Two new species of snail-eating snakes of the genus *Tropidodipsas* (Serpentes, Dipsadidae) from southern Mexico, with notes on related species

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Abstract

We describe two new species of *Tropidodipsas* related to the *T. fasciata* species group as defined by Kofron (1987), and provide morphological and molecular data to support the novelty of both species. A partial molecular phylogeny of the Mexican species of snail-eating snakes (Serpentes, Dipsadidae) is presented, and we discuss evolutionary relationships as supported by our molecular results. We analyze specific relationships of the new species described herein with their closest relatives. We present a distribution map for all species of *Tropidodipsas* and include photographs of living individuals of each species. Finally, we discuss other taxonomic changes based on our molecular phylogeny as well as conservation priorities of the new species.

Resumen

Describimos dos nuevas especies de *Tropidodipsas* relacionadas al grupo de *T. fasciata* definido por Kofron (1987), y presentamos evidencia morfológica y molecular que demuestran que estas especies son nuevas. Presentamos una filogenia molecular de las especies Mexicanas de caracoleras (Serpentes: Dipsadidae), y discutimos las relaciones generales según nuestros resultados. Analizamos las relaciones evolutivas de las nuevas especies con las especies más cercanas. Presentamos además mapas de distribución de todas las especies del género *Tropidodipsas* e incluimos fotografías en vida para cada especie. Finalmente discutimos otros cambios taxonómicos basados en nuestra filogenia molecular, así como prioridades de conservación para las nuevas especies.

Kurzfassung

Wir beschreiben zwei neue Arten von *Tropidodipsas*, die mit der von Kofron (1987) definierten *T. fasciata*-Artengruppe verwandt sind, und liefern morphologische und molekulare Daten, um die Neuheit beider Arten zu unterstützen. Eine partielle molekulare Phylogenie der mexikanischen Arten schneckenfressender Schlange (Serpentes, Dipsadidae) wird vorgestellt, und wir diskutieren evolutionäre Beziehungen, die durch unsere molekularen Ergebnisse gestützt werden. Wir analysieren spezifische Beziehungen der hier beschriebenen neuen Arten zu ihren nächsten Verwandten und besprechen andere taxonomische Veränderungen, die auf unserer molekularen Phylogenie basiert sind. Wir präsentieren eine Verbreitungskarte für alle Arten von *Tropidodipsas*, sowie Fotos von lebenden Exemplaren jeder Art. Abschließend geben wir Schutzprioritäten der neuen Arten an.

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Key Words
conservation, cloud forest, Geophis, Guerrero, Oaxaca, pine-oak woodland, Sierra Madre del Sur

Introduction
The herpetofaunal diversity of southern Mexico in the states of Guerrero and Oaxaca is among the richest in the country (Mata-Silva et al. 2015; Palacios-Aguilar and Flores-Villela 2018). However, the rugged sierras in this region are also some of the least explored areas in the country. This is partially due to the difficulty of accessing these mountain ranges because of the lack of good roads, combined with political and social conflicts and illegal narcotics production. Despite these issues, biologists continue to study these areas, and novel taxa are discovered in the region each year. In the past 15 years, 8 species of snakes as well as a new snake genus have been described from this region (Campbell and Flores-Villela 2008; Pavón-Vázquez et al. 2011; Campbell 2015; Campillo et al. 2016; Campbell et al. 2018; Canseco-Márquez et al. 2018; García-Vázquez et al. 2018; Hernández-Jiménez et al. 2019).

The snake family Dipsadidae is one of the most species-rich, and is distributed throughout the Americas, especially the Neotropics, as well as in southeast Asia (Figueroa et al. 2016; Pyron et al. 2011; Uetz 2021; Zheng and Wiens 2016). Among the Dipsadidae, a subset of genera (“goo eaters”) have specialized in feeding on invertebrate prey: Dipsas Laurenti 1768, Sibon Fitzinger 1826, Sibynomorphus Fitzinger 1843, Tropidodipsas Günther 1858 and Geophis Wagler 1830. Because of their diet specialization on slugs and snails, species of the genera Dipsas, Tropidodipsas and Sibon are referred to as snail-suckers (the name which we will use from now on).

Multiple authors have used morphological characters to study the taxonomy and systematic relationships among these snakes (Peters 1960; Kofron 1980, 1982, 1985a, 1985b, 1987, 1988, 1990; Wallach 1995; Smith et al. 2005), whereas Sheehy (2013), in an unpublished PhD dissertation, incorporated molecular data to resolve some long-standing questions regarding the systematics of the group. Although recent studies have combined morphological and molecular data to address the systematics of some members of the South American species groups of snail-suckers (Harvey 2008; Harvey and Embert 2008; Arteaga et al. 2018), multiple taxonomic questions remain unanswered, particularly regarding Mexican and Central American taxa. Some of these questions concern the generic placement of several North American species (e.g., Sibon sanniolus, Tropidodipsas sartorii species group, T. fischeri, Dipsas gaigeae, D. brevifacies), the validity of the genera in Mexico and Central America, and the definition of some of the wider ranging species (e.g., Tropidodipsas fasciata, T. sartorii).

While conducting herpetofaunal surveys in several mountain ranges in the states of Guerrero and Oaxaca, we obtained multiple individuals of two distinct species of snakes of the genus Tropidodipsas (Serpentes, Dipsadidae) that we could not allocate to any currently described species. After comparing these specimens to other taxa using both morphological and molecular data, we identified them as undescribed species which we describe here. To resolve the taxonomy of our recently discovered species of snail-suckers from southern Mexico, we used nuclear and mitochondrial loci to assess the phylogenetic placement of the new taxa among the snail-suckers genera. We additionally discuss some taxonomic issues with the Mexican species of snail-suckers.

Materials and methods
Taxonomic Sampling
Between 2004 and 2020 we collected multiple specimens of two undescribed species of snail-suckers in the states of Guerrero and Oaxaca, Mexico. Additionally, we collected other species of snail-suckers from across Mexico to serve as comparative material. Currently, thirteen species of snail-suckers are known from Mexico. These thirteen species are distributed among three genera: four are currently assigned to Sibon, two to Dipsas and seven to Tropidodipsas. Tropidodipsas fasciata and T. sartorii have two subspecies each in Mexico and T. fischeri has two subspecies, one of which occurs in Mexico. Of these, we sampled three of the four species of Sibon known from Mexico, both species of Dipsas and five out of the seven species of Tropidodipsas. We were not able to sample Tropidodipsas repleta Smith, Lemos-Espinal, Hartman & Chiszar, 2005; T. zweifeli Liner & Wilson, 1970; the subspecies T. fasciata kidderi; T. sartorii madougalli; Sibon linears Pérez-Higareda, López-Luna & Smith 2002; nor the population of T. cf. philippii from Oaxaca reported by Kofron (1987). Additionally, we sampled four species of Geophis which belong to two...
species groups that are hypothesized to be closely related to snail-suckers (Sheehy 2013).

We photographed all live snakes, including dorsal, lateral, and ventral profiles, and euthanized them with pentobarbital. We took tissue samples from muscle or liver upon death and preserved them in 96% ethanol. We fixed specimens in 10% formalin and transferred them to 70% ethanol for permanent storage.

The material collected was deposited at the Instituto de Investigaciones sobre los Recursos Naturales (INIRENA) of the Universidad Michoacana de San Nicolás de Hidalgo (UMSNH) in Morelia, Mexico; the Museo de Zoología, Facultad de Ciencias (MZFC) of the Universidad Nacional Autónoma de México (UNAM) in Mexico City; the Colección Herpetológica de la Facultad de Ciencias B ólogicas (CHFCB) of the Universidad Juárez del Estado de Durango (UJED) and the University of Texas at Arlington, Texas (UTA). Although we formally accessioned the specimens we collected, several specimens examined from both the MZFC and UTA collections have not been catalogued, in which case we list the original field numbers and the respective museum in which they were deposited. Original field number abbreviations are as follows: CIG (Christoph I. Grünwald) to be catalogued at MZFC; ENS (Eric N. Smith) to be catalogued at UTA; JAC (Jonathan A. Campbell) to be catalogued at UTA; JRV (Jacobo Reyes-Velasco) to be catalogued at UTA.

Museum acronyms throughout follow Sabaj-Pérez (2016). Representative specimens of snail-suckers and Geophis were examined in the following collections, in addition to specimens deposited in the collections reported above: the University of Kansas Natural History Collection (KU), Natural History Museum of Los Angeles County (LACM), Texas Cooperative Wildlife Collection (TCWC), University of Illinois Museum of Natural History (UIMNH). Specimen numbers for all material examined are provided in Appendix 1. We were not able to measure type specimens of some previously described taxa, so we used the measurements of the type specimens provided in the original descriptions and other published literature.

Taxonomy

We agree with Smith (1982) that the genus Sibon is a masculine noun so the specific epithet should also be masculine. We follow Kofron’s (1980, 1985a, 1985b, 1987, 1988, 1990) proposal for the majority of recognized species, with the following differences. We recognize the Tropidodipsas fasciata species group as proposed by Kofron (1987); however, we recognize Tropidodipsas as distinct from Sibon following Wallach (1995). We do not consider Sibon anthracops to belong to Tropidodipsas based on the phylogenetic evidence presented by Sheehy (2013). We recognize Tropidodipsas guerreroensis as a separate species rather than a subspecies of T. fasciata (see below). Thus, we consider Tropidodipsas to contain the T. fasciata species group which consists of five species (T. fasciata, T. guerreroensis, T. philippii and the two species described below) as well as two un-assigned species which we have not been able to sample (T. repleta and T. zweifeli) and T. fischeri which is tentatively retained in the genus pending more work (see below).

Measurements and character states

Our measurements and character states follow Kofron (1980) for all Dipsadidae and Downs (1967) for Geophis. Measurements and character states were taken from data published in Kofron’s succession of work on Mexican Dipsadidae such as Tropidodipsas Kofron (1980), Dipsas gaigeae and D. brevifacies (Kofron 1982), T. fischeri species group (Kofron 1985a), Tropidodipsas (Kofron 1987), T. sartorii species group (Kofron 1988), S. dimidiatums (Kofron 1990) as well as Davis (1953) for T. guerreroensis. Some measurements and characters for Central and South American Dipsadidae were taken from Lotzkat et al. (2012) and Arteaga et al. (2018). Abbreviations used in the text and tables are as follows: snout–vent length (SVL), tail length (TL), total length (TOL), head length (HL), head width (HW), snout length (SL), eye diameter (ED), chin shield (CS).

Scale counts were performed with the aid of a dissecting microscope. Measurements were taken with a ruler or digital calipers (Truper®, Mexico) under a dissecting microscope. Bilateral characters were scored on both left and right sides and given in that order, separated by a slash (/). Head length was measured from the tip of the snout to the posterior end of the parietals (following Peters 1964), head width was measured at the widest point of the head at the posterior part of the jaw. All scale dimensions were measured at their maximum.

To examine dentition characters, the maxilla and ectopterygoid were removed from the skull and cleansed in a dilute solution of Proteinase K for approximately one hour.

DNA extraction and amplification

We sequenced genetic data from two nuclear (DNAH3, NT5) and two mitochondrial (cytb, ND4) loci, and combined our data with previous studies of the group (Arteaga et al. 2018; Sheehy 2013).

We used 370 µL of Monarch gDNA Tissue Lysis buffer (New England Biolabs T3011L) and 20 µL of Proteinase K (New England Biolabs P8107S) to digest the tissue at 55°C overnight. We extracted whole genomic DNA from lysed samples using the Serapure bead extraction protocol of Rohland and Reich (2012) with modifications based on Glenn et al. (2019) for Sera-Mag SpeedBeads (Fisher Scientific 09-981-123).

We used polymerase chain reaction (PCR) to amplify two nuclear genes and two mitochondrial genes. The nuclear genes were DNAH3 using DNAH3-fl as the
forward primer and DNAH3_r6 as the reverse primer (Townsend et al. 2008) and NT3 using NT3_F3 as the forward primer and NT3_R4 as the reverse primer (Noonan and Chippindale 2006). The mitochondrial genes were cytb using Gludg-L as the forward primer (Palumbi 1996) and ATRCB3 as the reverse primer (Harvey et al. 2000) and ND4 using ND4 as the forward primer and ND4_Leu as the reverse primer (Arevalo et al. 1994). All four loci were amplified using a standard PCR protocol: 95°C for 3 min 30 sec, 35 cycles of 95°C for 30 sec, 51°C for DNAH3 and NT3 and 53°C for cytb and ND4 for 30 sec, 72°C for 1 min, followed by 15 min at 72°C and a final hold at 10°C. All PCR reactions were 30 µL and contained the following: 16.9 µL of water, 3 µL of 10X Reaction Buffer (New England Biolabs M0320L), 1.8 µL MgCl₂ (25mM), 0.6 µL dNTPs (2.5mM each; New England Biolabs N0446S), 0.3 µL DMSO, 0.6 µL of each primer (10 µM), 0.2 µL of Taq DNA Polymerase (New England Biolabs M0320L), and 6 µL of genomic DNA. PCR reactions were purified using 5 µL of Quick Cip (New England Biolabs M0525S) and incubating at 37°C for 30 min and inactivating the enzymes at 80°C for 15 min.

Sequencing was done in both directions using Eurofins Genomics LLC (Louisville, KY, USA). Forward and reverse reads were merged and trimmed in Geneious Prime v2020.2.1 (Biomatters Ltd., Auckland, NZ) and manually screened for errors and ambiguities. Heterozygous sites in nuclear genes were coded using International Union of Pure and Applied Chemistry (IUPAC) ambiguity codes. All sequences were deposited in GenBank. Information on all the new sequences as well as other sequences used in this study are found in Table 1.

Table 1. GenBank Numbers of sequences used in this study. New sequences are indicated in bold.

| Species                     | Locality         | Specimen number | ND4       | cytb      | NT3       | DNAH3    |
|-----------------------------|------------------|-----------------|-----------|-----------|-----------|----------|
| Chersodromus liebmannii     | Mexico: Oaxaca   | AMNO-2298       | JX398451  | JX398604  | JX398732  | JX293840 |
| Dipsas andiana             | Ecuador          | JM-79           | JX398453  | JX398607  | JX398744  | JX293843 |
| Dipsas articulata          | Costa Rica       | D-161           | JX398454  | JX398740  |           |          |
| Dipsas bicolor             | Costa Rica       | ASL-277         | JX398455  | JX398741  | JX293844  |          |
| Dipsas babriedgeyi         | Ecuador          | MZUTI-5414      |           | MH374984  |           |          |
| Dipsas catesby             | Ecuador          | MZUTI-5417      |           | MH374985  |           |          |
| Dipsas gaigeae             | Mexico: Colima   | JAC-28327       | JX398461  | JX398612  | JX293849  |          |
| Dipsas guatemala           | Mexico: Guatamala| JAC-28587       | JX398462  | JX398613  | JX293873  | JX293850 |
| Dipsas gracilis            | Ecuador          | JRV-30          | JX398464  | JX398614  | JX293878  | JX293851 |
| Dipsas indica              | Peru             | KU-204908       | JX398468  | JX398618  | JX293874  | JX293853 |
| Dipsas klebbai             | Ecuador          | QCAZ-12717      |           | MH375019  |           |          |
| Dipsas mikanii             | Brazil           | CTMZ-495        | JX398469  | JX398619  | JX293848  |          |
| Dipsas nicholi             | Panama           | JM-812          | JX398469  | JX398619  | JX293848  |          |
| Dipsas oswalldoboezi        | CEAZ-10369       |                 |           | MH374997  |           |          |
| Dipsas pavonina            | Brazil           | LSUMH-13989     | JX398470  | JX398620  | JX293849  | JX293855 |
| Dipsas peruana             | Peru             | LSUMH-15352     | JX398472  | JX398622  | JX293870  | JX293856 |
| Dipsas petersi             | Ecuador          | JM-72           | JX398555  | JX398695  | JX293818  | JX293898 |
| Dipsas pratti              | Venezuela        | MBU-UCV-6837    | JX398473  | JX398624  | JX293751  |          |
| Dipsas temporalis           | Panama           | JM-664          | JX398476  | JX398626  |           |          |
| Dipsas trinitatis           | Trinidad         | UWMZT-20110.25  | JX398479  | JX398629  |           |          |
| Dipsas turgidus            | Bolivia          | LSUMH-64588     | JX398556  | JX398696  | JX293819  | JX293899 |
| Dipsas variegata           | Suriname         | UTAR-15772      | JX398482  | JX398601  | JX293736  | JX293858 |
| Geophis bicolor            | Mexico: Jalisco  | INIRENA-2793    | MZ287388  | MZ287374  | MZ287422  |          |
|                            | Mexico: Michoacán| JAC-24684       | JX398487  | JX398637  | JX298759  | JX293862 |

Molecular Analysis

Alignments for each locus were performed in MAFFT version 7 (Katoh & Standley 2013). We used Geneious v9.1.6 (Biomatters Ltd, Auckland, NZ) to manually trim any regions of poor alignment and to make sure that protein-coding genes were in the correct reading frame. We then concatenated all genes with the use of FASconCAT-G v1.04 (Kück and Longo 2014). Our concatenated dataset consisted of 3,248 bp, of which 1987 bp correspond to two mitochondrial loci and 1261 bp to two nuclear loci.

We chose the best-fit model of nucleotide evolution for each locus with the use of the Bayesian Information Criterion (BIC) in PartitionFinder v1.1.1 (Lanfear et al. 2012). Table 2 shows the best-fit model for each locus. We partitioned our dataset by locus and by codon position. We then performed Bayesian phylogenetic inference (BI) in Mr. Bayes v3.2.2 (Ronquist et al. 2012) on the CIPRES science gateway server (Miller et al. 2011).

Our Bayesian phylogenetic analysis consisted of four runs of 10 million generations, sampling every 1,000th generation. Each run contained four chains, three heated and one cold. We checked for convergence between runs with the use of Tracer v1.6 (Rambaut et al. 2015) by visually inspecting overlap in likelihood and parameter estimates between runs, as well as effective sample sizes and potential scale reduction factor (PSRF) for each run. The individual runs had converged by 200,000 generations (based on the PSRF), so we conservatively discarded the first 25% of each run as burn-in. The runs were combined with the use of TreeAnnotator (Bouckaert et al. 2019) and visualized in FigTree v1.4.2 (Rambaut 2014).
| Species                      | Locality          | Specimen number | ND4        | cytb       | NT3       | DNAH3  |
|------------------------------|-------------------|-----------------|------------|------------|-----------|--------|
| Geophis nigrocinctus         | Mexico: Jalisco   | JAC-30704       | JX398488   | JX398638   | –         | –      |
| Geophis omissimorphanus      | Mexico: Guerrero  | ENS-11496       | –          | JX398639   | JX398760  | –      |
| Geophis tarascae             | Mexico: Michoacán | JAC-24692       | JX398489   | JX398640   | JX398761  | JX293870|
| Ninia atra                   | Colombia          | MHAJ-1452       | QG334659   | QG334553   | –         | QG334577|
| Ninia diademata              | Guatemala         | UTAR-42291      | –          | JX398645   | JX398764  | JX293864|
| Sibon annulatus              | CostaRica         | D-167           | JX398501   | JX398552   | JX398772  | JX293869|
| Sibon anthracops             | CostaRica         | ASL-198         | JX398506   | JX398567   | JX398778  | JX293872|
| Sibon angus                  | Costa Rica        | ASL-283         | JX398508   | JX398660   | JX398781  | JX293878|
| Sibon bevidigelyi            | Ecuador           | MZUTI-3269      | –          | MH374962   | –         | –      |
| Sibon carri                  | Guatemala         | UTAR-45493      | JX398514   | JX398665   | JX398786  | JX293876|
| Sibon dimidiius              | CostaRica         | B4S-62          | JX398515   | JX398666   | JX398787  | JX293877|
| Sibon lamari                 | Costa Rica        | No Number       | JX398520   | JX398671   | JX398791  | JX293879|
| Sibon longifrensis           | Costa Rica        | ASL-320         | JX398521   | JX398672   | JX398792  | JX293880|
| Sibon manzanarensi           | Honduras          | USNM-570455     | JX398524   | JX398685   | JX398795  | JX293883|
| Sibon merendonensis          | Guatemala         | MVZ-263880      | JX398526   | JX398675   | JX398797  | JX293884|
| Sibon miskitus               | Honduras          | USNM-570454     | JX398528   | JX398677   | JX398799  | JX293885|
| Sibon nebulatus              | Mexico: Chiapas   | INIRENA-278B    | MZ287387   | MZ287377   | Pending   | –      |
| Sibon perissasticum          | Panama            | SMF-88716       | JX398552   | JX398688   | JX398847  | JX293888|
| Geophis sanioius new comb.   | Mexico: Yucatan   | JAC-24409       | JX398533   | JX398692   | JX398815  | JX293895|
| Geophis anuliferus new comb. | Mexico: Guerrero  | JAC-27792       | JX398559   | JX398699   | –         | JX293914|
| Mexico: Colima               | JAC-30143         | –               | JX398561   | JX398701   | –         | –      |
| Tropidodipsas cf. fasciata   | Mexico: Yucatan   | INIRENA-278B    | –          | MM287421   | MM287421  | –      |
| Tropidodipsas fasciata       | Mexico: Oaxaca    | JAC-22920       | –          | JX398702   | –         | –      |
| Tropidodipsas fischi         | Mexico: Oaxaca    | JAC-30740       | JX398580   | JX398713   | –         | –      |
| Tropidodipsas guerrerienensis| Mexico: Guerrero  | CHFBC-032B      | MZ287390   | MZ287378   | –         | –      |
| Tropidodipsas guerrerienensis| Mexico: Oaxaca    | CHFBC-0335      | MZ287397   | MZ287379   | –         | –      |
| Tropidodipsas guerrerienensis| Mexico: Chiapas   | CHFBC-0352      | MZ287398   | MZ287380   | –         | –      |
| Tropidodipsas guerrerienensis| Mexico: Chiapas   | INIRENA-278B    | MZ287395   | MZ287381   | MZ287420  | MZ287403|
| Tropidodipsas guerrerienensis| Mexico: Oaxaca    | JAC-22545       | –          | JX398828   | –         | –      |
| Tropidodipsas guerrerienensis| Mexico: Oaxaca    | JAC-24267       | JX398594   | JX398724   | JX398839  | JX293999|
| Tropidodipsas guerrerienensis| Mexico: Guerrero  | JAC-27750       | JX398571   | JX39871  | –         | JX293908|
| Tropidodipsas papavericola   | Mexico: Oaxaca    | INIRENA-2805    | MZ287392   | MZ287382   | MZ287418  | MZ287400|
| Tropidodipsas papavericola   | Mexico: Oaxaca    | INIRENA-2801    | MZ287383   | MZ287419   | –         | –      |
| Tropidodipsas papavericola   | Mexico: Guerrero  | INIRENA-2802    | MZ287391   | MZ287384   | MZ287417  | MZ287401|
| Tropidodipsas papavericola   | Mexico: Guerrero  | INIRENA-2803    | MZ287393   | –         | –         | –      |
| Tropidodipsas philippi       | Mexico: Jalisco   | ENS-11639       | JX398569   | –         | JX293907  | –      |
| Tropidodipsas philippi       | Mexico: Yucatan   | JAC-24811       | JX398570   | JX398710   | JX398826  | –      |
| Tropidodipsas philippi       | Mexico: Michoacán | JAC-27923       | JX398572   | JX398712   | –         | JX293909|
| Tropidodipsas philippi       | Mexico: Michoacán | JAC-27923       | JX398573   | –         | –         | JX293910|
| Tropidodipsas philippi       | Mexico: Jalisco   | JAC-30601       | JX398577   | –         | –         | –      |
| Geophis sartori new comb.    | Mexico: San Luis  | INIRENA-278B    | MZ287390   | MZ287375   | MZ287412  | MZ287406|
| Geophis sartori new comb.    | Mexico: San Luis  | INIRENA-278B    | MZ287389   | MZ287376   | MZ287410  | –      |
| Geophis sartori new comb.    | Mexico: San Luis  | INIRENA-278B    | –          | –         | MZ287411  | MZ287408|
| Geophis sartori new comb.    | Unknown           | JAC-30401       | JX398583   | JX398716   | –         | –      |

Table 2. Best fit models of nucleotide evolution for each locus.

| Partition | Partition set | Best-Fit Model |
|-----------|--------------|----------------|
| 1         | cytb (first codon) | GTR + i + gamma |
| 2         | cytb (second codon) | HKY + i + gamma |
| 3         | cytb (third codon)  | GTR + i + gamma |
| 4         | DNAH3 (first codon) | JC |
| 5         | DNAH3 (second codon) | JC + gamma |
| 6         | DNAH3 (third codon) | HKY + gamma |
| 7         | ND4 (first codon)  | GTR + i + gamma |
| 8         | ND4 (second codon) | HKY + gamma |
| 9         | ND4 (third codon)  | GTR + gamma |
| 10        | NT3 (first codon)  | HKY + gamma |
| 11        | NT3 (second codon) | HKY + gamma |
| 12        | NT3 (third codon)  | HKY + gamma |
| 13        | tRNA            | HKY + gamma |
We consider a clade as highly supported if the posterior support value was greater than 0.95.

Additionally, we calculated uncorrected p-distances for the mitochondrial gene Cytochrome B (cytb) in the program MEGA X (Kumar et al. 2018). We calculated genetic distances of cytb once we completed the concatenated data set.

Results

Molecular phylogenetic results

We used a total of 79 individuals in our molecular phylogeny, including 76 snail-sucker individuals as well as three outgroup taxa. Of the individuals used, 18 specimens were novel sequences obtained by us, representing 9 taxa. As shown by previous studies (Sheehy 2013; Arteaga et al. 2018), the three currently recognized genera of snail-suckers form a monophyletic group, which also received high support in our analysis (Posterior Probability [pp] = 1) (Fig. 6).

Our results recovered a monophyletic Dipsas, including members previously allocated to Sibynomorphus, a South American genus previously synonymized with Dipsas (Arteaga et al. 2018). The only exception was D. gaigeae, a species restricted to western Mexico. This species and Tropidodipsas fischeri each formed monophyletic, highly supported, independent clades but their relationships with the remaining species of snail-suckers are poorly supported and remain ambiguous as they form a polytomy in a clade containing Tropidodipsas, Sibon and Geophis. Combined, these results were consistent with those reported by Sheehy (2013). We also recovered a strongly supported clade containing the majority of species of Tropidodipsas. This clade consisted of two main groups. The first one included both taxa from southern Mexico described herein, as well as an individual of T. cf. fasciata from Yucatan (with low node support, pp = 0.67). The other group in Tropidodipsias consisted of an individual of T. cf. fasciata from Tamaulipas (Fig. 7A), T. fasciata from Oaxaca, which was sister to T. guerrerensis + T. philippi (also with low node support, pp = 0.65). The fact that the three geographically distinct populations of T. fasciata fall in different positions in the tree suggest that this taxon needs further study and better sampling. Unfortunately, more material is currently unavailable, as most T. fasciata samples are from the Oaxaca (Isthmus of Tehuantepec) population. The type locality of T. fasciata has not been delimited beyond “Mexico” and thus it would be difficult to assign a specific clade to that name. Our suggestion is that a more detailed study of the T. fasciata species complex should be undertaken in the future.

The remaining snail-suckers grouped into two clades. The first clade is composed of multiple species currently allocated to three different genera: Sibon sanniolus from southeastern Mexico; the species Tropidodipsas annulifera and T. sartorii; multiple members of the genus Geophis. This clade had moderate support (pp = 0.88) and the node with T. sartorii and T. annulifera plus several Geophis was recovered with more robust support (pp = 0.93). From here on we refer to all these species as Geophis (see below). Several of the internal nodes were recovered as weakly supported (pp < 0.90). These results are similar to what was presented by Sheehy (2013) and Arteaga et al. (2018).

The last clade consisted of the majority of species in the genus Sibon, with the exception of S. sanniolus. The majority of nodes in this clade were recovered as highly supported (pp > 0.95). These results were in agreement with Sheehy (2013).

Our phylogenetic results support the novelty of the two species described herein, as we show that they are not conspecific with any previously described taxa and form monophyletic clades. Furthermore, our results suggest that these two new species are each other’s closest relative, and together with a sample of T. cf. fasciata from Yucatan they form a sister clade to the clade containing T. cf. fasciata, T. philippi and T. guerrerensis. Our analysis supports the validity of T. guerrerensis as a species as originally described, not a subspecies of T. fasciata, as it is more closely related to T. philippi and a sample of T. fasciata from Tamaulipas (CIG 819) than the T. fasciata from nearby Oaxaca (see below).

Systematic accounts

Tropidodipsas tricolor sp. nov.

http://zoobank.org/54130B5D-9C0D-4C8C-8417-00EB50E65216

Figs 1–2

Proposed standard English name: Tricolor Snailsucker

Proposed standard Spanish name: Caracolera tricolor

Holotype. (Fig. 1) INIRENA 2800 (original field number CIG 1837). Juvenile male, collected at 1.5 km east of Rio Verde, Municipio de Atoyac de Álvarez, (17.3131°, -100.1969°, datum=WGS84, 971 m a.s.l.), Guerrero, Mexico (Fig. 3A) on August 14th, 2020 by Miguel A. Peñaloza-Montaño, Jason M. Jones and Jacobo Reyes-Velasco.

Paratypes. (2) (Figs. 2C–D) INIRENA 2799 (original field number CIG 1596). Adult of unknown sex, DOR, collected at 26 km N of Putla Villa de Guerrero, on Putla Villa de Guerrero - Oaxaca Hwy., Municipality of Putla de Guerrero (17.1494°, -97.8709°, datum=WGS84, 1785 m a.s.l.), Oaxaca, Mexico, on September 4th, 2019 by Christoph I. Grünwald, Christopher M. Rodriguez and Carmen Mendoza-Portilla; INIRENA 2798 (original field number CIG 1863). Adult female, DOR, collected at 4.5 km NW of Mixtecapa, on the road to Malinaltepec, Municipio de Malinaltepec, (17.2539°, -98.6406°, datum=WGS84, 1815 m a.s.l.), Guerrero, Mexico on October 21st, 2020 by Tziuhtécatl Sánchez-Luna.

Diagnosis. Tropidodipsas tricolor sp. nov. is placed in the genus Tropidodipsas based on phyllogenetic evidence
(Fig. 6). It belongs to the *Tropidodipsas fasciata* group as defined by Kofron (1987) based on possessing a laterally compressed body, head distinctly wider than neck, protruding eyes of moderate to large diameter, vertebral and paravertebral scales not wider than other dorsal scales, 17 maxillary teeth, 18 dentary teeth, postmental scale absent. The species differs from all described species of *Tropidodipsas* based on the following combination of characters: (1) 15/15/15 rows of smooth dorsal scales with no enlarged vertebral row; (2) prefrontal broader than long, entering the orbit; (3) loreal square, equally long as high, not entering the orbit; (4) 183 ventral scales in males, 183 in females; (5) 78–79 divided subcaudals in males and 79 in females; (6) 19–22 reddish orange, light-edged bands on body, most with dark brown or black medial stippling, giving the snake a “tricolor” effect; 8 pale bands on tail; (7) ground color black or nearly so; (8) iris chocolate brown; (9) TL/SVL 0.31 in one male specimen, and 0.30 in one female specimen.

**Comparisons.** *Tropidodipsas tricolor* sp. nov. is most similar to *T. philippii*, *T. fasciata*, *T. guerreroensis* and the new species described below. It is the only species of snail-sucker in Mexico with a tricolor pattern, although similar patterns exist in Central American (e.g., *Sibon anthracops*) and South American (e.g., *Dipsas bobrid-gelyi*) snail-suckers. It is distinguished from other Mexican snail-suckers such as the *Geophis chalybeus* species group, the *Geophis omiletmanus* species group, *G. sartorii* new comb., *G. annuliferus* new comb., *Tropidodipsas fischeri* (Fig. 9C,D), *S. carri*, *G. sanniols* new comb. and *S. linears* by the laterally compressed body shape and the tri-colored pattern. Further, it differs from other snail-sucker species (character states in parenthesis) as follows: It differs from *S. anthracops* by possessing 15 scale rows at midbody (vs. 13), possessing a prefrontal that enters the orbit (vs. prefrontal does not enter orbit), loreal not entering the orbit (vs. loreal enters orbit). It may be distinguished from *S. dimidiatus* by its noticeably different tricolor banded dorsal coloration (vs. brown snake with dark blotches, spots or an undulating stripe) and by lacking a post-mental scale (vs. usually one or two post-mental scales present) and by possessing less than 80 subcaudals in males (vs. 96–144). It is distinguished from *Sibon nebulatus* (Fig. 9A,B) by possessing tricolor body bands (vs. a nebulated black, grey and white pattern with no red outlines in the pale bands), and by vertebral scales not enlarged (vs. row of vertebral scales 1.25–1.35 times larger than other dorsal scales).

This species differs from *Dipsas gaigeae* by having 15 dorsal scale rows at mid-body (vs. 13), possessing more than 180 ventral scales (vs. 155–169) and a loreal that does not enter orbit (vs. loreal entering orbit). Distinct from *D. brevijacies* by possessing more than 180 ventrals (vs. 162–180), always possessing the prefrontal in contact with orbit (vs. usually not), never possessing a loreal in contact with orbit (vs. usually in contact), usually possessing 1+2 temporals (vs. usually 2+3), and usually possessing one pair of infralabials in contact after the mental (vs. usually two pairs of infralabials in contact after the mental).

Within *Tropidodipsas*, *T. tricolor* sp. nov. differs from *T. fasciata* and *T. guerreroensis* by possessing 15 smooth dorsal scale rows (vs. 17 keeled scale rows) and by prefrontal entering orbit (vs. not entering orbit). It differs from the new species described below (*Tropidodipsas papavericola* sp. nov.; see below) by tricolor outline in pale dorsal bands (vs. unicolor pale body bands), 19–22 reddish orange body bands (vs. 25–33 pale body bands), by possessing a prefrontal which enters the orbit (vs. prefrontal not entering orbit), by possessing 78–79 subcaudal scales in males (vs. 69–76), by possessing one precaulal (vs. two), by possessing 2–3 postoculocals (vs. 1–2), 7–8 supralabials (vs. 5–7), 8–9 infralabials (vs. 6–7), eye–head length ratio 25% (vs. 17–21%), by pale throat coloration with black stippling concentrated toward anterior portion and a black mental (vs. pale throat coloration with random dark spots not concentrated in any specific region and a black and white mental), and a narrower head with less protruding eyes (vs. head noticeably wider than neck and eyes strongly protruberant). It differs from *T. philippii* (Fig. 8C,D) by presenting one precaulal (vs. 2–3), prefrontal entering orbit (vs. not entering orbit), supralabials 7–8 (vs. 6–7), 3 gular scales (vs. 5), a longer tail which is 30–31% of SVL in males (vs. 25–29% in males), consistently 19–21 pale body bands (vs. highly variable, from 9–44 according to Kofron (1980, 1987) but usually 11–21, according to our data (n=13), and eye–head length ratio of 25% (vs. 15–23%).

Genetic divergence in a 1,072-bp fragment of mitochondrial *cyt b* gene between *T. tricolor* sp. nov. and *T. papavericola* sp. nov. is 13–14%; between *T. tricolor* sp. nov. and geographically proximate *T. fasciata*, 14–15%; between *T. tricolor* sp. nov. and *T. guerreroensis*, 14–16%; and between *T. tricolor* sp. nov. and geographically proximate *T. philippii*, 15–16%.

**Description of holotype.** (Fig. 1) INIRENA 2800. Juvenile male. SVL 210 mm; TL 67 mm; TotL 277 mm. HL 8.5 mm (from tip to posterior border of parietal) and 7.7 mm (from snout to posterior commissure of mouth); HW 5.0 mm, head distinct from body, approximately 2 times wider than neck. Snout short, blunt, SL 2.7 mm, contained 3.1 times in the HL, rounded in dorsal and lateral profile. Loral region tall, canthus rostralis rounded. Rostral 1.4 times as broad as high (2.0 mm broad, 1.4 mm high); internasals broader than long (1.0 mm length, 1.6 mm width), rounded anteriorly, in lateral contact with anterior and posterior nasals; prefrontals relatively large, broader than long (1.7 mm length, 2.0 mm width), nearly rectangular, with posterior edge of scale pointed at supraocular and frontal suture, in median contact with each other and in lateral contact with postnasal. loreal, preocular (lower) and orbit; frontal 1.1 times longer than broad (2.8 mm long, 2.4 mm broad), pentagonal with angular tip posteriorly, in contact with prefrontals, supraoculocals and parietals. One moderately large preocular on each side. Single supraocular moderate, in contact with pre-
Figure 1. *Tropidodipsas tricolor* sp. nov. Holotype (INIRENA 2800). Holotype (INIRENA 2800) in life (A); Ventral aspect of Holotype (INIRENA 2800) in preservative (B,E); Dorsal aspect of Holotype (INIRENA 2800) in preservative.
Mental 2.1 times as broad as long (1.5 mm broad, 0.7 mm long), flat anteriorly, rounded posteriorly with posterior edge coming to a slight point at the suture of the first infralabials, separated from the anterior chinshields by the enlarged first pair of infralabials which are in contact with each other. Infracalabials 8 on both sides, 1–5 in contact with anterior chinshields and 5–6 in contact with posterior chinshields, sixth infralabial is largest. Anterior chinshields elongated, much longer than wide, left chinshield 2.4 mm long and 0.9 mm wide (2.7 times as long as wide) and right anterior chinshield 2.6 mm long and 1.0 mm wide (2.6 times as long as wide). Left posterior chinshield 1.9 mm long and 1.1 mm wide (1.7 times as long as wide) and right posterior chinshield 1.8 mm long and 1.1 mm wide (1.6 times as long as wide). Three gular scales. Infraoculars and scales in the chin region smooth. Distinct gular fold present, starting on the first ventral scale and running posteriorly to the twentieth ventral scale, then fading out by the twenty-fifth ventral scale. Dorsal scales in 15-15-15 rows, smooth throughout; apical pits not evident. Ventrels 183; anal plate single; subcaudal scales paired, 79 on both sides. Body shape laterally compressed. Pupil elliptical.

Coloration in life of the holotype (Fig. 1A). Dorsal coloration of head, body and tail black with 19 reddish orange, light-edged bands on body (not counting nuchal
band) and 9 bands on tail; red-orange color faded on four anteriormost bands. Nuchal band cream and involving latter fifth portion of the parietals and the anterior two and part of the third dorsal scale rows. Body bands 9–19 and all tail bands exhibit scattered dark stippling. The entire snake has a “tricolor” appearance. Pale bands on body and tail 2–4 scale rows long (mean = 3) and wider on the lateral surfaces due to the increased scale size. Head coloration predominately black on dorsal surface and pale cream ventrally. The pale nuchal coloration extends forward onto posterior part of the parietals, and beneath the temporals onto the supralabials. Anterior supralabial black, the others bicolor, posterior supralabial predominately cream on both sides but with black flecking in the middle. Dark body color almost complete ventrally, occasionally interrupted mid-ventrally by a pale line. Light dorsal bands are complete ventrally but with dark flecking towards posterior part of body. Reddish orange coloration of dorsal bands does not continue onto the venter. Anal plate bicolor, black on the margins and pale cream in the middle. First dark band on tail interrupted mid-ventrally by pale cream coloration, the other dark and pale bands on tail complete ventrally. Tail tip black above, pale orange below. Pupil pale gray, iris chocolate brown in life.

**Coloration in preservative of the holotype** (Figs 1B–E). General coloration tricolor. Reddish orange wedge-shaped markings have turned pale brown; the pale cream and orangish bands faded to white.

**Variation.** Meristic variation of the three available specimens is given in Table 3. One paratype (INIRENA 2798) possesses a tiny second preocular above the large lower preocular on both sides. This small scale does not prevent the prefrontal from entering the orbit as in the other specimens and may be a division of the lower preocular.

Two individuals were photographed by Peter Heimes in Guerrero in 2007. While these individuals were not collected or examined in detail, they are likely of this species. As we lack photos of variation of this new species in life, we have included photos of these two individuals which most probably belong to this species (Fig. 2A,B).

**Dentition.** One paratype (INIRENA 2798) has 17 maxillary teeth (including three empty sockets) on the right side. The same specimen had 18 dentary teeth (including one empty socket) on the right side.

**Distribution, habitat and ecology.** This species appears to be restricted to moderate elevations in Sierra Madre del Sur from central Guerrero to western Oaxaca (Fig. 10). Habitats include cloud forest, mesic pine-oak woodland, tropical evergreen forest, tropical semi-deciduous forest and combinations of those assemblages (Brown et al. 2007). Known from 700–2200 m a.s.l., though most localities are from 1700–2200 m a.s.l. with the exception of the Sierra de Atoyac where it has been found at 700–1000 m a.s.l. in mesic tropical evergreen forest (Fig. 3A). Specimens were found at night on low or medium (1.5–3.0 m) vegetation, or dead on the road.

**Etymology.** The specific epithet *tricolor* refers to the tricolor pattern of the black dorsal coloration interspersed by light dorsal bands of cream and reddish orange.

*Tropidodipsas papavericola* sp. nov.

http://zoobank.org/6D7EA79C-98E3-4287-92A9-E81C1831D6CA

Figs. 4–5

Proposed standard English name: Poppyfield Snailsucker

Proposed standard Spanish name: Caracolera amapolera

**Paratypes/ (5) (Fig. 5) INIRENA 2802 (original field number CIG 1496, Fig. 5C). Sub-adult male, collected AOR 18.1 km S of Puerto del Gallo on road from Nuevo Dehli to Puerto del Gallo, Municipio de Atoyac de Álvarez (17.4342°, -100.191°, datum=WGS84, 1654 m a.s.l.), Guerrero, Mexico on June 29th, 2019 by Christoph I. Grünwald, Miguel A. Peñaloza-Montaño, Eduardo Y. Barrera-Nava, Karen I. Morales-Flores and Janelle Morales-Flores. INIRENA 2803 (original field number CIG 1502). Sub-adult male collected DOR at 5 km S of La Laguna, on road from San Luis La Loma to Bajitos de la Laguna, Municipio de Técpan de Galeana (17.5137°, -100.7737°, datum=WGS84, 1686 m a.s.l.), Guerrero, Mexico on June 29th, 2019 by Jason M. Jones and Carlos Montaño-Ruvalcaba. INIRENA 2804 (original field...
Distinct from the superficially similar *Sibon nebulatus* (Fig. 9A, B) by vertebral row of dorsal scales same size as other dorsal scales (vs. vertebral row of dorsal scales 1.25–1.35 times larger than other dorsal scales), smaller adult body size of 450–560 mm in males (vs. up to 890 mm in males), fewer dark body bands, 26–35 on body (vs. 36–47 on body), head predominately dark brown or black (vs. head pattern speckled with dark and light marking) and absence of pink speckling in the dorsal pattern (vs. pink speckling present).

It differs from *Dipsas gaigeae* by having 15 dorsal scale rows at midbody (vs. 13), possessing 179–189 ventral scales (vs. 155–169), loreal that does not enter orbit (vs. loreal enters orbit), and 26–35 pale body bands (vs. 7–12). Distinct from *Dipsas brevifacies* by possessing 179–189 ventral scales (vs. 162–180), prefrontal and loreal not in contact with orbit (vs. variable in both), usually possessing 1±2 temporals (vs. usually 2±3), usually possessing one pair of infralabials in contact after the mental (vs. usually two pairs of infralabials in contact after the mental). Within *Tropidodipsas*, *T. papavericola* sp. nov. differs from *T. fasciata* and *T. guerreroensis* by possessing 15 smooth dorsal scale rows (vs. 17 keeled scale rows).

*Tropidodipsas papavericola* sp. nov. differs from *T. tricolor* sp. nov. by lacking the tricolor outline in the dorsal bands (vs. possessing a tricolor outline in dorsal bands), 26–35 dorsal dark bands on body (vs. 19–22), by possessing a prefrontal which does not enter orbit (vs. prefrontal entering orbit), 2 preoculars (vs. 1), 5–7 supralabials (vs. 7–8), 6–7 infralabials (vs. 8–9), less subcaudal scales, 69–76 in males (vs. 78–79 in males), by possessing a pale ventral coloration of head with irregularly scattered dark spots, and a black and white mental (vs. pale ventral coloration of head with heavy dark stippling anteriorly and black mental), and by a smaller eye - head length ratio of 17–21% (vs. 25%). It is most similar to *T. philippii* (Fig. 8C, D), from which it can be distinguished by having smooth dorsal scales (vs. slightly keeled), a longer tail which is 30–35% of SVL in males (vs. 25–29% in males), and less subcaudal scales, 69–76 in males (vs. 76–90 in males).

Genetic divergence in a 1,072-bp long fragment of the mitochondrial *cytB* gene between *T. papavericola* sp. nov. and *T. philippii* is 10–12%; between *T. papavericola* sp. nov. and *T. guerreroensis*, 10–13%; between *T. papavericola* sp. nov. and *T. fasciata* it is 10–12%.

**Description of holotype.** (Fig. 4) INIRENA 2801. Adult male. SVL 421 mm; TL 137 mm; TotL 558 mm. HL 13.8 mm (from tip to posterior border of parietal) and 13.5 mm (from snout to posterior commissure of mouth); HW 9.8 mm, head distinct from body, approximately 2 times wider than neck. Snout long, SL 5.7 mm, contained 2.4 times in HL, rounded in dorsal and lateral profile. Loreal region tall, canthus rostralis angular. Rostral 1.2 times as broad as high (3.3 mm wide, 2.7 mm high); internasals broader than long (1.7 mm length, 2.8 mm width), rounded anteriorly, in lateral contact with anterior and posterior nasals; prefrontals longer than broad, (3.8 mm length, 2.9 mm width), rectangular, in
Table 3. Meristic variation of *Tropidodipsas tricolor* sp. nov. and *Tropidodipsas papavericola* sp. nov.

| Specimen      | *T. tricolor* sp. nov. | *T. papavericola* sp. nov. |
|---------------|------------------------|----------------------------|
| Sex           |                        |                            |
| INIRENA 2800 | Male                   | Male                       |
| INIRENA 2798 | Female                 | Female                     |
| INIRENA 2801 | Male                   | Male                       |
| INIRENA 2802 | Male                   | Male                       |
| INIRENA 2803 | Male                   | Female                     |
| INIRENA 2804 | Male                   | Female                     |
| INIRENA 2810 | Male                   | Male                       |
| INIRENA 2805 | Male                   | Male                       |
| Snout-vent Length | 210                    | 210                        |
| Tail Length   | 67                     | 65                         |
| Total Length  | 277                    | 277                        |
| Head length   | 8.30                   | 13.62                      |
| Anterior Chinshield | Longer than wide | Longer than wide |
| Posterior Chinshield | Longer than wide | Longer than wide |
| Gulars       | 3                      | 3                          |
| Infrastr. in contact with 1st CS | 5                    | 5                          |
| Ventral Scales | 183                   | 183                        |
| Subcaudal Scales | 79 / 79               | 79 / 79                    |
| Pre-oculars (Side) | 1.1                   | 1.1                        |
| Post-oculars (Side) | 2.2                   | 3.3                        |
| Anterior Temporals (Side) | 1.1          | 1.1                        |
| Posterior Temporals (Side) | 2.2       | 2.2                        |
| Supralabials (Side) | 8.7                   | 7.8                        |
| Infrastrals (Side) | 8.8                   | 8.9                        |
| Post-nasals     | 0                     | 0                          |
| Pale Body Bands (Side) | 19.16             | 19.20                      |
| Pale Tail Bands (Side) | 9.9                  | 8.8                        |
| Black Body Blotches (Side) | 19.19            | 19.21                      |
| Black Tail Blotches (Side) | 8.8              | 8.8                        |
| Total Pale Body Blotch Count (Body & Tail) | 28.28          | 28.28                      |
| TL/TotL       | 0.24                   | 0.23                       |
| TL/SV/L       | 0.32                   | 0.30                       |
| Internal Length | 1.00                | 2.34                       |
| Internal Width | 1.60                  | 2.62                       |
| Loreal Length  | 0.9 / 1.0             | 1.16 / 1.10                |
| Loreal Width   | 1.0 / 1.0             | 1.26 / 1.12                |
| Loreal into orbit | No                  | No                         |
| Pre-frontal Length | 1.70                | 2.34                       |
| Pre-frontal Width | 2.00                | 2.64                       |
| Prefrontal into orbit | Yes              | Yes                        |
| Frontal Length | 2.80                  | 2.88                       |
| Frontal Width  | 2.40                  | 2.70                       |
| Parietal Length | 4.30                 | 5.54                       |
| Length Anterior Chinshields | 2.4 / 2.6     | 3.49 / 3.63                |
| Length Posterior Chinshields | 1.9 / 1.8 | 2.28 / 2.42                |
| Eye Diameter   | 2.10                  | 2.80                       |
| ED/HL         | 0.25                  | 0.21                       |
| Pre-Frontal Length/Width | 0.85            | 0.89                       |
| HL/SV/L       | 0.04                  | 0.04                       |
| Midbody-scale rows | 15                 | 15                         |
| Head scale rows | 15                   | 15                         |
| Caudal scale rows | 15                  | 15                         |
| Lateral compression | Yes = Extreme     | Yes = Extreme              |
| Anal plate     | Undivided             | Undivided                  |
| Penultimate supralabial enlarged | Yes        | Yes                        |
| First infralabials in contact | Yes     | Yes                        |
| Pale nuchal band | Yes                 | Yes                        |

median contact with each other and in lateral contact with postnasal, loreal, upper preocular and supraocular; frontal 1.1 times longer than broad, (4.4 mm long, 3.9 mm broad), pentagonal with angular tip posteriorly, in contact with prefrontals, supraoculars and parietales. Two moderately large preoculars on each side, upper larger than lower. Single supraocular large, in contact with upper preocular, prefrontal, frontal, parietales, and upper postocular. Two moderately large postoculars on each side, upper one larger than lower. Parietal 1.4 times as long as wide (6.0 mm long, 4.2 mm wide), the length of parietales 61% of HL, the common suture between parietales 4.4 mm, equal to frontal length. Nasal divided, prenasal slightly smaller than postnasal, with combined
Figure 4. *Tropidodipsas papavericola* sp. nov. Holotype (INIRENA 2801). Holotype (INIRENA 2801) in life (A); Ventral aspect of Holotype (INIRENA 2801) in preservative (B,E); Dorsal aspect of Holotype (INIRENA 2801) in preservative.
length of both nasals 2.6 mm, longer than loreal. Loreal small, slightly longer than high and almost square, 1.4 high, 1.7 mm long on left side, 1.6 mm high, 1.8 mm long on right side, not reaching orbit. Eye moderately large, 3.1 mm, 23% of HL. Supralabials 7 on both sides, first in contact with nasals, second and third in contact with loreal, fourth and fifth entering orbit, sixth largest and in contact with lower postocular, anterior and posterior temporals. Temporals 1 + 2. Five nuchal scales in contact with parietals.

Mental 2.4 times as broad as long (2.6 mm broad, 1.1 mm long), flat anteriorly, triangular posteriorly and separated from the anterior chinshields by the first pair of enlarged infralabials which are in contact with each other. Infracrals 7/8; on the left 1–5 in contact with anterior chinshields and fifth in contact with posterior chinshield; on the right 1–6 in contact with anterior chinshields and sixth in contact with posterior chinshield. Anterior chinshields irregular, both 2.3 times as long as wide (left 4.6 mm long and 2.0 mm wide, right 4.4 mm long and 1.9 mm wide). Left posterior chinshield 1.6 times as long as wide (3.5 mm long, 2.0 mm wide), and right posterior chinshield 1.3 times as long as wide (2.6 mm long, 2.0 mm wide). Four gular scales. Infracrals and scales in the chin region smooth. Dorsal scales in 15–15–15 rows, smooth throughout body; apical pits not evident. Ventral 180; anal plate single; 76 paired subcaudal scales. Body laterally compressed. Pupil elliptical.

Coloration in life of the holotype (Fig. 4A). Dorsal coloration of head, body and tail brown with 32/33 irregular dark brown bands on the body and neck which all have pale centers that are stippled with paler brown and pale cream. Pale brown nuchal band two body scales rows long but reduced to one at the mid-dorsal line; not involving the posterior part of the parietals. Dark bands on body and tail 3–7 scale rows long, widest mid-dorsally, and alternating on posterior half of body. Head coloration predominately dark brown on dorsal surface and with light brown mottling and pale cream on rostral and anterior labials. First three supraflabials bicolor pale cream and dark brown, posterior two supraflabials bicolor dark brown with pale brown. Ventral coloration of head pale cream with irregular dark brown blotches. Ventral coloration of body pale yellowish cream, with dark and pale bands incomplete on venter. The ventral pattern consists of longitudinal rows of dark spots that connect to...
form irregular longitudinal lines. Anal plate bicolor, dark brown and pale cream. All 15 dark bands on tail incomplete ventrally, dark coloration forming longitudinal lines ventrally on tail. Tail tip pale cream with black tip. Pupil black, iris dark chocolate-gray.

**Coloration in preservative of the holotype** (Figs 4B–E). General coloration bicolor. Dorsal surfaces of head, body and tail predominately dark brown with darker brown-black bands. Pale bands brown-gray, but whitish in a few places where *stratum corneum* has been removed (snake was going into shed when preserved). The dark bands remain brown-black and the pale cream faded to white.

**Variation.** Meristic variation of five available specimens given in Table 3. Variation in color and pattern detailed below (Fig. 5).

**Color in life.** A sub-adult male paratype from above San Luis La Loma, Municipio de Técutpan de Galeana, Guerrero (INIRENA 2802) had a pale brown body coloration with 31/29 dark brown bands on body and neck posterior to pale nuchal band and 14 on the tail. This specimen has light centers in all dark body bands, albeit limited to the lateral portions only. This implies that the irregular dark banding with light centers may be subject to ontogenetic variation, as this specimen is intermediate in size and pattern between the juvenile INIRENA 2810 and the large adult holotype INIRENA 2801. Neither the pale body nor the dark body bands, are complete ventrally, reaching only to the lateral edges of the ventrals. The venter was cream or pale tan mid-ventrally with two irregular black spots that do not form longitudinal rows or to pale nuchal band and 14 on the tail. This specimen has light centers in all dark body bands, albeit limited to the lateral portions only. This implies that the irregular dark banding with light centers may be subject to ontogenetic variation, as this specimen is intermediate in size and pattern between the juvenile INIRENA 2810 and the large adult holotype INIRENA 2801. Neither the pale body nor the dark body bands, are complete ventrally, reaching only to the lateral edges of the ventrals. The venter was cream or pale tan mid-ventrally with two irregular black spots that do not form longitudinal rows. Furthermore, the first well-defined dark band on the neck does not have a pale center, and on the second dark body band the pale center is interrupted mid-dorsally. The venter is white and covered with irregular black spots that do not form longitudinal rows of spots nor are fused into lines and dashes (Fig. 5C).

A juvenile male paratype from above San Luis La Loma, Municipio de Técutpan de Galeana, Guerrero (INIRENA 2804) has 29/27 dark bands on the body posterior to the pale nape band and 12 on the tail. This specimen differs from the holotype because it has a relatively well-defined pattern of dark and pale bands with regular edges. Furthermore, the first well-defined dark band on the neck does not have a pale center, and on the second dark body band the pale center is interrupted mid-dorsally. The venter is white and covered with irregular black spots that do not form longitudinal rows of spots nor are fused into lines and dashes (Fig. 5A, B). The venter was cream or pale tan mid-ventrally with two irregular black spots that do not form longitudinal rows. Furthermore, the first well-defined dark band on the neck does not have a pale center, and on the second dark body band the pale center is interrupted mid-dorsally. The venter is white and covered with irregular black spots that do not form longitudinal rows of spots nor are fused into lines and dashes (Fig. 5C).

**Color in preservative.** An adult male paratype from above Tepatlan de Galeana, Guerrero (INIRENA 2801) appears to have 14 maxillary teeth visible, however we did not remove the maxillary arch to avoid damaging the specimen. We observed *in situ* the maxillary arch and dentition of the holotype (INIRENA 2801): it has 14 maxillary teeth and 20–22 dentary teeth.

A sub-adult male paratype, from above Tepatlan de Galeana, Guerrero (INIRENA 2805) has 26 dark bands on body and neck posterior to the pale nuchal band, all with faded pale centers in the dark body bands, most of them extensive but incomplete mid-dorsally. It has 12 dark bands on the tail and a dark tail tip. Ventral coloration white with dark gray mottling, with no dark or pale dorsal bands complete on venter, but the dense mottling makes the dark dorsal bands on posterior portion of body connect in an alternating undulated pattern.

**Dentition.** The holotype (INIRENA 2801) appears to have 14 maxillary teeth visible, however we did not remove the maxillary arch to avoid damaging the specimen. We observed *in situ* the maxillary arch and dentition of the holotype (INIRENA 2801): it has 14 maxillary teeth and 20–22 dentary teeth.

An adult male paratype (INIRENA 2805) has 17 maxillary teeth (counting empty sockets) on the right side. We counted 22 dentary teeth on the right dentary of this same specimen.

**Distribution, habitat and ecology.** This species appears to be restricted to moderate elevations in the Sierra Madre del Sur in central Guerrero in two distinct areas (Fig. 10). Habitats include cloud forest, mesic pine-oak woodland, tropical evergreen forest, tropical semi-deciduous forest and combinations of these assemblages. Known from around 1600–2200 m a.s.l. in the Sierra de Tepatlan de Galeana, Sierra de Atayac and above Acapulco, in the foothills in the area encompassed between Acahuizotla and El Treinta. Specimens were found at night during the rainy season on low to medium (1.5–3.0 m) vegetation or crossing the road.

**Etymology.** The specific epithet *papaver + cola* refers to the living among poppy plants (genus *Papaver*) which are illegally planted throughout the range of this species for the extraction of opium gum.

**Discussion**

**Generic identity of Tropidodipsas annulifera, T. sartorii and Sibon sanniolus**

Our molecular analyses confirm the results of Sheehy (2013) that some species previously included in *Tropi-
**Figure 6.** Bayesian phylogenetic inference of members of the Dipsadidae based on two mitochondrial loci and two nuclear loci. All nodes with support of less than 0.5 are collapsed, while those with posterior support equal to 1 are marked with a black dot. New species described here in bold.

dodipsas and Sibon (T. annulifera, T. sartorii and S. sanniolus) are actually more closely related to members of the genus Geophis than to other snail-suckers. However, we disagree with Sheehy’s (2013) recommendation that new genera should be erected for each of these species. Instead, we consider it more appropriate for taxonomic stability to regard these three species as members of the genus Geophis, with the following new combinations: Tropidodipsas annulifera Boulenger 1894 = Geophis annuliferus comb. nov. (Boulenger, 1894); Tropidodipsas sartorii Cope 1863 = Geophis sartorii comb. nov. (Cope 1863); and Sibon sanniolus (Cope 1866) = Geophis sanniolus comb. nov. (Cope 1866). We refrain from assigning the two species of the Tropidodipsas sartorii species group (sensu Kofron 1988; Smith et al. 2005) for which we do not have genetic material (T. repleta and T. zweifeli) to Geophis, and provisionally retain them in Tropidodipsas, although it has been suggested by previous authors that a close phylogenetic relationship between these taxa is likely (e.g., Kofron 1988). It should be noted that synonyms of both G. annuliferus comb. nov. and G. sartorii comb. nov. have previously been described in the genus Geophis. Geophis tecpanecus Dugès 1896 was synonymized with G. annuliferus comb. nov. by Scott (1967) and Geophis annulatus Peters 1870 was synonymized with G. sartorii comb. nov. by Boulenger (1894).
Content of the genus *Tropidodipsas*

As revised here, the genus *Tropidodipsas* thus comprises eight species: *Tropidodipsas fasciata* Günther 1858, *T. fischeri* (Boulenger, 1894), *T. guerreroensis* Taylor, 1939 (see comment below), *T. papavericola* sp. nov., *T. philippi* (Jan, 1863), *T. repleta* Smith, Lemos-Espinal, Hartman & Chiszar, 2005, *T. tricolor* sp. nov. and *T. zweifeli* Liner & Wilson, 1970. The two species described herein are each other’s closest relative and comprise a clade together with samples assigned to *T. fasciata* from the Yucatán Peninsula (Fig. 7B). These two species together with *T. philippi*, *T. guerreroensis*, and the various populations assigned to *T. fasciata* form a species group that we continue to recognize as the *T. fasciata* species group (Kofron 1987, Wallach 1995). The phylogenetic relationships of *Tropidodipsas repleta* and *T. zweifeli* are unknown as no genetic data currently exist for these species. With the removal of *Geophis annuliferus* new comb. and *G. sartorii* new comb. these two species become unassignable to a species group. Genetic material of these two species is needed to determine whether their relationship is indeed with *Tropidodipsas*, or whether they should also be placed in *Geophis*. The generic affinities of *Dipsas gaigae* (Oliver, 1937) and *Tropidodipsas fischeri* (Boulenger, 1894) are not clear as they render their respective genera paraphyletic, and these two species probably deserve their own monotypic genera as suggested by previous authors (Kofron 1985b; Fernandes 1995; Sheehy 2013). However, this is beyond the scope of this paper and we provisionally retain them in their respective genera.

Identity of *Tropidodipsas guerreroensis*

*Tropidodipsas guerreroensis* was described by Taylor (1939) as a member of the genus *Tropidodipsas* that he considered closely related to *T. fasciata*. Later, Álvarez del Toro and Smith (1956) relegated it to a subspecies of *T. fasciata*, an arrangement followed by Kofron (1980, 1987). Mertz et al. (2010) reported a specimen (UTADC 3701 = JRV-31) purportedly of this form from western Guerrero as a range extension of *T. fasciata* in the southern Oaxaca. The genetic distances of the mitochondrial gene cytb between the westernmost specimens of *T. guerreroensis* on Hwy. 134 in Guerrero and the easternmost specimen from near Candelaria Loxicha in Oaxaca is less than 6%, suggesting that all four specimens belong to one cohesive lineage, albeit with considerable intraspecific divergence, probably the result of isolation-by-distance.

In contrast, the easternmost individual of *T. guerreroensis* (Candelaria Loxicha, Oaxaca) has a genetic distance of 12.4–12.5% compared to nearby *T. fasciata* from various localities in the Isthmus of Tehuantepec, Oaxaca (Fig. 8A, B). For comparison, the western population of *T. guerreroensis* (Hwy. 134, Guerrero) has a genetic distance of 6.9–8.5% from the nearest sampled *T. philippii* in Michoacán, which suggests a closer relationship to *T. philippii* than to *T. fasciata*, as shown by our phylogenetic analyses, or recent gene flow. Thus, we suggest that *T. guerreroensis* represents a diagnosable, monophyletic species that is closest related to *T. philippii*.

These results show that among Mexican snail-suckers inter-population intra-specific genetic distances of the cytb mitochondrial gene range from 3–7%, whereas inter-specific genetic distances of closely related species (such as *T. philippii*, *T. guerreroensis*, and *T. fasciata*) range between 7–10% and other species have genetic distances >10%. It is notable that in our results intraspecific distances of cytb are less than 0.7% in *T. papavericola* sp. nov., less than 0.4% in *T. fasciata*, but 3–6% in *T. guerreroensis*.

Unassigned museum specimens from southern Oaxaca

Kofron (1980) described a specimen (LACM 104321) from the Sierra Madre del Sur of Oaxaca as “*Sibon sp. cf. philippii*”. He apparently considered the specimen to be of enough significance that he provided a detailed description, including scalation and color pattern, and even gave comparisons of how it differed from other *Tropidodipsas philippii*. He also recognized the uniqueness of the habitat compared to that of other *T. philippii*, but he refrained from designating a name for this form. Later, Kofron (1987) included this specimen, along with three others from Oaxaca (KU 137655, UCM 49372, UIMNH 73681) in his “*Sibon philippii* (= *Tropidodipsas philippii*). He did not give any further explanation as to why he placed these specimens in *T. philippii*, and the scale counts and ranges given by him for *T. philippii* include these specimens.

It is important to note that these southern specimens are over 750 km from the nearest populations of *T. philippii* in Michoacán, and originate from elevations between...
1600–2100 m a.s.l., whereas T. philippii is known from elevations usually below 1500 m a.s.l. We have reviewed detailed photographs of three of these specimens (LACM 104321, KU 13766, UCM 49372) and it is likely that these high-elevation populations are referable to T. papa
ericola sp. nov. or represent an undescribed species, closely related to it. We were not able to examine the other specimen (UIMNH 73681) from southern Oaxaca due to institution closures related to the Covid-19 pandemic. Unfortunately, genetic material is not available for the high-elevation populations of Tropidodipsas from the Sierra Madre del Sur of Oaxaca. Future collecting is necessary to properly assign these populations to a species.

The occurrence of Geophis sartorii new comb. along the west coast of Mexico

Sheehy (2013) included a specimen of Geophis sartorii new comb. (JAC 30401) purportedly from the vicinity of El Tuito, Jalisco, in his phylogenetic analyses. Geophis sartorii new comb. is restricted to eastern Mexico, so a specimen of this species from Jalisco greatly increased the known range of the species. The sample did indeed group with G. sartorii new comb. in Sheehy’s genetic analysis and not with the superficially similar G. annuliferus comb. nov., which is known from that region of Jalisco. We inquired about this specimen at the UTA collection and were informed that it was collected by Mr. Paulino Ponce-Campos. We contacted Mr. Ponce-Campos about the provenance of this specimen, and he told us that he specifically remembers providing the UTA field crew with a specimen of G. sartorii new comb. that he had collected in the state of San Luis Potosí and given them while they were in Jalisco.

Considering that Sheehy (2013) does not list any Geophis sartorii new comb. as coming from San Luis Potosí, we suggest it is very likely that this is the correct provenance of JAC 30401 and the locality listed by Sheehy (2013) was confused with another specimen with origins in Jalisco. This is further supported by the fact that the genetic distances of a fragment of the mitochondrial gene cytb between JAC 30401 and two G. sartorii new comb. (INIRENA 12783–84) which we collected in San Luis Potosí is 0.0. We have included this specimen (JAC 30401) as “locality unknown” and strongly doubt that G. sartorii new comb. is present in the state of Jalisco. However, Geophis sartorii macdougalli new comb. was
recently reported from the Pacific versant of Guerrero by Blancas-Hernández et al. (2019). It would be interesting to verify the identity of this Guerreran population with molecular tools and assess how it is related to populations of *G. sartorii* new comb. on the Atlantic versant of Mexico and around the Isthmus of Tehuantepec. If the Guerrero population indeed belongs to *G. sartorii* new comb., then the presence of additional populations of *G. sartorii* new comb. on the Pacific versant of Mexico can be expected.

**Conservation priorities for the new *Tropidodipsas***

The species of *Tropidodipsas* described above are the two species with the smallest known range in the genus, and likely are also the two species needing the most conservation attention. *Tropidodipsas papavericola* sp. nov. can be considered micro-endemic. It is known from only one biogeographical formation (#48 – Guerreran Sierra Madre del Sur Mixed Temperate Woodland) as defined by Grünwald et al. (2015). While it is currently known from four distinct localities, all fall within this biogeographical formation or right at its lowermost limit.

Widespread illegal logging is present at two of the localities (pers. obs.) and illicit poppy farming is present at or near all four localities. As poppy prices drop due to international supply and demand, and competition from synthetic drugs such as fentanyl, illicit poppy farmers tend to fluctuate to other businesses. This often leads to illegal logging (pers. obs.) and thus widespread habitat destruction. We suggest that this new species be awarded the highest conservation category possible by the Mexican government, and that studies are undertaken to determine its vulnerability to the habitat destruction caused by illegal logging. Also, further sampling should focus on southern Oaxaca to determine whether populations from the Sierra Madre del Sur above San Gabriel Mixtpec are assignable to this species.

*Tropidodipsas tricolor* sp. nov. is apparently widely distributed along the windward slopes of the Sierra Madre del Sur of Guerrero and Oaxaca. It has been collected in three biogeographical formations (#12 – Guerreran Tropical Dry Forest & Savanna; #48 – Guerreran Sierra Madre del Sur Mixed Temperate Woodland; #49 – Malinaltepec - Putla Sierra Madre del Sur Mixed Temperate Woodland) as defined by Grünwald et al. (2015). The type locality experiences small-time agricultural disturbance and some logging.
Figure 9. Similar looking but not closely related snail-suckers from southern Mexico. *Sibon nebulatus* from Municipio de Ixtlahuacán, Colima (A); *Sibon nebulatus* from Municipio de Las Margaritas, Chiapas (B); *Tropidodipsas fischeri* from Municipio de Rayón, Chiapas (C); *Tropidodipsas fischeri* from Municipio de Unión Juárez, Chiapas (D).

Figure 10. Distribution map of species in the *Tropidodipsas fasciata* species group in Mexico. Triangles represent type localities, type localities are only mapped when known with certainty.
but for now seems to be free of any major threats to the persistence of the habitat. The two localities in the Sierra Madre de Guerrero, near el Páezclár (Municipality of Leonardo Bravo) and near La Cienega (Municipality of Malinaltepec), are threatened by logging, but it does not appear as widespread as is the case with T. papavericola sp. nov. The only known locality of T. tricolor sp. nov. in Oaxaca is threatened by deforestation for small-scale agriculture. We did not see signs of illegal logging or drug cultivation in this area, however in general the habitat in this area is more disturbed than in the vicinity of the type locality.

The Environmental Vulnerability Score (EVS) was developed by Wilson and McCranie (1992) for use with amphibians in Honduras. The EVS system was later applied to the amphibians and reptiles of Mexico by Wilson et al. (2013). It was further modified by Porras et al. (2013) to better apply to animals outside of Honduras. Grünwald et al. (2015) applied the EVS system as modified by Porras et al. (2013) and defined Biogeographical Formations specific to reptiles and amphibians for the country of Mexico. These more-inclusive Biogeographical Formations replaced the “Forest Formations” initially outlined by Wilson and McCranie (1992) in their application to Honduras. Herein we apply the EVS system as outlined by Porras et al. (2013) and Grünwald et al. (2015) to the Tropidodipsas fasciata species group as defined above.

\[
\begin{align*}
T. fasciata & = 3 + 3 + 4 = 10 \\
T. guerrerensis & = 4 + 6 + 4 = 14 \\
T. papavericola sp. nov. & = 4 + 8 + 4 = 16 \\
T. philippii & = 3 + 7 + 4 = 14 \\
T. tricolor sp. nov. & = 4 + 6 + 5 = 15
\end{align*}
\]

The IUCN categories for assigning conservation status are the most commonly used scheme to assess the degree of extinction risk for taxa at the species level (Porras 2013). The criteria used for this assessment are stipulated in the Guidelines for Using IUCN Red List Categories and Criteria (Version 8.1; August 2010). We evaluate the species of the T. fasciata species group (as defined herein) below:

\[
\begin{align*}
T. fasciata & = \text{Least Concern} \\
T. guerrerensis & = \text{Least Concern} \\
T. papavericola sp. nov. & = \text{Near Threatened} \\
T. philippii & = \text{Least Concern} \\
T. tricolor sp. nov. & = \text{Least Concern}
\end{align*}
\]

Our evaluation of T. papavericola sp. nov. as near threatened is based on the limited extent of its known distribution. While known from more localities than just the type locality, all known localities fall within a 90 km radius of the type locality in the same physiographic province. This fact, coupled with the moderate habitat destruction for small scale agriculture present at all collecting localities, supports our evaluation of this species as “near threatened”.

**Funding**

Partial funding for the field work was provided by Consejo de Ciencia y Tecnología del Estado de Durango (COCYTED) grant number COCYTED 2018 01: 15088. Further funding for field work was provided by Herp.mx A.C. and Biencom Real Estate. Funding for molecular data generation was provided by the University of South Alabama through startup funds to JLS. IMH was funded by a Summer Undergraduate Research Fellowship from the Office of Undergraduate Research at the University of South Alabama to conduct lab work.

**Competing interests**

The authors have declared that no competing interests exist.

**Acknowledgements**

First and foremost, we thank our field crew for their courageous enthusiasm to go out in the field in Guerrero, including but not limited to Brandon T. La Forest, André J. Grünwald, Ámbar Lanomy Grünwald, Karen I. Morales-Flores, Janelle Morales-Flores, Carmen Mendoza-Portilla, Nadia Pérez-Rivera, Alejandro Lara, William Mertz and Tziuhtécatl Sánchez Luna. We thank Peter Heimes, Tziuhtécatl Sánchez Luna and Abelino Uriostegui for graciously providing photos of live individuals in the field. Nefti Camacho (LACM), Emily Braker and Christy McCain (UCM), Jeff Weinell and Rafe Brown (KU), David Kizirian (AMNH) and Toby Hibbits (TCWC) all provided detailed photographs of preserved specimens under their care. We are indebted to María de los Ángeles Palma Irizarry of the Secretaría de Medio Ambiente y Recursos Naturales (SEMARNAT) for providing collecting permits (SGPA/DGVS/002288/18) and to Dr. Juan Miguel Borja-Jiménez for extending his collecting permit to us. We thank Robert Hansen, Arthur Tiutenko, Günter Gollmann and two anonymous reviewers for reviewing our manuscript and providing valuable input.

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**Appendix 1**

**Specimens examined**

*Dipsas brevifacies.— Mexico: Yucatan • 36.4 km S of Valladolid on Hwy. 295, Municipio de Tixcacalcuipul, 25 m a.s.l., INIRENA 2791, CIG 1841 • 1.4 km E of fair complex on Calle 253, Merida, Municipio de Merida, 11 m a.s.l., INIRENA 2792, CIG 1844.

*Dipsas gaigeae.— Mexico: Colima • 3.0 km N of Ixtlahuacán Rd. on Hwy. 54 frontage road, Municipio de Tecomán, 322 m a.s.l., JAC 28327 • 6.4 km N of Ixtlahuacán Rd. on Hwy. 54 frontage road, Municipio de José Azueta, 130 m a.s.l., JRV 0030.

*Geophis sieboldi* comb. nov.— Mexico: Colima • El Mixcuate, on Colima–Minatitlán Road, Municipio de Villa de Álvarez, 569 m a.s.l., JAC 30142 • Guerrer: 24.1 km NE of Hwy. 200 on Hwy. 134, Municipio de José Azueta, 330 m a.s.l. JAC 27792.

*Geophis bicolor.— Mexico: Colima • 10 km (airline) NNW of Quesería, Municipio de Cuauhtémoc, 2128 m a.s.l., INIRENA 2795–97, CIG 1786–88 • Jalisco:...
3 km (airline) NNW of Cumbre de Guadalupe, Municipio de Talpa de Allende, 2095 m a.s.l., INIRENA 2793–94, CIG 1576–77 • 9.2 km SW of Talapa on road to San Gabriel, Municipio de Tapalpa, 2012 m a.s.l., INIRENA 2808, CIG 1850 • Plan de Cervantes, on Valle de Juárez – Santa María del Oro Rd., Municipio of Quiputan, 2291 m a.s.l. INIRENA 2809, CIG 1851 • Michoacán: 3.2 km NW of Apo, Municipio de Tancitaro, 2010 m a.s.l., JAC 24684.

*Geophis nigrocinctus.* — Mexico: Jalisco • Cerro Tettilla, 20.1 km (airline) W of Talpa de Allende, Municipio de Talpa de Allende, 2466 m a.s.l., JAC 30704 • Michoacán: Between Paso Malo and Rancho Las Torrecillas, on Coacóman-Dos Aguas Rd., Municipio de Coacóman de Vázquez-Pallares, 2115 m a.s.l., CIG 0568.

*Geophis omnilemanus.* — Mexico: Guerrero • Omiltemi, Municipio de Chilpancingo de los Bravo, 2115 m a.s.l., ENS 11496.

*Geophis sanniolus* comb. nov. — Mexico: Yucatán • 2.2 km E of Homún on road to Huhl, Municipio de Homún, 16 m a.s.l., INIRENA 2790, CIG 1842 • 12.7 km S of Hwy. 180. 180 on road to Táahdzibichén, Municipio de Yaxcabá, 30 m a.s.l., JAC 24409 • 2.8 km S of Tixcacalcupul on Hwy. 295, Municipio de Tixcacalcupul, 26 m a.s.l., INIRENA 2789, CIG 1839.

*Geophis sartorius* comb. nov. — Mexico: San Luis Potosí • 1.1 km SW of Huichihuayan, on road to El Nacimiento, Municipio de Huehuetlán, 90 m a.s.l., INIRENA 2784, CIG 1758 • 2.0 km NE of Xilitla on Hwy. 120, Municipio de Xilitla, 552 m a.s.l., INIRENA 2785, CIG 1759 • 2.2 km NE of Xilitla on road to El Túnel, Municipio de Xilitla, 679 m a.s.l., INIRENA 2783, CIG 1518 • *Yucatan:* 6.7 km S of Tixcacalcupul on Hwy. 295, Municipio de Tixcacalcupul, 26 m a.s.l., INIRENA 2786, CIG 1840 • Unknown: JAC 30401.

*Geophis tarascae.* — Mexico: Colima • 7 km (airline) NNW of Montitlán, Municipio de Cuahutémoc, 1846 m a.s.l., INIRENA 2807, CIG 1631 • Jalisco: 1.9 km S of El Montoso, Municipio de Quiputan, 1969 m a.s.l., INIRENA 2806, CIG 1372 • Michoacán: 2.5 km S of southern edge of Uruapan, on Hwy. 37 libre toward Lombardía, Municipio de Uruapan, 1563 m a.s.l., JAC 24692.

*Sibon nebulatus.* — Mexico: Chiapas • 0.8 km SW of Ejido Morelos, Municipio de Huixtla, 1185 m a.s.l. INIRENA 2788, CIG 0788 • Colima: Road from Comala to Minatitlán, 739 m a.s.l., JAC 30102 • Hwy. 54 frontage road, near La Salada, 301 m a.s.l., JAC 30124 • Michoacán: 9.4 km NNW of Caleta de Campos, Municipio de Aguila, 15 m a.s.l., INIRENA 2787, CIG 1481.

*Tropidodipsas fasciata.* — Mexico: Chiapas • 13.5 km (airline) NW of Rizo de Oro, Municipio de Cintalapa, elev. unknown, JAC 22920 • Oaxaca: 33.6 km SSE Matias Romero, Municipio de Asunción Ixaltepec, LACM 103765 • 17.3 km W of Zanatepec, Municipio de Santiago Niltepec, 56 m a.s.l., LACM 38212 • 3.2 km N of Tehuantepec, Municipio de Santo Domingo Tehuantepec, 32 m a.s.l., LACM 114067 • 60.8 km WNW of Tehuantepec on Hwy. 190, Municipio de Magdalena Tequisistlán, LACM 74042 • 51.2 km NW of Magdalena Tequisistlán turnoff on Hwy. 190 to Oaxaca, Municipio de Nejapa de Made-ro, LACM 38211 • 2.9 km W of Hwy. 195 on road to Almoloyas, Municipio de El Barrio de la Soledad, 270 m a.s.l., JAC 30740.

*Tropidodipsas cf. fasciata.* — Mexico: Tamaulipas • Gómez-Farias Rd., at the Ojo de Agua turnoff, Municipio de Gómez Farias, 243 m a.s.l., CIG 0819 • *Yucatan:* 15.4 km NW of Huenumá on road to Sisal, Municipio de Huenumá, 5 m a.s.l., INIRENA 2780, CIG 1843.

*Tropidodipsas fischeri.* — Mexico: Chiapas • Selva Negra, Municipio de Rayón, 1895 m a.s.l., CHFCB-0352 • Chichihuites, Municipio de Unión Juárez, 2090 m a.s.l., CHFCB-0332, 0335.

*Tropidodipsas guevaraensis.* — Mexico: Guerrero • Acahuizotlán, 853 m a.s.l., KU 61242, TCWC 7477–80 • 1.6 km W Acahuizotlán, 853 m a.s.l., TCWC 7481–82 • 2.8 km SW of Rincón de la Parotas, on Atoyac de Álvarez to Puerto del Gallo Rd., Municipio de Atoyac de Álvarez, 273 m a.s.l., INIRENA 2781, CIG 1857 • Oaxaca: 12 km S of Candelaria Loixcha on Hwy. 175, Municipio de Candelaria Loixcha, 291 m a.s.l., JAC 24267 • 1.6 km SE of Cacahuatpec, Municipio de San Juan Cacahuatpec, 360 m a.s.l., UIMNH 52958.

*Tropidodipsas papavericola* sp. nov. — Mexico: Guerre-ro • 12.5 km S of Puerto del Gallo on road from Nuevo Delhi to Puerto del Gallo, Municipio de Atoyac de Álvarez, 1914 m a.s.l., INIRENA 2801, CIG 1495 • 18.1 km S of Puerto del Gallo on road from Nuevo Delhi to Puerto del Gallo, Municipio de Atoyac de Álvarez, 1654 m a.s.l., INIRENA 2802, CIG 1496 • 5 km S of La Laguna, on road from San Luis La Loma to Bajitos de la Laguna, Municipio de Técpan de Galeana, 1686 m a.s.l., INIRENA 2803 CIG 1502 • Bajitos de la Laguna, Municipio de Técpan de Galeana, INIRENA 2804, CIG 1632 • Jaguar Research Facility, Municipio de Técpan de Galeana, INIRENA 2805, CIG 1457 • 4.2 km S of La Laguna, on San Luis San Pedro – La Laguna Rd., Municipio de Técpan de Galeana, 1620 m a.s.l., INIRENA 2810, JRV 0362.

*Tropidodipsas philippii.* — Mexico: Colima • 2 km E of Hwy. 54 frontage road on road to Ixtlahuacán, Municipio de Ixtlahuacán, 346 m a.s.l., INIRENA 2782, CIG 1503 • San Gabriel, Municipio de Ixtlahuacán, 490 m a.s.l., CIG 0902 • 2 km S of Minatitlán, on Hwy. 98, Municipio de Minatitlán, 712 m a.s.l., JAC 28262 • 16–24 km SW Colima, LACM 59146 • *Jalisco:* 8.6 km N of El Tuito on Hwy. 200, Municipio de Cabo Corrientes, 702 m a.s.l., ENS 11639 • *Nayarit:* Las Mesas, Municipio de Tepic, 338 m a.s.l., JAC 24811 • Michoacán: 2.4 km N of Hwy. 200 on road to Ostula, Municipio de Aquila, 138 m a.s.l., JAC 27923 • *Sinaloa:* 10.1 km NE of Concordia turnoff on Hwy. 40-D cuota, on road to Durango,
Municipio de Concordia, 287 m a.s.l., JAC 30601 • 85.3 km N of Mazatlán, LACM 7118 • 57.9 km N of Mazatlan, KU 73640; 50.9 km N Mazatlan, LACM 7119 • between Escuinapa and Palmilla, LACM 7117 • Teacapan, LACM 7116.

**Tropidodipsas sp. cf. philippii.** — **Mexico: Oaxaca** • 5.1 km S of Jalatengo, Municipio de Candelaria Loxicha, 1390 m a.s.l., KU 137655 • Santa Rosa, Distrito Lachao, Municipio de San Juan Lachao, UCM 49372 • 27.4 km S of Juchatengo, Municipio de San Juan Lachao, 1829 m a.s.l., LACM 104321.

**Tropidodipsas tricolor** sp. nov. — **Mexico: Guerrero** • 1.5 km east of Río Verde, Municipio de Atoyac de Álvarez, 971 m a.s.l., INIRENA 2800, CIG 1837 • 4.5 km NW of Mixtecapa, on road to Malinaltepec, Municipio de Malinaltepec, 1815 m a.s.l., INIRENA 2798, CIG 1863 • **Oaxaca**: 26 km N of Putla Villa de Guerrero, on Putla Villa de Guerrero - Oaxaca Hwy., Municipio of Putla de Guerrero, 1785 m a.s.l., INIRENA 2799, CIG 1596.

### Supplementary material 1

**Table S1**

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Data type: species data

Explanation note: Morphological differences between the new species of *Tropidodipsas* and other snail-suckers from Mexico.

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