB 89311; same collection data as for proceeding; all ZMB specimens collected by Sebastian Kirchhof. Elevation data for the holotype and paratypes was obtained using the GPS, or, if not available, from Google Earth (earth.google.com) using georeferenced GPS coordinates from the collecting localities.

**Diagnosis.** Distinguished from *P. lineocellata*, *P. laticeps* and *P. burchelli* by having 10 longitudinal ventral scale rows (vs. 12 or more). It is distinct from *P. benguelensis*, *P. gaerdesi*, *P. breviceps*, *P. namaquensis* and *P. husabensis* in usually possessing a semi-transparent lower eyelid with a brille formed by 2–4 scales (brille formed by a single scale in *P. benguelensis* and *P. gaerdesi*, lower eyelid with eight opaque scales in *P. husabensis* and opaque and scaly in *P. breviceps* and *P. namaquensis*); in some rare cases *P. branchi* sp. nov. may possess a single transparent scale in lower eyelid, those individuals can be distinguished from *P. benguelensis* and *P. gaerdesi* by color and dorsal patterning (see below). Dorsal patterning is characterized as being uniformly gray from the mid-back towards the head with a reddish hindbody (posterior half of body) and with a series of pale to bright yellow spots or ocelli on lower flanks, distinguishing it from *P. rubens* (dorsum and tail uniform red-brown to brick red, lacking conspicuous markings with only a hint of a slightly brighter dorso-lateral line on each side), *P. mayeri* sp. nov., *P. haackei*, *P. huntleyi*, *P. undata* (dorsum contains bold stripes or other longitudinal elements), *P. gaerdesi* (never with lateral ocelli, and speckled with very small black or light dots) and *P. inornata* (spots on flanks are typically pale green, not yellow). It can further be distinguished from all other *Pedioplanis* (except *P. inornata*) in typically having a pair of distinct dark markings on the face, one through the eye and extending onto the supralabials directly below and one more posterior, near the corner of the mouth. The new species is significantly smaller than *P. inornata* (*P. branchi* mean adult SVL = 44.4 mm, max. 49.1 mm, versus *P. inornata* mean adult SVL 47.3 mm, max. 54.0 mm for specimens sampled here; to 56.0 mm elsewhere [Bauer and Shea 2006]). The maturity of specimens was confirmed by their mostly or completely fused long bone epiphyses. We also provide several diagnostic characters based on the mitochondrial gene ND2. *Pedioplanis branchi* sp. nov. can be distinguished from all other members of the *P. undata* species complex except *P. laticeps* and *P. undata* in being characterized by having the amino acid histidine instead of tyrosine at base pair 359 due to a codon change at that position. It can also be distinguished from all other *Pedioplanis* species except *P. laticeps* in possessing a thymine at base pair 300, rather than an adenine (*P. benguelensis*, *P. burchelli*, *P. gaerdesi*, *P. haackei*, *P. huntleyi*, *P. husabensis*, *P. inornata*, *P. mayeri* sp. nov., *P. namaquensis*, *P. rubens*, and *P. undata*) or a cytosine (*P. breviceps* and *P. lineocellata*).

**Description of holotype.** Body relatively slender (SVL 47.5 mm); interlimb distance 16.8 mm; femur 9.5 mm; tibia 8.7 mm; humerus 5.4 mm; forearm 5.7 mm; body length 26.9 mm from groin to collar; collar-snout length 19.3 mm; fourth finger length 4.7 mm; fourth toe length 10.4 mm; head narrow and elongated (head width 55% of head length) with slight constriction at base of rostrum and narrow, pointed snout (width at rear of frontonasal 2.2 mm, width at front of eye 4.4 mm); head length 11.8 mm; head width 6.6 mm; lower jaw length 10.2 mm; eye–ear distance 4.4 mm; eye–nostril distance 4.0 mm; 1.1 mm between the nostrils; complete original tail 92.3 mm, with epidermis missing from posterior portions. Rostral semicircular; contacting nasals; nasals in contact medially, unaunched relative to rostral and frontonasal; postnasal contacts nasal, frontonasal, anterior loreal, and enters the nostril; nostrils circular in shape; two loreals, anterior loreal half the length of the posterior loreal; two
preoculars; prefrontals in median contact; frontal large, with a narrow posterior projection, bordered anteriorly by the prefrontals and posteriorly by the frontoparietals; frontoparietals in broad medial contact; interparietal in contact anteriorly with both frontoparietals, laterally with the parietals, and posteriorly with the occipital; occipital trapezoidal in shape; two supracroculars, both in medial contact with the frontal and frontoparietals, preceded anteriorly by 8R/8L granules (on left side two granules in contact with prefrontal and one in contact with frontal; on right side two granules in contact with prefrontal and three in contact with frontal); single row of granules dividing anterior supraciliary from supraciliaries, and double row of granules dividing posterior supraocular from supraciliaries; two small scales between last supraciliary and parietal; six supraciliaries on each side, the anteriormost longest; lower eyelid with transparent bristle formed of two larger, black-edged scales, with a row of five smaller scales beneath; five supralabials anterior to subocular and three supralabials posterior to subocular, on both sides; subocular bordering the lip, its lower edge shorter than its upper; 6R/6L infralabials; first infralabial in contact with the second infralabial, the first chin shield, and the mental; four enlarged pairs of chin shields, with the first three in medial contact and the posterior-most largest; no enlarged temporals; tympanum sunk; no scales projecting significantly past margin of ear opening; enlarged narrow scale at anterodorsal margin of ear opening. 30 gular scales in a straight line between symphysis of chin shields and median collar plate; collar free, comprising nine enlarged plates (median kite-shaped, projecting posteriorly) and extending onto side of neck as a crease that terminates midway up the lateral side; dorsal scales small, juxtaposed, granular, without keels, lateral scales larger towards ventrals; 49 rows of granular scales around the midbody; ventral plates in 10 longitudinal and 31 transverse rows (from collar to groin); plates of the outermost rows squarish, ventral rows usually twice as wide as long; single transverse row of ventrals across chest just behind collar longer than broad; nine enlarged precloacal scales, irregular in shape, median ones larger; centralmost enlarged precloacal in contact with six other enlarged precloacals; scales on upper surface of forearm large, smooth, and overlapping, without keels; scales on lower surface of forearm with a series of enlarged plates, at least twice the width of scales on upper forearm; scales on upper surface of tibia rhombic, subimbricate and distinctly keeled, below with a series of enlarged plates; scales on upper surface of femur granular and smooth with enlarged scales along the anterior margin; 12R/12L femoral pores; subdigital lamellae under fourth toe 26R/28L; scales on tail obliquely rectangular in shape, and strongly keeled.

**Color in alcohol.** Dorsum of head gray-tan with distinct black speckles of varying size on the head shields; lateral surface of head with three bold, black vertical bars, one at the anterior margin of eye, one originating on the ventral eyelid and one just posterior to the orbit, all

*Figure 7. Pedioplanis branchi sp. nov., adult male holotype (CAS 214788) from south of Karibib in the Erongo Region of Namibia. (a.) Dorsal and (b.) ventral views of the whole body; scale bars indicate 10 mm; (c.) dorsal, (d.) ventral) and (e.) lateral views of the head; scale bar indicates 2.0 mm.*
Coloration (in alcohol). In comparison to the life coloration, the row of spots along the flanks that appear yellow in life appear gray-blue in preserved specimens and may be conspicuous or faint, and the reddish-brown color of the posterior body and tail appears darker and more grayish.

Distribution. *Pedioplanis branchi* sp. nov. is endemic to the Erongo Region in Namibia. Its range stretches from just south of the Swakop River in the South, where it occurs in parapatry with *P. hasubensis*, through the pro-Namib, to the Ugab River and the Brandberg in the north and Mount Erongo and Otjimbingwe in the east, probably occurring in parapatry with *P. mayeri* all along its northeastern border (see Fig. 6 for map of locality records).

Etymology. The specific epithet is a patronym formed in the genitive singular honoring our friend and colleague, the British-born South African herpetologist, William Roy Branch (1946–2018), in recognition of his many contributions to African herpetology and in remembrance of many happy trips in the field together.

Natural history. *Pedioplanis branchi* inhabits Namibian savanna woodlands and the gravel plains and rocky outcrops in the Namib Desert (Deserts & Xeric Shrublands biome; Olson et al. 2001). The species is active on harder substrates, gravel and bare rock (including marble, granite, basalt), mostly with only sparse vegetation cover and only rarely forages on softer sandy substrates (e.g. in small washes on slopes) or in more grassy patches (Fig. 9). Median annual precipitation within the distribution range of *P. branchi* reaches a maximum of only 187 mm (minimum = 16 mm, median = 142 mm), while mean annual temperatures are similar to those in

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Table 2. Morphological data for *Pedioplanis branchi* sp. nov. type specimens. CAS = California Academy of Sciences, MCZ = Museum of Comparative Zoology, ZMB = Museum für Naturkunde. Abbreviations for character values are as follows: C = In contact; S = Separated by granule; SF = Separated by frontal and frontonasal; RGN = Tail regenerated. See Materials and Methods for character abbreviations and descriptions.

| Type Status | Holotype | Paratype | Paratype | Paratype | Paratype | Paratype | Paratype | Paratype |
|-------------|----------|----------|----------|----------|----------|----------|----------|----------|
| Museum No.  | CAS 214788 | ZMB89310 | ZMB89311 | CAS 214790 | CAS 214792 | CAS 214794 | ZMB 89305 | ZMB 89316 |
| Sex         | Male     | Male     | Male     | Male     | Male     | Male     | Male     | Male     |
| SVL (mm)    | 46.6     | 48       | 42.5     | 50.0     | 40.0     | 49.0     | 49.0     | 44.1     |
| TaL (mm)    | 92.3     | 88.4     | 111.2    | 107.0    | 107.0    | 99.0     | 99.0     | 94.1     |
| C           | SF       | C        | C        | C        | C        | C        | C        | SF       |
| PF          | 8        | 12       | 14       | 11       | 11       | 14       | 11       | 9        |
| G           | 1        | 2        | 2        | 2        | 2        | 2        | 2        | 1        |
| Su          | 6        | 5        | 6        | 6        | 6        | 7        | 7        | 6        |
| IO          | C        | C        | C        | C        | C        | C        | C        | C        |
| IF          | 7        | 6        | 6        | 7        | 6        | 6        | 7        | 6        |
| Gu          | 30       | 30       | 26       | 30       | 26       | 30       | 30       | 28       |
| SuL         | 26       | 29       | 28       | 26       | 25       | 25       | 27       | 29       |
| Fe          | 11       | 10       | 12       | 12       | 10       | 13       | 13       | 11       |
| PV          | 24       | 25       | 24       | 24       | 24       | 24       | 23       | 24       |

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Dorsal coloration and pattern varied considerably across examined specimens. Dorsalum of head ranges from uniformly gray to distinctly speckled with black, sometimes with bold dark blotches or vertical bars. Body dorsal gray anteriorly and red-orange posteriorly, sometimes bearing light or dark spots that can be arranged in longitudinal rows, or are irregularly scattered, or may form horizontal bars or chevrons across the posterior dorsum near the tail base; belly uniform cream. The flanks may be patterned with light or dark irregular markings, sometimes forming vertical bars or chevrons, and are always lined with a more or less distinct longitudinal row of yellow spots or yellow-centered ocelli bordered by a broken ring of black granules, which range from 0.5–1.0 mm in diameter. The anterior surface of the thighs sometimes tinged with yellow; limbs and feet uniformly gray (front) or reddish-brown (hind), or speckled with light or black dots or blotches that may form bars. Dorsum of tail is orange-red, often with dense, black speckles.
the distribution of *P. mayeri* (median = 21 °C, range 19.4–22.5 °C). The species occupies lower elevations, from 77 m a.s.l. near the coast to a maximum of ca. 1190 m a.s.l. near the southeasternmost corner of the distribution along the Swakop River Canyon (median = 968 m a.s.l.). The largest recorded individual was a male (SVL = 50 mm). Breeding season appears to be similar to other southern African *Pedioplanis* spp. in spring, gravid females were found in December. Sympatric diurnal lizards include, *Trachylepis sulcata*, *Rhoiptopus boultoni*, *Agama planiceps*, *Matobosaurus validus*, and *Varanus albigularis*, among others.

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**Figure 8.** Life photographs of representative individuals of *Pedioplanis branchi* sp. nov. highlighting color pattern variation within the species. (a.) Lateral and (b.) dorsal whole body images of an adult specimen from the Chuos Mountains (Chuosberg) (SK376.2014; not included in study), note the dense, dark speckling on the dorsum, and bold dark lateral markings. (c.) Dorsal and (d.) lateral whole body images of an adult specimen (SK402.2014; not included in study) from the northward extension of the Swakop River Canyon, note the more uniformly gray and red dorsum lacking speckling and fainter yellow spots along the flanks. Lateral (e.) head and (f.) whole body images of an adult specimen from the northward extension of the Swakop River Canyon; note the black, vertical bar beneath the lower eyelid, the bold, dark lateral markings on the body, and the distinct longitudinal row of yellow spots along the flanks.

**Figure 9.** Habitat of *Pedioplanis branchi* sp. nov. (a.) Collection site for ZMB 89310 (Farm Friedhelm Sack, Erongo Region, Namibia) from the extended Swakop River Canyon on a marble/calc silicate slope with sparse grass cover and isolated shrubs. (b.) Collection site for ZMB 89612 (not included in study) from a small hill slope in the Tsiseb conservancy in the Pro-Namib, 20 km south of the Brandberg (visible in the background) depicting a sparsely vegetated scree slope with a *Commiphora* shrub.
**Conservation status.** Although *P. branchi* has an extent of occurrence below 20,000 km² it occurs in an area with low human population density, generally low intensity land use, and a high proportion of suitable habitat. Populations occur in protected areas within Namib-Naukluft Park and Tsisseb Conservancy, and possibly Dorob National Park. Applying IUCN criteria we consider *P. branchi* to be Least Concern.

**Discussion**

*Pedioplanis* phylogeny and the recognition of two new species within the *P. undata* species complex

In order to resolve the systematics and biogeography of the *Pedioplanis undata* species complex we analyzed a new and greatly extended multi-locus molecular dataset comprising 455 samples, from all 13 currently recognized species of *Pedioplanis* and eight outgroup taxa. The results from our phylogenetic analyses show that interspecific relationships within the genus *Pedioplanis* are largely congruent with previous studies, but with several notable exceptions. We recover with strong support the phylogenetic affinities of three species that have heretofore remained problematic, *P. benguelensis*, *P. husabensis* and *P. rubens*. Within the *P. undata* species complex our results support the recognition of two highly divergent lineages and multiple instances of polyphyly.

Historically, *P. benguelensis* was thought to be closely related to *P. namaquensis* due to similarities in dorsal patterning (Bocage 1867; Laurent 1964; Makokha et al. 2007). More recently, Conradie et al. (2012) recovered *P. benguelensis* as sister to the Angolan species pair *P. huntleyi* + *P. haackei* and the *P. undata* complex, with *P. husabensis* falling outside of the entire grouping; however these results did not receive high nodal support in their analyses.

Our results confirm the sister relationship between the Angolan taxa and members of the *P. undata* complex, however we recover *P. husabensis* and *P. benguelensis* as sequentially more distant sister taxa to this clade (Fig. 1b, c). Previous phylogenetic studies have obtained conflicting topologies for the placement of *P. rubens*. Maximum likelihood analyses by Makokha et al. (2007) found that *P. rubens* forms a clade with *P. undata* (actually *P. mayeri* sp. nov.) that is sister to *P. gaerdesi* + *P. inornata*. However, their BI analysis recovered *P. rubens* as nested within the *P. undata* complex, a result corroborated by Conradie et al. (2012) and by this present study (Figs 1c, 2a).

We identify two instances of polyphyly within the *P. undata* complex. First, *P. inornata* as previously defined comprises two divergent lineages, one that occurs in the eastern and southern parts of Namibia, and in the Northern Cape of South Africa (= *P. inornata*) and another comprising individuals from central-western Namibia (= *P. branchi* sp. nov.). The two lineages are not each other’s closest relatives; rather, *P. branchi* sp. nov. is sister to *P. gaerdesi*. This result was corroborated by the nuclear data, in which the concatenated phylogeny recovered these clades as being distinct from one another. Interestingly, some *P. branchi* sp. nov. individuals possess a single (instead of two) large semi-transparent scale (“brille”) in the lower eyelid, which caused them to be identified as *P. gaerdesi* in the field given that this scale character is currently used to diagnose the species (Conradie et al. 2012). We suggest that this morphological character alone is inadequate for diagnosing species within the *P. undata* complex. Makokha et al. (2007) identified the presence of a *P. inornata* “Central” clade (= *P. branchi* sp. nov.), however, due to limited sampling they were unable to identify its geographic extent. Here we provide evidence that this clade is limited to an arid region north of the Kuiseb River and south of the Ugab River. These results corroborate Mayer and Berg-er-Dell’mour’s (1987) hypothesis that within *P. inornata* there exist two geographically and morphologically distinct forms, one restricted to west-central Namibia that possesses a gray forebody and red-orange hindbody with yellow lateral spots (*P. inornata*—“Central” = *P. branchi* sp. nov.; body coloration depicted in Fig. 8), and a more broadly distributed form in southern Namibia that is characterized by brownish or reddish coloration with greenish lateral spots, *P. inornata*—“South” (= *P. inornata*).

Another equally striking result is the discovery that *P. undata* comprises two genetically distinct and polyphyletic lineages. The first clade, *P. undata* “North” (= *P. mayeri* sp. nov.) includes individuals primarily from northern Namibia, (although a single sample was found as far south as Nauchas), and is sister to all remaining members of the *P. undata* species complex. The second lineage, *P. undata* “South” (= *P. undata* sensu stricto), is more deeply nested within the complex and is closely related to *P. inornata*, a result recovered by both the nuclear and mtDNA data.

**Diversification and geographic context**

Geographic isolation of populations through vicariance can result from various events, including both geological (formation of rivers, valleys, mountains, dunes) and climatic (drastic environmental change resulting from cooling, warming, or changes in precipitation resulting in habitat expansion or contraction). Southern Africa has experienced a complex paleoclimatic and geological past. In particular, the climatic oscillations of the Upper Miocene and the Pliocene-Pleistocene era are proposed to have been influential in the evolution and diversification of southern African fauna (e.g. Bauer 1999; Matthee and Flemming 2002; Montgelard and Matthee 2012).

In addition, multiple potential geographical barriers have been hypothesized to be responsible for vicariance and subsequent cladogenesis. These include the Great Escarpment, the sand seas of the Namib Desert, shifting Kalahari sands, and rivers such as the Kuiseb, Zambezi,
and Orange (e.g. Lamb and Bauer 2000; Matthee and Flemming 2002; Scott et al. 2004; Bauer et al. 2006).

The current distribution ranges of members of the *P. undata* complex (see Fig. 6) do not indicate that the aforementioned barriers today have a substantial effect in restricting gene flow. For example, the sister taxa *P. inornata* and *P. undata* both occur east and west of the escarpment and instead possess a north-south border (*P. undata* also occurs both north and south of the Kuiseb), and the sister pair *P. branchi* and *P. gaerdesi* both occur west of the escarpment with a north-south border not obviously coincident with any geographical barrier. Nevertheless, during the estimated times of major cladogenesis events within the genus (ca. 8–3.8 million years ago (MYA); Makokha et al. 2007; Garcia-Porta et al. 2019) these potential physical barriers of gene flow may have been less porous and only later, in concert with the Plio-Pleistocene/Holocene climatic changes, may have permitted passage.

During the Early Miocene conditions in southern Africa were wetter and warmer than today. Since then southern Africa has undergone stepwise cooling and drying. Diekmann et al. (2003) suggested that arid periods occurred in southern Africa between 11.6 and 10.7 MYA and between 9.7 and 7.7 MYA. The region experienced a third major cooling step with increased aridification at the end of the Pliocene (around 2.8 MYA) (Van Zinderen Bakker 1986; Partridge 1998; Diekmann 2003). This coincided with a major uplifting of the subcontinent (the African superswell; Lithgow-Bertelloni and Silver 1998) resulting in major changes of erosional patterns due to rejuvenation and reactivation of drainage systems and increased fluvial activity (Partridge et al. 2006). These drastic climatic changes may have resulted in newly available open habitats for *Pedioplanis* species occupying more arid habitats like *P. branchi*, facilitating subsequent radiation and then likely to range contractions and fragmentation among ancestral *Pedioplanis* spp. with subsequent cladogenesis, scenarios that have been suggested for e.g. southern African *Pachydactylus* geckos (Bauer 1999; Heinicke et al. 2017) and lacertids generally (Garcia-Porta et al. 2019).

The major uplift together with changes in drainage systems during less arid periods at the end of the Pliocene seems to coincide well with the split of *P. inornata* and *P. undata*, indicating that the now ephemeral Kuiseb and/or Swakop River might have been effective impediments to gene flow between these two sister taxa in the past. Today, in the east there are populations of *P. undata* occurring south of the mostly dry Kuiseb, indicating that gene flow between the two taxa is possible. For *P. mayeri* (the basal taxon of the clade) and *P. gaerdesi*, the escarpment has been suggested to form a geographical barrier (Bauer et al. 1993). During our study we detected a narrow area of potential sympathy north of Palmwag and – based on museum specimens – near Sesfontein, suggesting that while the escarpment might have formed a barrier in the past it no longer does.

Evolution of body coloration and patterning within the *P. undata* species complex

The *P. undata* species complex has historically proven taxonomically challenging, in part due to the fact that most species have been distinguished chiefly by their body coloration and patterning (Branch 1998; Conradie et al. 2012). Our phylogenetic results show both that divergent taxa share similar color and patterning characteristics, and that intraspecific variation can be considerable.

Dorsal striping may be a plesiomorphic condition within the *P. undata* complex given that *P. mayeri*, the sister taxon to the remainder of the complex, as well as the Angolan outgroup species *P. huntleyi* and *P. haackei*, are all typically striped. The absence of dorsal striping in *P. rubens*, *P. inornata* and *P. gaerdesi* would suggest that stripes were lost in lineages that subsequently invaded more open habitats (further discussed below) whereas the ancestral, striped condition was retained within *P. undata*.

Members of the *P. undata* species complex are chiefly allopatrically distributed in regions of Namibia that are ecologically distinct; we suggest that apparent color variation both among and within species may be a result of differences in habitat and associated selective pressures. Uniformly and plainly colored taxa including *P. branchi*, *P. gaerdesi*, *P. inornata* and *P. rubens* all occur in arid regions where they are camouflaged on bare, sandy or rocky substrates that lack dense vegetation (Arino et al. 2012; Stone and Thomas 2013). In the case of *P. rubens*, which has a very restricted range in the Waterberg Plateau, individuals are reddish and are found exclusively on red sandstone, whereas *P. gaerdesi* and *P. branchi* are found in similar habitats in the pro-Namib and adjacent rocky outcrops of the Namib, respectively, and each possesses a light gray dorsum, especially anteriorly. Likewise, striped patterning in *P. mayeri* and northern populations of *P. undata* may also be a consequence of shared similarities in habitat type. Both taxa occur in inland localities in northern Namibia, a region that receives 200–500 mm of rainfall annually (Rohde and Hoffman 2012) and in habitats predominantly characterized by annual and perennial grasses (Arino et al. 2012; Wang et al. 2012). Among squamates the correlation between dorsal striping and the use of grassy substrates has been hypothesized to reflect the use of disruptive coloration in order to reduce the likelihood of detection by predators during escape (Wolf and Werner 1994; Farallo and Forstner 2012).

This hypothesized link between ecology and body coloration in the *P. undata* species complex may be further demonstrated by the pronounced intraspecific variation evident within the sister taxa *P. inornata* and *P. undata*. For example, within *P. inornata*, typical specimens possess a predominantly gray dorsum, however we encountered localized red morphs on red, hard-packed soils ca. 60 km west of Mariental, and dark morphs on basalt rock piles just north of the same town. *Pedioplanis undata* collected in northern localities that are characterized by substantial grass cover (Harris et al. 2014) exhibit bold dorsal stripes.
that appear nearly indistinguishable from those exhibited by *P. mayeri* (see Fig. 4 for images of dorsal color patterning typical of the species), whereas individuals collected just south of the Khomas Hochland, where grasses are more sparsely distributed, possess faint gray dorsal stripes; furthermore, individuals collected near the Gobabeb Training and Research Centre on vegetationless, schist rock outcrops had almost patternless, nearly uniformly gray dorsums.

Taken together, our phylogenetic results and field observations strongly suggest that ecology is a significant evolutionary driver of body coloration and patterning within the *P. undata* species complex, as has also been demonstrated previously for e.g. *P. lineocellata* (Branch 1998; Edwards 2013; Childers 2015).

**Conclusion**

The results of our study indicate that species diversity within Namibian *Pedioplanis* is greater than previously thought and that this diversity is concentrated within the *P. undata* species complex. These results settle the long-standing hypothesis originally put forth by Mayer and Berger-Dell’mour (1987) on the existence of additional lineages within the complex and confirm that it comprises six species rather than the four traditionally considered.

Based on present species distributions and divergence date estimates derived from previous phylogenetic studies on African lacertids (Makokha et al. 2007; Garcia-Porta et al. 2019) we propose several biogeographic scenarios that may have led to cladogenesis in the *P. undata* species complex. We hypothesize that species diversification beginning at ca. 8–3.8 MYA may have been precipitated by major climatic changes in combination with the formation of biogeographic barriers across the subcontinent, causing once widespread populations to become isolated and restricting gene flow until cladogenesis ultimately ensued. Among these isolated populations, differences in habitat and associated natural selection pressures may have produced phenotypic differences among them including the evident diversity in body color and patterning between species.

This study highlights the need for dense geographic sampling among broadly distributed groups that have radiated recently and rapidly, as is the case for the genus *Pedioplanis*. Furthermore, without fine-scaled microhabitat data it is difficult to accurately assess ecological differences and to test our hypothesis of convergent evolution in this group, and we recommend future researchers prioritize these critical data.

The majority of organismal research in Namibia has so far mainly focused on the Namib Desert region in the west (Herrmann and Branch 2013), where species discovery has accelerated over the past decade due in part to the implementation of molecular phylogenetics. However, this study highlights the need for increased biodiversity surveys of the central interior portion of Namibia, as our results suggest that squamate endemicity in this region may have been previously underestimated.

**Acknowledgements**

We thank the permit issuing authorities in Angola, South Africa and Namibia for allowing the collection of specimens within their respective jurisdictions. Curators and collections managers of the institutions cited in Tables 1 and 2 and Suppl. material 1 are thanked for their generous access to specimens and provision of tissue samples and data. We are especially grateful to Carol Spencer (MVZ), Lauren Scheinberg (CAS), José Rosado (MCZ), Frank Tillack and Mark-Oliver Rödel (MN) for their assistance with specimen loans and access to collection facilities. Luke and Ursula Verburt, Johan Marais, Mathew Heinicke, William Branch, Tomas Kleinert, Johannes Penner, Rodrigo Megia Palma, Laura Zamorano, Ammon Corl, Donald Miles, Lee Fitzgerald, Christy Hipsey and H. Berger-Dell’mour provided essential field assistance. Thank you, Kristin Mahlow and Johannes Müller (both MN), for micro-CT scans of the ZMB specimens and logistic support. Thank you to Elisa Yang and Tomo Yoshi- no (both MVZ) for assistance with GenBank data accessioning. AMB was supported by NSF grant EF1241885, subaward 13–0632. SK was supported by the German Academic Exchange Service (DAAD) and the Elsa-Neumann-Stipendium des Landes Berlin (Humboldt University of Berlin, Germany).

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Pedioplanis undata complex
Appendix 1

Voucher specimens used for morphological analyses.

CAS = California Academy of Sciences, MCZ = Museum of Comparative Zoology, ZMB = Museum für Naturkunde. Specimens in bold were not sequenced; X-rayed or CT scanned specimens are denoted with superscript (micro-CT scans available on the MiN Data Repository, DOI: 10.7479/4ig6-ae46; X-ray images available on MorphoSource.com, DOI: 10.17602/M2/M350312, M350317, M350322, M350327, M35036, M350352, M350362, M350367, M350372, M350377, M350384, M350391, M350396, M350401, M350406, M350411, M350416,
M350421, M350427, M350432, M350437, M350442.
Type specimens are denoted with asterisks (*** = holotype; * = paratype). All raw data is available upon request from the corresponding author (JLC).

Pedioplanis branchii sp. nov. – CAS 126221*, 173423*, 173443*, 173446, 214787*, 214788**, 214789, 214790*, 214792*, 214793*, 214794**; MCZ-R184162 (x-ray only); ZMB 89302, 89303, 89304, 89305, 89307, 89308, 89309, 89310***, 89311**, 89312, 89313, 89314, 89315, 89316**, 89317, 89318, 89319, 89320, 89321, 89613.

Pedioplanis gaerdesi – CAS 206964, 214599, 214729, 214730, 214741, 214742, 214743, 214744, 214745, 214746; MCZ-R184294, 184959, 185875, 185883, 185885, 185887, 188237; ZMB 89328, 89329, 89330, 89331, 89332, 89612.

Pedioplanis inornata – CAS 193404, 200012, 200039, 200096, 201873, 203522; MCZ-R183772, 184767, 185890, 193103, 193104, 193105, 193293, 193311, 193370; ZMB 89333, 89334, 89336, 89337, 89338, 89339, 89341.

Pedioplanis mayeri sp. nov. – CAS 214643*, 214644*, 214645*, 224010, 224013; MCZ-R185870*, 185872*, 188223, 190213*, 193125*, 193178*, 193179**, ZMB 80391, 89349**, 89350*, 89624, 89625.

Pedioplanis rubens – ZMB 84748, 89351, 89615, 89616, 89617, 89618, 89619, 89620, 89621, 89622, 89623.

Pedioplanis undata – MCZ-R188247, 188248, 193272, 193276, 193376; ZMB 91631, 89626, 89627, 89628, 89629, 89630, 89631.

Supplementary material 1

Sampling list for all specimens used in phylogenetic analyses

Authors: Jackie L. Childers, Sebastian Kirchhof, Aaron M. Bauer
Data type: excel table
Explanation note: Locality information and GenBank accession numbers are provided for all tissue vouchers used in this study. We also provide the specimen ID numbers and mtDNA clade assignments for members of the Pedioplanis undata species complex that are depicted in Fig. 2a, b. Institutional abbreviations for catalog numbers are as follows: CAS = California Academy of Sciences, MCZ = Museum of Comparative Zoology, NHMW = Naturhistorisches Museum Wien, PEM = Bayworld (Port Elizabeth, SA), ZMB = Museum für Naturkunde. Collector and associated field number abbreviations are as follows: ABA-ABD-ABE = Werner Mayer, AMB = Aaron M. Bauer, JM = Johan Marais, JVV = Jens V. Vindum, KTH = Krystal Tolley, MB-MBUR = Marius Burger, MH = Michael Cunningham, N = Anamarija Zagar, RBH = Raymond B. Huey, SK = Sebastian Kirchhof, SVV = Stuart V. Nielson, USH = University of Stellenbosch Herpetology, WC = Werner Conradie. AMB, MCZ-A, and MCZ-Z specimens lacking institutional registration numbers are currently awaiting cataloging at the National Museum of Namibia.

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Link: https://doi.org/10.3897/zse.97.61351.suppl1

Supplementary material 2

Mensural data for P. undata species complex specimens examined.

Authors: Jackie L. Childers, Sebastian Kirchhof, Aaron M. Bauer
Data type: excel table
Explanation note: Minimum and maximum values (mm) for each species are reported for each character, see Materials and methods for character abbreviations and descriptions. Only adults were examined, and only individuals with intact tails were used to obtain tail length values (TaL); see Appendix 1 for a list of all specimens examined.

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Link: https://doi.org/10.3897/zse.97.61351.suppl2

Supplementary material 3

Meristic and scalation data for P. undata species complex specimens examined.

Authors: Jackie L. Childers, Sebastian Kirchhof, Aaron M. Bauer
Data type: excel table
Explanation note: Abbreviations for character values are as follows: C = In Contact; S = Separated by granule; SF = Separated by frontal and frontonasal; SP = Separated by parietals. Data were collected from juvenile and adult specimens. See Materials and methods for character abbreviations and descriptions; see Appendix 1 for a list of all specimens examined.
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Link: https://doi.org/10.3897/zse.97.61351.suppl3

Supplementary material 4

Table S1. Primer pairs

Authors: Jackie L. Childers, Sebastian Kirchhof, Aaron M. Bauer
Data type: excel table
Explanation note: Primers pairs used in this study; PCR annealing temperatures (°C) are provided.

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Link: https://doi.org/10.3897/zse.97.61351.suppl4

Supplementary material 5

Table S2. Bayesian partitioning schemes

Author: Jackie L. Childers
Data type: excel table
Explanation note: Bayesian partitioning schemes for the concatenated mtDNA (Partition nos. 1–4) and concatenated nDNA (Partition nos. 5–8) datasets based on the results of our PartitionFinder v.2.1.1 analyses, in which best-fitting models of evolution were selected using the Bayesian Information Criterion (BIC) score.

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Link: https://doi.org/10.3897/zse.97.61351.suppl5

Supplementary material 6

Table S3. Summary statistics for select meristic variables for *P. undata* species complex

Author: Sebastian Kirchhof
Data type: excel table
Explanation note: Summarized results of Shapiro-Wilk normality test, Levene’s test, Kruskal-Wallis rank sum test and ANOVA for selected meristic variables for adults (SVL ≥ 40 mm) of all species of the *P. undata* species complex. Columns show character (see Materials and Methods for abbreviations and descriptions), Shapiro-Wilk W (Shapiro_W) and corresponding p-value (Shapiro_p), Levene’s F (Levene_F) and corresponding p-value (Levene_p), Kruskal-Wallis chi-square statistics (KW_chi-squared) and corresponding p-value (KW_p), ANOVA F (ANOVA_F) and corresponding p-value (ANOVA_p) and degrees of freedom (df). Differences are considered significant with p ≤ 0.05 and significance codes are as follows: * = p < 0.5; ** = p < 0.01; *** = p < 0.001.

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Link: https://doi.org/10.3897/zse.97.61351.suppl6

Supplementary material 7

Table S4. Summarized results of Wilcoxon rank sum tests for the meristic characters

Author: Sebastian Kirchhof
Data type: excel table
Explanation note: Summarized results of Wilcoxon rank sum tests for the meristic characters with significant Kruskal-Wallis test results (number of subdigital lamellae SuL and number of femoral pores on right leg Fe). All possible species pairs (species pair) for species of the *P. undata* species complex were tested. Columns show Wilcoxon W (Wilcox_W) and corresponding p-value (Wilcox_p) and significant differences (p ≤ 0.05) are indicated with * and significance codes are as follows: * = p < 0.5; ** = p < 0.01; *** = p < 0.001.

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Link: https://doi.org/10.3897/zse.97.61351.suppl7
Supplementary material 8

Figure S1. Linear relationships with 95% confidence level intervals of mensural characters between *P. mayeri* (= *P. undata* "North") and *P. undata*

Author: Sebastian Kirchhof
Data type: PNG image
Explanation note: Linear relationships with 95% confidence level intervals of mensural characters with snout–vent length (SVL) for adult (SVL ≥ 40 mm) *P. mayeri* (= *P. undata* “North” sensu Mayer and Berg-Dell’mour 1987) and *P. undata*. Linear regression equations are provided when the slope coefficient was significant (p ≤ 0.05), in case of non-significance this is indicated by “not sig.” abbreviations; descriptions of characters are provided in Materials and methods.
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Link: https://doi.org/10.3897/zse.97.61351.suppl8

Supplementary material 9

Figure S2. Linear relationships with 95% confidence level intervals of mensural characters between *P. branchi* (= *P. inornata* "Central") and *P. inornata*

Author: Sebastian Kirchhof
Data type: PNG image
Explanation note: Linear relationships with 95% confidence level intervals of mensural characters with snout–vent length (SVL) for adult (SVL ≥ 40 mm) *P. branchi* (= *P. inornata* “Central” sensu Makokha et al. 2007) and *P. inornata*. Linear regression equations are provided when the slope coefficient was significant (p ≤ 0.05), in case of non-significance this is indicated with “not sig.” abbreviations; descriptions of characters are provided in Materials and methods.
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Link: https://doi.org/10.3897/zse.97.61351.suppl9