Phylogenomics of paleoendemic lampshade spiders (Araneae, Hypochilidae, Hypochilus), with the description of a new species from montane California

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Abstract

Hypochilus is a relictual lineage of Nearctic spiders distributed disjunctly across the United States in three montane regions (California, southern Rocky Mountains, southern Appalachia). Phylogenetic resolution of species relationships in Hypochilus has been challenging, and conserved morphology coupled with extreme genetic divergence has led to uncertain species limits in some complexes. Here, Hypochilus interspecies relationships have been reconstructed and cryptic speciation more critically evaluated using a combination of ultraconserved elements, mitochondrial CO1 by-catch, and morphology. Phylogenomic data strongly support the monophyly of regional clades and support a ((California, Appalachia), southern Rocky Mountains) topology. In Appalachia, five species are resolved as four lineages (H. thorelli Marx, 1888 and H. coylei Platnick, 1987 are clearly sister taxa), but the interrelationships of these four lineages remain unresolved. The Appalachian species H. pococki Platnick, 1987 is recovered as monophyletic but is highly genetically structured at the nuclear level. While algorithmic analyses of nuclear data indicate many species (e.g., all H. pococki populations as species), male morphology instead reveals striking stasis. Within the California clade, nuclear and mitochondrial lineages of H. petrunkevitchi Gertsch, 1958 correspond directly to drainage basins of the southern Sierra Nevada, with H. bernardino Catley, 1994 nested within H. petrunkevitchi and sister to the southernmost basin populations. Combining nuclear, mitochondrial, geographical, and morphological evidence a new species from the Tule River and Cedar Creek drainages is described, Hypochilus xomote sp. nov. We also emphasize the conservation issues that face several micro-endemic, habitat-specialized species in this remarkable genus.
Keywords
Conservation, mountains, multispecies coalescent, short-range endemism, Sierra Nevada, southern Appalachians, taxonomic over-splitting, ultraconserved elements

Introduction

Discovering and delimiting cryptic species boundaries is, almost by definition, challenging. When the multispecies coalescent (MSC) is applied to species delimitation, species boundaries are explored by estimating gene trees and accounting for species tree/gene tree discordance using MSC models. Critical to this approach is discerning the boundary between population-level versus species-level divergence, as a core assumption of most MSC models is that species are panmictic and without population structure (Degnan and Rosenberg 2009). It is now well-established that, in systems with high natural population genetic structuring, MSC-based delimitation methods can conflate structure at the population level with divergence at the species level (Carstens et al. 2013; Sukumaran and Knowles 2017; Chambers and Hillis 2019; Mason et al. 2020). Many empirical studies indicate that, used alone, MSC methods can drastically over-split taxa (e.g., Niemiller et al. 2012; Satler et al. 2013; Hedin et al. 2015; Derkarabetian et al. 2019; Hundsdörf er et al. 2019).

Population genetic structuring is rather ubiquitous in nature. Species with strict or semi-strict habitat or microhabitat preferences will naturally occur discontinuously over a landscape. Combine this natural habitat fragmentation with limited dispersal ability, and populations will evolve to be genetically different, to various degrees (Templeton et al. 1990; Coates et al. 2018; Marshall et al. 2021). Under some circumstances, arrays of parapatric or allopatric populations which are diverging genetically might remain morphologically quite similar, particularly when microhabitat preferences are strong (Stockman and Bond 2007; Bernardo 2011; Fišer et al. 2018). This combined suite of circumstances corresponds to what we refer to as a “no gene flow” or “non-adaptive radiation” speciation model (Gittenberger 1991; Kozak et al. 2006). Non-adaptive speciation is quite common in nature (e.g., Reilly and Wake 2005; Leavitt et al. 2007; Emata and Hedin 2016; Singhal et al. 2018; Derkarabetian et al. 2022), and challenges species delimitation. This species delimitation problem lies at one end of a spectrum of difficult scenarios for species delimitation (a high gene flow model representing an opposite, but equally challenging, scenario). Non-adaptive speciation represents a conundrum for species delimitation, as genetic data combined with many currently available models will likely over-split taxa, while other lines of evidence needed to confirm or reject this over-splitting (e.g., morphological evidence, etc.) is difficult to uncover in these same taxa (Derkarabetian et al. 2022).

The spider genus Hypochilus Marx, 1888 represents a challenging system for species delimitation, combining allopatric geographic distributions, morphological conservatism, and high genetic structuring. Hypochilus is a Nearctic genus representing one of two described genera in the family Hypochilidae, a family of true spiders which retain
many interesting plesiomorphic traits (Forster et al. 1987; Alberti and Coyle 1991; Catley 1994). Commonly known as lamplight spiders, Hypochilus spiders are microhabitat specialists occurring in shaded, mesic, rock outcrop habitats (Catley 1994; Hedin and Wood 2002; Keith and Hedin 2012). Hypochilus includes ten described species from three disjunct montane regions: the southern Appalachians, the southern Rocky Mountains, and the California mountains (Fig. 1). Described species are mostly exclusively allopatric, and within montane regions where species are in close geographic proximity, occur in parapatric patchworks. These spiders are textbook examples of so-called short-range endemic (SRE) taxa, including species with naturally small geographic distributions (often defined as less than 10,000 km²; Harvey 2002, Harvey et al. 2011). Several Hypochilus species occupy severely limited geographic distributions; for example, H. bernardino Catley, 1994 is only known from a handful of locations in a single mountain range (San Bernardino mountains of southern California). Multiple restricted-distribution species also warrant conservation attention, particularly in the face of climate change; this conservation focus also highlights the need for rigorous species delimitation (e.g., Hedin 2015).

As spiders with extraordinarily low vagility, one would expect deeper phylogenetic relationships in Hypochilus to closely mirror geography, with phylogenetic predictions following geography. However, previous studies have suggested that this may be an oversimplification. Both morphological and mitochondrial data suggest that the geographically separated California and Appalachian mountain faunas are sister lineages (Catley 1994; Hedin 2001), although this inferred relationship is sensitive to varying combinations of data and analyses (Hedin 2001). Also, monophyly of the California fauna has been questioned, as both morphological and mitochondrial tree topologies sometimes recover an Appalachian clade within a larger paraphyletic Californian group (Hoffman 1963; Hedin 2001). Given the impressive geographic disjunction between these faunas (Fig. 1), such a pattern would be biogeographically compelling, if verified. Regional non-monophyly has been well-established in other north-temperate, habitat-specialized arthropods (e.g., Brachycybe millipedes – Brewer et al. 2012; Sabacon harvesters – Schönhofer et al. 2013; travunioid harvesters – Derkarabetian et al. 2018; leptonetid spiders – Ledford et al. 2021), so this biogeographic pattern is certainly possible.

As commonly found in SRE taxa, prior intraspecific genetic research in Hypochilus has revealed ubiquitous and extensive genetic structuring. In Appalachia, extreme mitochondrial genetic divergence occurs within and among five described species over small geographic distances (Hedin 2001; Hedin and Wood 2002; Keith and Hedin 2012). Within H. pococki Platnick, 1987, recovered as paraphyletic on mitochondrial gene trees, highly divergent, geographically cohesive “microclades” have been discovered (Keith and Hedin 2012). In California, mitochondrial CO1 sequences for H. petrunkevitchi Gertsch, 1958 from the Merced versus Kaweah River basins reveal extreme intraspecific genetic divergences (> 15% divergent; Hedin 2001). Past mitochondrial studies have attributed extensive genetic structuring to limited female-biased gene flow (Hedin and Wood 2002; Keith and Hedin 2012), but whether such genetic
patterns extend to the nuclear genome and are less pronounced because of male-based gene flow remains unknown.

In this research we used phylogenomic data to resolve *Hypochilus* species relationships within and among montane regions. We also explored putative cryptic diversification within Appalachian *H. pococki* and Californian *H. petrunkevitchi*. In Appalachia, previous mitochondrial-based species delimitation using a Generalized Mixed Yule Coalescent (GMYC) model (Fujisawa and Barraclough 2013) appears to severely over-split species (Keith and Hedin 2012), likely due to strong genetic structuring. We asked whether similar results applied to nuclear single nucleotide polymorphism (SNP) datasets derived from ultraconserved elements (UCEs), using additional MSC methods (Bayes Factor Delimitation and TR2 rooted triplet analysis). We combine these molecular data and analyses with SEM images of the male pedipalp for large samples of *H. pococki* and *H. petrunkevitchi*, and gather UCE CO1 mitochondrial “by-catch” data for *H. petrunkevitchi*. Based on evidence derived from a combination of nuclear phylogenomic and mitochondrial data, geography, and adult male and female morphology, we describe a new SRE species from the southern Sierra Nevada of California.
**Materials and methods**

**Molecular sampling**

Specimens representing the genus *Ectatosticta* Simon, 1892 from China, the sister genus to *Hypochilus* (and the only other hypochilid genus described), were used to root all phylogenies (UCE data from Ramírez et al. 2021). The monophyly of the family, and two constituent genera, is strongly supported by both morphology (Forster et al. 1987; Alberti and Coyle 1991; Catley 1994; Li et al. 2021) and phylogenomics (Fernández et al. 2018; Ramírez et al. 2021). Phylogenomic data were gathered for forty-three *Hypochilus* specimens, representing all ten described species (see Suppl. material 1; Fig. 1). A priori species identifications were based on male morphology in combination with geography (Forster et al. 1987; Catley 1994), and we included many samples from prior genetic research (Keith and Hedin 2012). Multiple specimens per species were sampled, chosen to maximize the breadth of geographic coverage within species (Fig. 1). Our southern Rocky Mountain samples included collections from near the respective type localities (Gertsch 1964; Catley 1994) of both *H. bonneti* Gertsch, 1964 and *H. jemez* Catley, 1994, plus two geographically intermediate populations from southern Colorado of uncertain species identity. For species delimitation focal taxa (*H. pococki* and *H. petrunkevitchi*), larger sample sizes were used to obtain more fine-scale genetic data on potentially cryptic species (Fig. 1). For *H. pococki*, sampling included representatives of the five mitochondrial haplogroups identified in Keith and Hedin (2012). For *H. petrunkevitchi*, specimens were sampled from multiple drainage basins, including the Merced River (*YOSE*), San Joaquin River (*SAN*), Kings River (*KING*), Kaweah River (*KAW*), Tule River (*TULE*), and Cedar Creek (*CEDAR*) drainages.

**UCE data collection**

For almost all specimens (with tissues stored at -80 °C), DNA extraction was performed using a Qiagen DNEasy kit from leg tissues. Sequence capture libraries were prepared using an ultraconserved elements capture protocol for arachnids (Starrett et al. 2017; Hedin et al. 2019), with arachnid probes designed by Faircloth (2017). Sequencing was done at the Brigham Young University DNA Sequencing Center on an Illumina HiSeq 2500 150 cycle paired-end sequencing platform. Published data for two *H. pococki* specimens (H595 and H232, from Starrett et al. 2017), one *H. kastoni* Platnick, 1987 (G2519, Hedin et al. 2019), and *Ectatosticta* (Ramírez et al. 2021) were used from prior studies. Raw reads were filtered using the illumiprocessor wrapper (Faircloth 2013) within PHYLUCE v1.6 (Faircloth 2016), after which cleaned reads were assembled using Trinity v2.0.6 (Grabherr et al. 2011) and Velvet v1.0.19 (Zerbino and Birney 2008) on the HPC Cluster at UC Riverside. These assemblies were combined and resulting contigs were matched to probes with minimum identity and minimum coverage values of 80 (--min-identity 80 --min-coverage 80). UCE loci were aligned and trimmed within PHYLUCE using MAFFT (Katoh and Standley...
A 50 percent occupancy matrix (623 loci) was generated from the pipeline above (here called the “unfiltered” matrix). A second matrix was further filtered to remove duplicate and potentially non-homologous sequences. Previous work has shown that arthropod UCEs are mostly located within exonic regions (Bosser and Danforth 2018), and the arachnid probe set is no exception (Hedin et al. 2019). In fact, the arachnid probe set can target separate exons from the same protein as separate loci (Hedin et al. 2019), perhaps violating assumptions such as independence and linkage of loci. An annotated list of the arachnid UCEs from Hedin et al. (2019) was used to identify duplicate loci. A total of 73 duplicate loci was found, with the longest locus in each instance retained while the rest were discarded. Using the annotated list, those loci identified by Hedin et al. (2019) as including potential paralogs were also identified and discarded ($n = 2$), leaving a matrix containing a total of 550 loci. Finally, this filtered dataset was trimmed again with more stringent Gblocks settings (--b1 0.5 --b2 0.85 --b3 4 --b4 8), resulting in a “filtered and trimmed” matrix. This last step was conducted to isolate as much of the purely exonic region as possible for each locus.

**Generic-level phylogenomics**

Interspecific relationships were reconstructed using all three UCE data matrices (“unfiltered”, “filtered”, and “filtered and trimmed”), utilizing both concatenation and coalescent-model approaches. Data were partitioned by locus, with optimal models selected using PartitionFinder2 (Lanfear et al. 2012) on the CIPRES portal HPC Cluster. Concatenated maximum likelihood trees were reconstructed using RAxML v8.2.12 (Stamatakis 2014) and IQ-TREE v1.5.0 (Nguyen et al. 2015). Using IQ-TREE 2 we also calculated gene (gCF) and site concordance (sCF) factors. For every node of a reference tree, gCF can be defined as the percentage of “decisive” gene trees containing that node, while sCF can be defined as the percentage of decisive sites (in an alignment) supporting a node (Minh et al. 2018). The latter support metric is particularly useful when individual gene trees are uncertain, perhaps because individual alignments are short. Concordance factor calculations were performed using the topology from IQ-TREE which has an identical topology to the RAxML tree and mostly agrees with the SVDquartets reconstruction (see Results). The latter is a coalescent-model topology reconstructed using SVDquartets v1.0 (Chifman and Kubatko 2014) implemented in PAUP* (Swofford 2003), set for 1M quartets with 500 bootstrap replicates for all runs.

**Nuclear species delimitation**

Nuclear single nucleotide polymorphism (SNP) data were extracted from UCE loci following the methods of Harvey et al. (2016) and a combination of tools and methods from vcf tools (Danecek et al. 2011) and a modified version of the best practices approach for variant isolation with GATK v4.0.0.0 (Van der Auwera et al. 2013).
Separate datasets were created for Californian *H. petrunkevitchi* plus *H. bernardino* specimens, and for samples of *H. pococki*. These matrices started with cleaned read data containing only relevant samples and used a highest coverage reference specimen (H_petrunkevitchi_G2543, H_pococki_H551). Genetic structure and sample clustering were explored using k-means clustering in STRUCTURE v2.3.4 (Pritchard et al. 2000), with unlinked SNPs. Multiple K values (K = 1–10) were run for 100,000 generations with each K value replicated 10 times. Optimal K values were determined following Pritchard et al. (2000) and Evanno et al. (2005), using the online resource CLUMPAK (Kopelman et al. 2015).

Using multiple data sources (phylogenomic results, STRUCTURE results, geography for *H. petrunkevitchi*, and mitochondrial haplogroup membership for *H. pococki*), alternative species models were generated and compared using nuclear SNP datasets in the program SNAPP (Tables 1, 2; Bryant et al. 2012). These analyses also included outgroup data (other *Hypochilus* species), allowing us to test hypotheses of *H. pococki* as a single species (current taxonomy) and *H. petrunkevitchi* lumped with *H. bernardino* (see Tables 1, 2). The averaged marginal likelihoods of duplicate runs were compared for alternative species models using Bayes Factor Delimitation for genomic data (*BFD*) (Leaché et al. 2014); we followed the recommendations of Kass and Raftery (1995) in interpreting Bayes factor values.

A rooted triplet species delimitation approach was also implemented using the Python2 compatible version of the program TR2 (Fujisawa et al. 2016). Here, nuclear

### Table 1. Alternative species model comparison results for *H. petrunkevitchi*, from SNAPP. Alternative models were compared to current taxonomy and ranked with 1 as the most favorable and 5 as the least. Bayes factors were calculated as (BF = 2 × (model 1 – model 2)) where negative values represent support for model 2 (alternative model) and positive values are support for the null model (current taxonomy).

| Model       | Species                  | Partitioning                  | MLE  | MLE 2 | BF   | Rank |
|-------------|--------------------------|-------------------------------|------|-------|------|------|
| Every Tip   | 16                       | Every specimen as a species   | -105 | -105.2| -7056| 1    |
| Basins      | 8                        | *H. kastoni*, *H. bernardino*, CEDAR, TULE, KAW, KING, SAN, YOSE | -1781 | -1781 | -3704| 2    |
| STRUCTURE   | 6                        | *H. kastoni*, *H. bernardino*, TULE+CEDAR, KAW+KING, SAN, YOSE | -1939 | -1939 | -3388| 3    |
| Current Taxonomy | 3                        | *H. kastoni*, *H. petrunkevitchi*, *H. bernardino* | -3633 | -3633 | - | 4    |
| Collapse    | 2                        | *H. kastoni*, *H. petrunkevitchi* + *H. bernardino* | -4223 | -4223 | 1180 | 5    |

### Table 2. SNAPP results for *H. pococki*. Models were compared to current taxonomy and ranked with 1 as the most favorable and 4 as the least.

| Model       | Species                  | Partitioning                  | MLE  | MLE 2 | BF   | Rank |
|-------------|--------------------------|-------------------------------|------|-------|------|------|
| Every Tip   | 16                       | Every specimen as a species   | -62.9| -62.8 | -5030| 1    |
| Mitochondrial | 6                        | *H. thorelli*, WEST+bone+alark, CENT, VA, ELK, NE | -3917 | -3917 | -4462 | 2    |
| STRUCTURE   | 5                        | *H. thorelli*, WEST+bone+alark, CENT, ELK+NE, VA | -4338 | -4340 | -3620| 3    |
| Current Taxonomy | 2                        | *H. thorelli*, *H. pococki* | -6148.9 | -6148 | - | 4    |

 kappa
gene trees are decomposed into partially rooted triplets and congruence is assessed among triplet topologies using a likelihood model testing framework. Input gene trees were constructed in RAxML using rapid bootstrap analysis (-f a) with 200 bootstrap replicates for each gene tree with 550 UCE loci from the “filtered and trimmed” dataset. With the intent to detect patterns of increasing support for increasing species number (i.e., over-splitting), models in which every tip was categorized as a species were included in both SNAPP and TR2 runs.

**CO1 phylogeny, distances, and GMYC**

Mitochondrial Cytochrome c oxidase subunit I (CO1) data for the California taxa were captured from UCE “by-catch” for purposes of phylogenetic and distance analyses, particularly considering the extreme CO1 distances observed in Hedin (2001). A consensus reference sequence was created from specimens of *H. petrunkevitchi* and *H. bernardino* (from Hedin 2001), which was then used as a custom database for a BLASTN search for extracting CO1 sequences from UCE contigs. The BLASTN search and subsequent alignment was performed using Geneious Prime (2020.2). Sequence alignments were partitioned by codon position using MODELFINDER (Kalyaanamoorthy et al. 2017) and a phylogeny was constructed using IQ-TREE with 1000 ultrafast bootstrap replicates. A pairwise mitochondrial distance matrix was generated in PAUP* (Swofford 2003) using a Kimura two-parameter model of nucleotide substitution (Kimura 1980).

A Generalized Mixed Yule Coalescent (GMYC) model was used to algorithmically delimit Californian species using CO1 data, in a manner similar to the approach of Keith and Hedin (2012) for Appalachian *H. pococki*. A GMYC model assumes a difference in intra and inter-specific branching patterns under maximum likelihood and estimates a threshold for the transition from intraspecific (a coalescent process) to interspecific (speciation) branching on an ultrametric tree. Both a single threshold and multiple thresholds can be estimated in different approaches with the model. To this end, an ultrametric CO1 tree was generated using the *chronopl* function from the APE library in R version 4.0.2 (Paradis and Schlkipe 2019), with both single and multiple threshold models performed on the GMYC web server (https://species.h-its.org/gmyc/; Fujisawa and Barraclough 2013).

**Morphological study**

Although *Hypochilus* is a strongly morphologically conserved genus, current species were described and are diagnosed using subtle morphological variation, mostly in male genitalia (Hoffman 1963; Forster et al. 1987; Catley 1994). Using SEM we examined male palpal morphology for California taxa (*H. kastoni, H. bernardino, multiple lineages of H. petrunkevitchi*), and for all primary lineages of *H. pococki*. We lacked adult males for the southern Rocky Mountain populations of uncertain species identity, so could not study them at this time. Male pedipalps preserved in 80% EtOH were
transferred to pure EtOH (200 PF, ≥ 99.5%) for at least 10 min. Following EtOH dehydration, samples were placed in a multilayered sample holder and dried to the critical point using a Tousimis SAMDRI790. Transitional and intermediate fluids used were CO₂(l) and pure EtOH, respectively. Dried palps were mounted on aluminum stubs fitted with sticky carbon conductive spectro-grade tabs. Following mounting, samples were run through an EMS Quorum Q150T sputter coater and covered with 6 nm platinum nanoparticles. SEM micrographs of retrolateral and prolateral views were taken using a vertical stage on an FEI Quanta 450 FEG. Female spermathecal organs were imaged using a Visionary Digital Imaging System, comprising a Canon EOS 5D Mk II DSLR mounted to an Infinity Optics microscope tube. Spermathecal organs were dissected from specimens using fine forceps, immersed for 2–5 minutes in BioQuip specimen clearing fluid (http://www.bioquip.com), then imaged in this fluid on depression slides.

For taxonomic descriptions, morphological measurement details follow Catley (1994: figs 1–4):

| PTW/PTL | maximum width of male pedipalpal tibia in retrolateral view/length of tibia in retrolateral view; |
| CdL     | male palpal conductor length in retrolateral view; |
| AME     | diameter of anterior median eye pupil; |
| PTAŁ    | length of male palpal tarsus in retrolateral view; |
| CTpr    | number of promarginal cheliceral teeth; |
| CTre    | number of retromarginal cheliceral teeth. |

Measurements were taken from alcohol-preserved specimens using an Olympus SZ40 dissecting microscope fitted with an ocular micrometer, and converted to millimeters; raw measurements are provided in Suppl. material 2, and summarized in Table 5.

**Results**

**UCE processing and generic-level phylogenomics**

Original UCE raw reads have been submitted to the SRA (PRJNA760946), with summary statistics presented in the Suppl. material 1. Data matrices and resulting .tre files are deposited at Dryad (https://doi.org/10.5061/dryad.g1jwstqs). The “unfiltered” matrix included 623 UCE loci with an average length of 783 base pairs (bp) and a concatenated length of 734,881 bp (189,387 parsimony informative (PI) sites). The “filtered” matrix included 550 loci with an average length 921 bp and concatenated length of 506,689 bp (145,037 PI sites), while the “filtered and trimmed” matrix contained 550 loci with an average length of 591 bp and concatenated length of 325,452 bp (81,911 PI).
Concatenated ML and SVDquartets analyses of the above three matrices recover nearly identical *Hypochilus* relationships, except for some nodes in the Appalachian and Rocky Mountain clades (Figs 2, 3; https://doi.org/10.5061/dryad.g1jwstqsd). These analyses confirm regional faunas as monophyletic, rejecting CA paraphyly, and recover Appalachia and California clades as sister taxa. Within Appalachia, a sister species relationship between the geographically disjunct *H. thorelli* Marx, 1888 and *H. coylei* Platnick, 1987 is strongly supported across all analyses, consistent with mitochondrial evidence (Hedin 2001; Keith and Hedin 2012), but contrary to morphological evidence which groups *H. coylei* with the geographically adjacent *H. sheari* Platnick, 1987 (Huff and Coyle 1992; Catley 1994). The monophyly of all currently recognized Appalachian species is supported, contradicting mitochondrial results which had previously suggested *H. pococki* paraphyly. Certain parts of the Appalachian topology include low bootstrap values, low gene and site concordance values, and discordant topologies among analyses. Whereas ML analyses place *H. pococki* as sister to other Appalachian species, SVDquartets nests *H. pococki* well within the clade with *H. gertschi* Hoffman,
Figure 3. Concordance Factor values. Phylogeny reconstructed from partitioned RAxML analysis of “filtered and trimmed” UCE matrix, with gCF / sCF values. The lowest gCF and sCF values, for two unresolved nodes which collapse to a four lineage polytomy within the Appalachian clade, are highlighted by red text.
As sister to other Appalachian species (Fig. 2). A California clade is consistently recovered, with *H. kastoni* sister to remaining lineages. *Hypochilus bernardino* is nested prominently within *H. petrunkevitchi*, and for this reason analyses examining species boundaries included samples of all three species (see below).

**Nuclear species delimitation**

Nuclear SNP datasets included 670 unlinked SNPs for the California sample (allowing 20% missing data) and 655 unlinked SNPs for the Appalachian sample (19% missing data). Overall, STRUCTURE analyses reveal strong population structure for both samples, with inferred genetic clusters congruent with phylogenomic clades (Fig. 4). Within California there is little evidence for mixed ancestry, with genetic populations of *H. petrunkevitchi* also appearing to be structured by river basin (Fig. 4). Best K as determined by the Pritchard et al. (2000) method is decisive for K = 5 while the Evanno method is less conclusive, supporting a scenario of K = 2 with almost equal but slightly lower support for K = 4 (https://doi.org/10.5061/dryad.g1jwstqsd). When *H. kastoni* samples were included

![Image of STRUCTURE results](image_url)

**Figure 4.** UCE STRUCTURE results, with optimal and suboptimal K clusters shown in relation to UCE RAxML phylogeny and geography. For *H. pococki*, the distribution of mitochondrial “microclades” follows Keith and Hedin (2012).
in STRUCTURE runs, the Pritchard method recovered the same population structure scheme with the addition of a separate *H. kastoni* “population”, while the Evanno method recovered stronger support for K = 2 in which *H. kastoni* and *H. bernardino* comprised a single genetic population. Because we viewed this latter K = 2 result as spurious (see Janes et al. 2017), we generally preferred optimal K values as inferred by the Pritchard method.

There is more evidence for mixed ancestry in *H. pococki* STRUCTURE analyses, using a best K from both Pritchard (K = 4) and Evanno (K = 3) methods. The Elk and Northeast “microclades” (ELK and NE are lumped as a single genetic population, while one member from the mitochondrial Central (CENT) group clusters with the WEST population (Fig. 4), discordant with previously delineated mitochondrial groups (Keith and Hedin 2012). The geographically isolated BONE population, not sampled in previous mitochondrial studies, clusters with disjunct WEST specimens (Fig. 4).

Alternative species hypothesis models were generated and compared using SNAPP and TR2 for both *H. petrunkevitchi* (five models) and *H. pococki* (four models). Pritchard-based best K schemes were used for the STRUCTURE derived species models. The most-favored SNAPP model for *H. petrunkevitchi* (Table 1) is one where every tip represents a species; the next best supported model is the “Basins”, 8-taxon model. SNAPP results for *H. pococki* were similar and favored every tip as a species as the best model, with a pattern of more speciose models being more favored (Table 2). The rooted triplet TR2 approach also found this pattern of favoring more species-rich models over current taxonomy with the most species-rich model, every tip as a distinct taxon, being the most favored (Tables 3, 4).

**Table 3.** TR2 results for *H. petrunkevitchi*; ranking of the models with 1 being the most favored and 6 being the least favored.

| Model        | Species                        | Partitioning                      | Score  | Rank |
|--------------|--------------------------------|-----------------------------------|--------|------|
| Every Tip    | 16                             | Every specimen as a species       | 171.62 | 1    |
| Basins       | 8                              | *H. kastoni*, *H. bernardino*, TULE, CEDAR, KAW, KING, SAN, YOSE | 213.95 | 2    |
| STRUCTURE    | 6                              | *H. kastoni*, *H. bernardino*, TULE+CEDAR, KAW+KING, YOSE, SAN   | 348.23 | 3    |
| Current Taxonomy | 3                              | *H. kastoni*, *H. petrunkevitchi*, *H. bernardino* | 9334.94 | 4    |
| Collapse     | 2                              | *H. kastoni*, *H. petrunkevitchi* + *H. bernardino*  | 25926.79 | 5    |
| One species  | 1                              | *H. kastoni* + *H. petrunkevitchi* + *H. bernardino* | 30938.48 | 6    |

**Table 4.** TR2 results for *H. pococki*; ranking of the models with 1 being the most favored and 5 being the least favored.

| Model                 | Species                                    | Partitioning                           | Score  | Rank |
|-----------------------|--------------------------------------------|----------------------------------------|--------|------|
| Every Tip             | 16                                         | Every specimen as a species            | 344.10 | 1    |
| Mitochondrial         | 7 (*includes BONE* H. thorelli, WEST, CENT, ELK, NE, VA, BONE) | 366.66 | 2    |
| STRUCTURE             | 5                                          | H. thorelli, WEST+Bone+Alark, CENT, ELK+NE, VA | 819.85 | 3    |
| Current               | 2                                          | H. thorelli, *H. pococki*              | 9930.73 | 4    |
| Collapse              | 1                                          | H. thorelli + *H. pococki*             | 17961.09 | 5    |
CO1 phylogeny, distances, and GMYC

The CO1 by-catch phylogeny, using *H. kastoni* and *H. bernardino* as possible outgroups, shows strong support (BP = 100) for a clade including *H. bernardino* and *H. petrunkevitchi* together (Fig. 5). Within this clade, recovered mitochondrial lineages are consistent with UCE optimal K = 5 STRUCTURE lineages, but the interrelationships of these mitochondrial lineages are not resolved. Considering this phylogenetic uncertainty (i.e., collapsing poorly-resolved nodes), the mitochondrial results are not strictly inconsistent with nuclear results. Pairwise mitochondrial distance values are extremely high among primary lineages (Fig. 5 inset), ranging from 12%–15%. Divergence values within lineages are lower, except for the combined SAN + KING + KAW lineage (> 12% divergence); this obviously reflects significant mitochondrial divergence across drainage basins within this more broadly-distributed lineage. Similarly, there is evidence for structuring across drainage basins within the combined TULE + CEDAR lineage (Fig. 5 inset). As shown previously for Appalachian *H. pococki* (Keith and Hedin 2012), implementation of a GMYC model using CO1 data appears to collapsed as the same species in multi-threshold GMYC analyses; all other branches supported as separate species (e.g., n = 13 for multi-threshold model). **Inset** – K2P distances within and among primary mitochondrial lineages.

**Figure 5.** Maximum likelihood mitochondrial tree. Black circles designate clusters of sequences collapsed as the same species in multi-threshold GMYC analyses; all other branches supported as separate species (e.g., n = 13 for multi-threshold model). **Inset** – K2P distances within and among primary mitochondrial lineages.
severely over-split species. Specimens from the same geographic location are collapsed as the same species, but all unique geographic locations are delimited as distinct species (multi = 13, single = 14; Fig. 5 inset).

**Morphology**

Morphological data for *H. pococki* and California taxa are presented below in the Discussion and Taxonomy sections, respectively.

**Discussion**

**Broad-scale phylogenomics and biogeography**

Our results confirm the Catley (1994) hypothesis of a sister relationship between highly disjunct California and Appalachian faunas, sister to a more geographically central Rocky Mountain clade. Support was unequivocal for this topology, recovered in all analyses from all UCE matrices (Figs 2, 3; https://doi.org/10.5061/dryad.g1jwstqsd). These regional relationships in *Hypochilus* are contrary to patterns seen in the salamander genus *Aneides*, a taxon which also includes montane-associated species from California, the southern Rocky Mountains, and Appalachia. Molecular phylogenetic research in *Aneides* has recovered a sister relationship between California and southern Rocky Mountain species, sister to Appalachian taxa (Vieites et al. 2007). Inter-regional divergences in *Aneides* are estimated to have occurred roughly in the timeframe spanning the Eocene to the Oligocene, perhaps coincident with periods of global warming allowing for intercontinental dispersal events (Vieites et al. 2007).

We hypothesize that *Hypochilus* has a more ancient history, and that this timing difference might also explain the unique phylogenetic topology seen in *Hypochilus*. Divergence time estimates for *Hypochilus* are hindered by a lack of direct fossil evidence, with current age estimates for the genus derived from broader examinations of diversification dates for spiders. For example, using published transcriptome data, Magalhaes et al. (2020) estimated divergence between *H. gertschi* and *H. pococki* (both Appalachian taxa) during the early Paleogene, with a very large confidence interval. Given approximately polytomous relationships within Appalachia (see below), this point estimate would correspond to a crown group age for the Appalachian radiation, and thus implies older divergences at the base of *Hypochilus*, perhaps during the Cretaceous. We note here that many other non-entelegyne araneomorph spider lineages are at least this ancient (both within extant families and sometimes within extant genera), as estimated from molecular clock analyses (e.g., sicariids – Magalhaes et al. 2019; leptonetids – Ledford et al. 2021), but also known directly from Upper Cretaceous Burmese amber fossils (e.g., psilodercids – Magalhaes et al. 2021). Also, the combination of Cretaceous fossil evidence in the context of living spider families suggests that non-entelegyne araneomorph lineages (akin to *Hypochilus* and
Hypochilidae) dominated spider diversity at this time (Wunderlich 2008; Magalhaes et al. 2020).

Hypothesized Cretaceous-age divergences for *Hypochilus* are complicated by the presence of the Western Interior Seaway of North America, a major transcontinental marine barrier in place from ~105–65 mya (Blakey and Ranney 2017). Although an east / west vicariance hypothesis associated with the Western Interior Seaway seems attractive, the Rocky Mountain orogeny took place after the withdrawal of the Western Interior Seaway (Blakey and Ranney 2017). We speculate that *Hypochilus* is either much older than imagined (ages exceeding 105 mya), or that diversification (and dispersal) was spurred soon after the withdrawal of the Western Interior Seaway. Lacking the discovery of relevant fossils, future work could aim to more precisely estimate rates of nuclear gene molecular evolution in *Hypochilus*, in order to better understand the origin and timing of *Hypochilus* diversification events. Also, inclusion of a transcriptome representing the Rocky Mountain clade could be incorporated into the well-calibrated Magalhaes et al. (2020) dataset, allowing for a crown group age estimate for the entire genus.

**Appalachian diversification and potential cryptic species**

Within Appalachia, nuclear UCE data strongly support a monophyletic *H. pococki*, contra mitochondrial paraphyly as in Keith and Hedin (2012). Although all currently described species in Appalachia are recovered as monophyletic, and the geographically disjunct *H. thorelli* and *H. coylei* are strongly supported as sister species (see also Hedin 2001; Keith and Hedin 2012), our nuclear datasets otherwise do not resolve species relationships, with an overall topology consistent with a four-lineage polytomy. Gene and site concordance factor values at two key unresolved interspecific nodes take lower values than seen anywhere else in *Hypochilus*, including all nodes within species (Fig. 3). Gene CF values are 9–13 for these nodes, meaning that only ~10% of the UCE alignments support these nodes. Site CF values that hover around minimum values (30%) illustrate that the data are essentially equivocal regarding three possible resolutions of a quartet for both of these unresolved nodes (Lanfear 2018). This incongruence and lack of resolution possibly points to a non-adaptive radiation where lineages became rapidly isolated from one another due to environmental factors, but the nature of incongruence requires more study.

Nuclear STRUCTURE results confirm distinct genetic groups within *H. pococki* (Fig. 4); however, these genetic groups do not correspond exactly to the previously described mitochondrial “microclades”. In particular, the Alarka Mountain specimen from the mitochondrial Central (CENT) clade of Keith and Hedin (2012) instead groups with the nuclear WEST genetic cluster (Fig. 4). This is important because the geographic boundaries of previously defined mitochondrial clades were hypothesized to coincide with riverine barriers (e.g., the Little Tennessee River separating the CENT versus WEST mitochondrial clades, etc.; see Keith and Hedin 2012: fig. 2). In fact, most phylogeographic studies in the southern Appalachians have primarily relied upon mitochondrial evidence to define geographic groupings (e.g., references in Keith and
Hedin 2012). Future studies that include dense geographic sampling of nuclear lineages will be important here, with *H. pococki* representing a prime candidate. More generally, if gene flow across cryptic lineages is promoting mitonuclear discordance, this system might provide interesting insight into how cryptic lineages interact at areas of contact. Also, areas of parapatric contact can be used to understand the degree to which gene flow is restricted across cryptic lineage boundaries, providing strong and direct tests of species status (see Singhal et al. 2018).

Although SNAPP and TR2 show higher support for increasing species numbers within *H. pococki* (i.e., a many species hypothesis), nuclear STRUCTURE results and consideration of male pedipalp morphology suggest more conservative species numbers. In their diagnosis, Forster et al. (1987) stated that *H. pococki* males “can be recognized by the flaplike tip of the palpal conductor”. We thus focused particular attention on this structure when searching for morphological differences that might distinguish primary genomic lineages (e.g., VA, ELK + NE, CENT, WEST), and included representatives of all such lineages in our SEM surveys. We did not examine female variation (e.g., in spermathecal morphology), but this is another character system to search for morphological differentiation. We observed minimal differences in male palpal morphology across *H. pococki* populations and genomic lineages (Figs 6–8). One possible difference is the shorter secondary coil of the conductor tip observed in ELK specimens (Fig. 6), but we note that ELK itself is well nested within the primary K = 4 lineages (Fig. 4). The discord between nuclear genomic data (and analytical results) which suggest many species, versus morphology which suggests few to one species, is a conspicuous example of the cryptic species challenge, and also focuses attention on patterns of morphological stasis. Despite high genomic divergences and ample evolutionary time, morphological change in *Hypochilus* remains conservative. We might expect conserved *Hypochilus* somatic morphology because of selective constraints on both niche evolution and morphological differentiation, under a model of phylogetic niche conservatism (Keith and Hedin 2012; Fišer et al. 2018). The fact that we also observe similar conservatism in genitalia, where at least genetic drift in isolated populations is expected, is compelling.

**A new Hypochilus species from montane California**

Both nuclear and mitochondrial genetic structuring is very prominent in the California region, and our analyses show that this structure generally follows a pattern of relatedness by drainage basin (Figs 4, 5, 9). The observed divergence within *H. petrunkevitchi* was not surprising as it has previously been noted as having high levels of intraspecific mitochondrial variation (Hedin 2001). As also found for Appalachian samples, both SNAPP and TR2 show a trend of increasing support for models with increasing numbers of species (Tables 1, 3). GMYC analysis of CO1 data similarly delimits all unique geographic locations as distinct species (Fig. 5 inset). We contend that not all local populations can represent unique species, and instead view this as another example of algorithmic over-splitting.
Figure 6. *H. pococki* male palp (conductor) comparison. For each specimen, left panel = prolateral view, right panel = retrolateral view. NE lineage A, B Green Mtn (MCH 01_162) C, D Boone Fork (MCH 01_159); ELK lineage E, F Elk River (MCH 01_155) G, H Linville Gorge (MCH 01_165). Shorter secondary coil for ELK specimens highlighted by arrows; VA lineage I, J Cliff Mtn (MCH 04_028) K, L Guest River (MCH 04_027); CENT lineage M, N Hickory (MCH 01_144) O, P Wagon Road Gap (MCH 01_181); WEST lineage Q, R Alarka (MCH 02_168) S, T Starr Mtn (MCH 02_156) U, V Backbone Rock (MCH 04_025) W, X Chunky Gal Mtn (MCH 02_142). Detailed specimen information provided in Suppl. material 2.
Based on bootstrap values, nuclear data strongly support the hypothesis that \textit{H. bernardino} is phylogenetically nested within \textit{H. petrunkevitchi} (bootstrap = 100 for all matrices across all analyses; Figs 2, 3; https://doi.org/10.5061/dryad.gljwstqsd). Acknowledging that bootstrap values can provide an inflated view of support (Lanfear 2018; Minh et al. 2018), we considered several other concordance factor values for this node (gCF = 34.8, sCF = 48.3, Fig. 3). From a gene (individualUCE locus) perspective, of 155 alignments that could have included the branch of interest (gN = 155), 34.84% or ~ 50 alignments (= gCF) showed the Fig. 3 topology, with very few alignments supporting an alternative topology at high frequency (gDF1 = 3.23, gDF2 = 8.39). Similarly, from a site perspective, of ~ 200 decisive sites for the quartet of interest (sN = 199.42), approximately half support the Fig. 3 topology (sCF = 48.28), with fewer supporting alternative resolutions (sDF1 = 18.33, sDF2 = 29.4). Overall, we view these values (see Lanfear 2018), in concert with bootstrap values, as strongly supporting the paraphyly of \textit{H. petrunkevitchi} with respect to \textit{H. bernardino}.

Based on this phylogenomic pattern we see two obvious taxonomic alternatives. The first is to sink \textit{H. bernardino} into a broadly distributed, highly genetically-structured \textit{H. petrunkevitchi}. The second is to elevate and formally describe the distinct genetic lineage that is sister to \textit{H. bernardino}. We prefer and argue for the latter approach, for
reasons concisely summarized as follows: 1) *H. bernardino* is already described based on a diagnostic morphology (Catley 1994, and revised diagnosis below), 2) *H. bernardino* is geographically-localized, known only from a single mountain range in southern California, highly disjunct from more northern populations (Fig. 9), and 3) *H. bernardino* is supported both as a distinct nuclear and mitochondrial genetic lineage (Figs 2–5). By these multiple measures of genetic, morphological, and geographic distinctiveness, we view *H. bernardino* as an evolutionary lineage on a unique and independent trajectory. Catley (1994) described *H. bernardino* from the southern section of the San Bernardino mountains of southern California. The few known populations are separated from the southern Sierra Nevada by hundreds of kilometers of mostly inappropriate habitat (lower elevations, fewer granite outcrops), where these spiders (or their distinctive webs) have never been found (Fig. 9). This lack of records includes not only our own extensive field work in the intervening region, but also that of the many arachnologists that have conducted research in California, as well as modern-day tools such as iNaturalist. This sort of geographic disjunction has been found in some other taxa, for example, the iconic *Ensatina* salamander complex, where the geographic disjunction is known as “Bob’s Gaps” (Jackman and Wake 1994). In this instance, separated populations have been described as separate subspecies (*E. eschscholtzii klauberi* in the Tehachapi mountains distinct from *E. eschscholtzii croceater* in the San Bernardino...
Figure 9. Southern Sierra Nevada topography map with genetic and morphological sample locations (see Suppl. material 1 and Suppl. material 2). Geographic gaps and other notable geographic features mentioned in the text are highlighted.

mountains; Jackman and Wake 1994), but *Ensatina* taxonomy is generally regarded as being fairly conservative.

Accepting *H. bernardino* as an independently evolving lineage, the current taxonomy must be updated to reflect unique lineages discovered within *H. petrunkevitchi*. 
Conservatively, we retain the northern lineages that include the type locality (male holotype of *H. petrunkevitchi* from Cedar Grove, Fresno County = KINGS lineage) as *H. petrunkevitchi*. This lineage is distributed across the Merced, San Joaquin, Kings and Kaweah River basins (Fig. 9). Again, accepting *H. bernardino* as a unique species, our data indicate that populations from the Tule River and Cedar Creek drainage basins need to be recognized as a new species, which we formally describe below. Specimens from these drainage basins represent new locality records and the southern-most known observations of *Hypochilus* spiders in the Sierra Nevada (Fig. 9). More generally, this part of the southern Sierra Nevada is a well-known area of active speciation, with many short-range endemic arthropods and vertebrates (Bond 2012; Jockusch et al. 2012; Papenfuss and Parham 2013; Satler et al. 2013; Leavitt et al. 2015; Emata and Hedin 2016; Starrett et al. 2018; Bennett et al. 2021; Weng et al. 2021). In this regard, discovering a new species in the southern Sierra Nevada is not surprising.

**Cryptic species concept**

We define species as evolutionary lineages on a “unique and independent trajectory”. Following Davis et al. (2021), we consider species to be cryptic “if they depend on additional sources of data to formulate the delineation hypothesis prior to establishing diagnostic morphological characters”. This definition applies well to the new species description below, which is motivated by a combination of genetic divergence and uniqueness, phylogenetic pattern (paraphyly w.r.t. *H. bernardino*) and geographic allopatry, which has prompted us to take a closer look at morphology. Additionally, the definition allows for an initial hypothesis of morphological crypsis that does not preclude future downstream studies that in fact find morphological (or other) differences, making the species no longer technically “cryptic”. In many instances, species are morphologically cryptic because of their youth, where morphological divergence has not yet caught up with molecular divergence. But as noted above, the measured mitochondrial differences in *Hypochilus* are among the highest known in spiders (a clade with ~ 50,000 described species), and we hypothesize relatively ancient divergences in the genus. These data and arguments are inconsistent with recent evolutionary divergence, and we instead favor a model of long-term phylogenetic niche conservatism constraining morphological evolution (Fišer et al. 2018), as argued above.

Overall, we view our new cryptic species hypotheses as conservative, consistent with perspectives that genomic data should be interpreted conservatively when describing new species (Coates et al. 2018). In *Hypochilus*, this is particularly true since genomic diagnosability is ubiquitous, and extends to the level of localized populations, as reflected in nuclear and mitochondrial algorithmic delimitations. Our hypotheses below do not recognize all genetically divergent lineages as species and allow for varying degrees of population divergence within described species (e.g., *H. petrunkevitchi*, new species below). In particular, there is evidence that the Yosemite Valley (Merced River) population is on a unique evolutionary trajectory due to its disjunct distribution and because samples from the isolated valley floor routinely fall out as a divergent
group in both nuclear and mitochondrial analyses (Figs 2–5). The single sampled population from the San Joaquin drainage is similarly genetically divergent and geographically isolated. This disjunction might be natural, as spiders have never been collected in the apparently suitable granite outcrop habitats between San Joaquin locations and the Yosemite Valley, despite concerted collecting efforts (Fig. 9). In the context of an integrative taxonomic framework, we weigh conservation considerations, extreme geographic allopatry, and well-supported paraphyly as particularly important. Genomic diagnosability is important but not decisive, with morphological diagnosability as least important, again reflecting phylogenetic niche conservatism. Regarding conservation value, we argue that sinking *H. bernardino* into a broadly distributed, highly genetically-structured *H. petrunkevitchi* would immediately decrease the conservation value and importance of the former.

**Taxonomy**

*Hypochilus* Marx, 1888

**Diagnosis.** following Forster et al. (1987).

*Hypochilus bernardino* Catley 1994

Figs 9–13

*Hypochilus petrunkevitchi* Gertsch 1958: Forster et al. 1987: 22 (San Bernardino county records).

*Hypochilus bernardino* Catley 1994: 10, figs 7, 11, 25, 33, 36–39.

**Material examined.** F from Forsee Creek (SDSU_G2893), Ms from East Fork Mountain Home Creek (SDSU_G2929–2932), see Suppl. material 2.

**Diagnosis.** Following from the original diagnosis of Catley (1994), we paid closest attention to the length of the PTaL (should be shorter in *H. bernardino*), and the PTW/PTL (should be shorter and more thickened proximally in *H. bernardino*). We found that PTaL overlaps with northern populations (Table 5), and is therefore not diagnostic. The PTW/PTL ratio is generally smaller in *H. bernardino*, but there is some overlap with northern populations, again calling into question the diagnostic value of this character. We did find that the male CdL is consistently shorter in *H. bernardino* (Table 5), and hypothesize this as a new morphological character diagnostic for the species. Again, consistent with a hypothesis of phylogenetic niche conservatism imparting morphological stasis, the species is only weakly morphologically diagnoseable. The disjunct geographic distribution (Fig. 9) and hundreds of diagnostic nucleotide changes (alignments at https://doi.org/10.5061/dryad.g1jwstqsd) can also be used to recognize this species.
Table 5. Morphological measurements. PTW/PTL (maximum width of male pedipalpal tibia in retrolateral view/length of tibia in retrolateral view), CdL (male palpal conductor length in retrolateral view), AME (diameter of anterior median eye pupil), PTaL (length of male palpal tarsus in retrolateral view), CTpr (number of promarginal cheliceral teeth), CTre (number of retromarginal cheliceral teeth). Raw measurements provided in Suppl. material 2.

|                  | PTW/PTL | CdL       | AME    | PTaL  | CTpr | CTre |
|------------------|---------|-----------|--------|-------|------|------|
| *H. bernardino*  | 0.253–0.267 | 0.475–0.50 | 0.1–0.125 | 0.82–1.0 | 4–5  | 2–3  |
| *H. petrunkevitchi* YOSE lineage | 0.259–0.292 | 0.625 | 0.10–0.125 | 0.925–1.0 | 5 | 2 |
| *H. petrunkevitchi* KING lineage | 0.278 | 0.60 | 0.10 | 0.875 | 5 | 2 |
| *H. petrunkevitchi* KAW lineage | 0.274–0.307 | 0.575–0.675 | 0.10–0.125 | 0.925–1.15 | 4–5  | 2 |
| *H. xomote* sp. nov. | 0.256–0.338 | 0.525–0.575 | 0.10–0.125 | 0.775–1.075 | 4–5  | 1–2 |

**Genetic data.** SRA Accession numbers: SAMN21239435–SAMN21239437.

**New records.** California, San Bernardino County, San Bernardino Mountains, Camp Creek east of Forest Falls, 34.0760, -116.8876, coll. M. Hedin, 10 July 1993 (SDSU_H0025–0027, 3I). San Bernardino County, San Bernardino Mountains, Hwy 38, tributary into East Fork Mountain Home Creek, in culvert and tunnel under highway, 34.1198, -116.9768, coll. E. Ciaccio, 4 August 2018 (SDSU_G2897–2899, 3I). San Bernardino County, San Bernardino Mountains, Hwy 38, tributary into East Fork Mountain Home Creek, in culvert and tunnel under highway, 34.1198, -116.9768, coll. E. Ciaccio, 27 Sept 2018 (SDSU_G2929–2932, 4M). San Bernardino County, San Bernardino Mountains, Hwy 38, Forsee Creek, along stream and tunnel under highway, 34.1574, -116.9315, coll. E. Ciaccio, 4 August 2018 (SDSU_G2893–2896, F, 3I). See also Suppl. material 2 for locality (including elevation) and natural history information for specimens examined.

**Remarks.** Catley (1994) hypothesized the following diagnostic features, based on comparisons to near-type locality *H. petrunkevitchi*:

“The species most closely resembles its sister species *Hypochilus petrunkevitchi* in general coloration, eye dimensions, and male pedipalpal morphology. Males can be recognized by the apex of the conductor which is more loosely whorled (fig. 24) than in *H. petrunkevitchi*, the shorter pedipalpal tarsus, a greatly reduced distal palpal (conductor) apophysis (fig. 25), and a median palpal apophysis that is significantly smaller than *H. petrunkevitchi*, with no notch (fig. 7). The short palpal tibia is strongly incrassate proximally. Females ... are particularly difficult to separate from *H. petrunkevitchi* females, the former possessing similar but smaller convoluted spermathecal ducts (fig. 11).”

Our comparisons of near-type *H. bernardino* to larger samples (Suppl. material 2) of more northern populations in California (not including the distinctive *H. kastoni*) suggests the following character trends. Regarding the shape of the apex of the conductor, we find no consistent difference in tightness of the whorls (Fig. 10), a feature that we also found difficult to characterize. The small distal conductor apophysis is likewise inconsistently absent or barely present across populations (Fig. 10). We also could not discern a consistent difference in the shape of the median palpal apophysis (Figs 11, 12), with our SEM imaged *H. bernardino* specimens appearing much like the
original drawings of *H. petrunkevitchi* (Catley 1994: fig. 8). Differences in the degree of sclerotization at the base of this apophysis also makes the narrowness somewhat subjective to score, at least in some specimens. Finally, we found female spermathecal morphology to be highly conserved (Fig. 13); it is possible that the median ducts are more convoluted in *H. bernardino* than in northern populations, but this difference is subtle given our sampling.

**Distribution and habitat.** Known only from two primary forks of a single drainage basin (headwaters of Santa Ana River, and Mill Creek, a large tributary of the Santa Ana), south side of the San Bernardino Mountains of southern California (Fig. 9). The Forsee Creek population, near the headwaters of the Santa Ana River, represents a new record for this species. We suspect that additional populations likely exist in the narrow
canyons that lead into the Santa Ana River, for example, Bear Creek, Warm Springs Canyon, etc. In our recent collections we have found spiders in webs on large, sheltered granite boulders near streams, and in stream culverts beneath a primary highway.

**Conservation.** We view *H. bernardino* as a short-range endemic taxon with a precarious future, deserving of special conservation attention and monitoring efforts. Over 25 years ago, Catley (1994) discussed how populations may have been negatively affected by drought. More recently, large fires have burned the forests of the Mountain Home Creek drainage (e.g., 2018 Valley fire). The loss of forest canopy cover is expected to result in fundamental changes in microhabitat conditions, and in concert with increasing global temperatures, calls for continued close monitoring of *H. bernardino* populations.

*Hypochilus petrunkevitchi* Gertsch, 1958

Figs 9–13

*Hypochilus petrunkevitchi* Gertsch 1958: 11, figs. 5, 7, 15, 17, 21; Lehtinen 1967: 431, fig. 14; Forster et al. 1987: 21, figs 68–72; Catley 1994: 7, figs. 8, 13, 24.

**Material examined.** Fs from Ladybug Trail (SDSU_G2275), Mineral King Road (SDSU_G2485), Providence Creek (SDSU_G2508), Mill Creek (SDSU_G2543), Huntington Lake Road (SDSU_G2514, SDSU_G2557), Yosemite Falls (SDSU_G2564); Ms from Atwell-Hockett Trail (SDSU_G2260), Big Fern Springs (SDSU_G2262), Ladybug Trail (SDSU_G2274), Mehrten Creek (SDSU_G2285), Mineral King Road (SDSU_TAC000192), South Fork Kaweah River (SDSU_G2279), Mill Flat (SDSU_G2254), and Yosemite Falls (SDSU_G2568, SDSU_G2569); see Suppl. material 2.

**Diagnosis.** We found that the male palpal conductor (CdL) is consistently longer in *H. petrunkevitchi* than in both *H. bernardino* and the new species below, although barely for the latter (Table 5), and larger sample sizes might negate this difference. It is clear that all southern Sierran populations retain a very similar morphology, with minor morphological divergence associated with evolution in the southern Transverse ranges (*H. bernardino*).

**Genetic data.** SRA Accession numbers: SAMN21239424–SAMN21239431.

**New records.** Merced River drainage (YOSE): California, Mariposa County, Yosemite NP, Big Oak Flat Rd., bridge over Tamarack Creek, 37.7278, -119.7143, coll. E. Ciaccio, M. Hedin, A. Rivera, 29 Sept 2017 (SDSU_G2561–2563, 3I). Mariposa County, Yosemite NP, vic Bridalveil Falls, 37.7167, -119.6519, coll. E. Ciaccio, 3 August 2017 (SDSU_G2515–2518, 4I). Mariposa County, Yosemite NP, near Yosemite Falls, 37.7491, -119.5965, coll. E. Ciaccio, M. Hedin, A. Rivera, 29 Sept 2017 (SDSU_G2564–2566, 2I, 2F, 2M). Mariposa County, Yosemite NP, near Yosemite Falls, 37.7491, -119.5965, coll. M. Hedin, K. Crandall, 27 June 1992 (SDSU_H0015–H0016, 2I). San Joaquin River drainage (SAN): Fresno County, Sierra NF,
Figure 11. California taxa male palp comparison, prolateral views. H. kastoni A Ney Springs (SDSU_TAC000191); H. bernardino B Mtn Home (SDSU_G2931) C Mtn Home (SDSU_G2932); H. xantomote D Tule River (SDSU_G2289) E Alder Creek (SDSU_G2600) F Belknap Springs (SDSU_G2300); H. petrunkevitchi KINGS lineage G Mill Flat (SDSU_G2554); H. petrunkevitchi KAW lineage H Mineral King Road (SDSU_TAC000192); H. petrunkevitchi YOSE lineage I Yosemite (SDSU_G2568). Median apophysis highlighted by arrows. For specimens E and I the bulb has rotated during specimen prep; these two images are retrolateral views, subsequently flipped in Photoshop. Detailed specimen information provided in Suppl. material 2.

Huntington Lake Rd., Balsam Creek turnout, 37.1884, -119.2591, coll. E. Ciaccio, 31 July 2017 (SDSU_G2510–2514, 1I, 4F). Fresno County, Sierra NF, Snowslide Creek on Huntington Lake road, 37.2029, -119.2367, coll. E. Ciaccio, 10 Sept 2017 (SDSU_G2557–2560, 1I, 3F). Kings River drainage (KING): California, Fresno County, Sierra NF, McKinley Grove Rd., Bear Creek turnout, 37.0411, -119.1202, coll. E. Ciaccio, B. Hernandez, S. Torres, J. Waters, 24 July 2017 (SDSU_G2557–2560, 4I, 1M). Fresno County, Sierra NF, McKinley Grove Big Trees Area, 37.0224, -119.1066, coll. E. Ciaccio, 10 Sept 2017 (SDSU_G2555, G2556, 1I, 1F). Fresno County, Bretz Mill, Providence Creek, 37.0427, -119.2371, coll. E. Ciaccio, 31 July
2017 (SDSU_G2505–2509, 2M, 3F). Fresno County, Kings Canyon NP, dam at Sheep Creek on Don Cecil Trail, 36.7840, -118.6784, coll. E. Ciaccio, 9 Sept 2017 (SDSU_G2551, 1I). Fresno County, Kings Canyon NP, Road’s end permit station, Bubs Creek/Zumwalt Meadow trail jct, 36.7918, -118.5871, coll. E. Ciaccio, 9 Sept 2017 (SDSU_G2549–2550, 11, 1F). Fresno County, Mill Flat OHV staging area, Mill Flat Creek, 36.7452, -119.0047, coll. E. Ciaccio, 30 July 2017 (SDSU_G2502–2504, 3I). Fresno County, Sequoia NF, Mill Flat OHV Staging Area, 36.7471, -119.0046, coll. E. Ciaccio, 9 Sept 2017 (SDSU_G2552–2554, M, 2F). Fresno County, Sequoia NF, Ten Mile Rd., at bridge N of Hume Lake, 36.7838, -118.9006, coll. E. Ciaccio, 8 Sept 2017 (SDSU_G2546–2548, 1F). Fresno County, Sequoia NF, Ten Mile Rd., Landslide Creek turnout, 36.7625, -118.8801, coll. E. Ciaccio, 29 July 2017 (SDSU_G2497–2501, 11, 2M, 2F). Fresno County, Sequoia NF, Hwy 245 at Mill Creek, ~1 mi S of Hwy 180 Jct, 36.7145, -118.9879, coll. E. Ciaccio, 30 July 2017 (SDSU_G2542–2545, 2I, 2F). Kaweah River drainage (KAW): Tulare County, Sequoia NP, Hwy 198, Big Fern Springs, 36.5382, -118.7751, coll. M. Hedin, 12 July 1993 (SDSU_H0020–H0022, 3I). Tulare County, Sequoia NP Hwy 198, Big Fern Springs, 36.5382, -118.7751, coll. E. Ciaccio, 17 August 2016 (SDSU_G2261–G2265, 1M, 3F, 1I). Tulare County, Sequoia NF, Forest Rte 14S11, Boulder Creek turnout, 36.7342, -118.7736, coll. E. Ciaccio, 29 July 2017 (SDSU_G2492–G2496, 1M, 4F). Tulare County, Sequoia NP, Hwy 198 near Lodgepole CG, Marble Fork Kaweah River, 36.6037, -118.7392, coll. E. Ciaccio, 29 July 2017 (SDSU_G2487–G2491, 4M, 1F). Tulare County, Sequoia NP, Mineral King Rd., Squirrel Creek pullout, 36.4428, -118.7694, coll. E. Ciaccio, 28 July 2017 (SDSU_G2482–G2486, 3M, 1F, 1I). Tulare County, Sequoia NP, Atwell-Hockett Trail, bridge on trail, 36.4584, -118.6564, coll. E. Ciaccio, 16 August 2016 (SDSU_G2256–G2260, 3M, 1F, 1I). Tulare County, Sequoia NP, bridge over Marble Fork on road to Crystal Cave, 36.5759, -118.7860, coll. E. Ciaccio, 17 August 2016 (SDSU_G2266–G2270, 4F, 1I). Tulare County, Sequoia NP, Ladybug Trail, upstream from bridge at start of trail, 36.35005, -118.76238, coll. E. Ciaccio, B. Hernandez, S. Torres, 3 Sept 2016 (SDSU_G2273–G2278, 2M, 3F, 1I). Tulare County, Sequoia NP, Middle Fork Trail, 36.5416, -118.7074, coll. E. Ciaccio, 18 August 2016 (SDSU_G2271, 1I). Tulare County, Sequoia NP, Middle Fork Trail, Mehrten Creek, 36.5457, -118.6920, coll. E. Ciaccio, B. Hernandez, S. Torres, 4 Sept 2016 (SDSU_G2284–G2288, 2M, 2F, 1I). Tulare County, Sequoia NP, Middle Fork Trail, near Mehrten Creek, 36.5456, -118.7036, coll. E. Ciaccio, 16 August 2016 (SDSU_G2272, 1I). Tulare County, Sequoia NP, Mineral King Road, turnout on the road, 36.45346, -118.6923, coll. E. Ciaccio, 18 August 2016 (SDSU_G2251–2255, 5F). Tulare County, Sequoia NP, Mineral King Road, crossing of Redwood Creek, W of Atwell Mill CG, 36.4533, -118.7036, coll. M. Hedin, 24 August 2009 (SDSU_TAC000192-TAC000193, 1F, 1M). Tulare County, Sequoia NP, South Fork Kaweah River, jct of Cedar Creek and South Fork Kaweah River, 36.3551, -118.7335, coll. E. Ciaccio, B. Hernandez, S. Torres, 4 Sept 2016 (SDSU_G2279–G2283, 1M, 3F, 1I). Tulare County, Sequoia NP, below Atwell Mill CG, along Kaweah River, 36.4584, -118.6561, coll. M. Hedin, 23 August 2009 (SDSU_H0842-H0844, 1F, 1M, 1I). See
also Suppl. material 2 for locality (including elevation) and natural history information for specimens examined.

**Remarks.** Catley (1994) provided no formal diagnosis for *H. petrunkevitchi* (and original diagnoses only compared *H. petrunkevitchi* to the easily distinguishable *H. kastoni*), but offered the following differences from *H. bernardino* in his keys to males and females - length of male PTaL relatively long (0.99–1.10 mm); index of shape of male palpal median apophysis (= vertical height of distal edge of apophysis × length of ventral border of apophysis) large (0.02–0.03) and strongly notched at proximal edge; apex of the conductor in tight whorl with large inwardly directed distal apophysis. Spermathecal bulbs large, diameter of largest not less than 0.11 mm; median ducts of greater length than lateral ducts. We have commented above on the shape of the apex of the male conductor (Fig. 10), the shape of the male palpal median apophysis (Figs 11, 12), and a PTaL which overlaps in length (Table 5). Similarly, we view the relative size of spermathecal bulbs, and relative length of median versus lateral ducts as qualitatively similar (Fig. 13).

**Distribution and habitat.** *H. petrunkevitchi* was previously known from a handful of locations in the west-central Sierra Nevada, and our work has greatly expanded our distributional knowledge for this species (Fig. 9). All previous taxonomic publications involving this species examined only northern specimens (Kaweah River drainage and northwards), which here retain the name *H. petrunkevitchi*.

**Conservation.** Of all the basins in the southern Sierra Nevada, the Kaweah and Kings populations appear to occupy the most contiguous habitat, as reflected in both nuclear phylogenies and STRUCTURE results (Figs 2–5, 9). The Yosemite and San Joaquin populations, which are geographically isolated and particularly genetically divergent, deserve close monitoring. While all Yosemite populations lie within the boundaries of Yosemite National Park, this does not strictly assure future persistence. Populations in Yosemite Valley occur only in deep breakdown “caves”, and in our experience spiders are not abundant. Both known San Joaquin populations occur in habitats that have recently burned as part of the devastating 2020 Creek Fire, again likely changing the nature of the canopy structure (and thus microclimatic conditions) in this area.

**Hypochilus xomote** Hedin & Ciaccio, sp. nov.

http://zoobank.org/7AF45D16-59AC-4E3D-B846-3DD14D4B8BBE

Figs 9–14

**Type material.** **Holotype** male (SDSU_TAC000658) from California, Kern County, upstream of Cedar Creek campground, off Hwy 155, Sequoia National Forest, 35.7508, -118.5807, elevation ~ 1520 meters, coll. M. Hedin, 4 October 2021 (MCH 21_091). Deposited at the University of California Davis Bohart Museum of Entomology. **Paratype** females (SDSU_TAC000659, TAC000660) and paratype male (SDSU_TAC000661) from same collecting event (MCH 21_091). Deposited at the Denver Museum of Nature and Science.
**Etymology.** *xomote*, from the Native American Yowlumni tribal word for south, providing a name for the southern-most known *Hypochilus* populations in the California Sierra Nevada. The X of *xomote* is pronounced as a “breathy, hissy sort-of H” (Vera and Clark 2002). Language translation from the Tule River Yokuts Language Project (Vera and Clark 2002), representing the language of the Yowlumni Yokuts. Members of the larger Yokuts people historically occupied the southern San Joaquin Valley and adjacent Sierran foothills, including the Tule River basin; the Yowlumni occupied a smaller region near the valley outlet of the Kern River (see Fig. 9).

**Diagnosis.** CdL of intermediate length (Table 5), longer than *H. bernardino* but shorter than *H. petrunkevitchi*, although barely so for geographically adjacent KAW populations of *H. petrunkevitchi*.

**Genetic data.** SRA Accession numbers: SAMN21239432–SAMN21239434.

**Description.** Male holotype – Total length 7.5. Cephalothorax 3.0 long, 2.4 wide: clypeus 0.10. Eye diameters: AME 0.125, ALE 0.225, PME 0.175, PLE 0.20.

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**Figure 12.** California taxa male palp comparison, retrolateral views. *H. kastoni A* Ney Springs (SDSU_TAC000191); *H. bernardino B* Mtn Home (SDSU_G2931) C Mtn Home (SDSU_G2932); *H. xomote D* Tule River (SDSU_G2289) E Alder Creek (SDSU_G2600) F Belknap Creek (SDSU_G2300); *H. petrunkevitchi* KINGS lineage G Mill Flat (SDSU_G2554); *H. petrunkevitchi* KAW lineage H Mineral King Road (SDSU_TAC000192); *H. petrunkevitchi* YOSE lineage I Yosemite (SDSU_G2568). For specimens E and I the bulb has rotated during specimen prep; these two images are prolateral views, subsequently flipped in Photoshop. Detailed specimen information provided in Suppl. material 2.
Chelicerae pale yellow to white, dusky markings at base; promarginal cheliceral teeth 5, cheliceral formula 52314, retromarginal cheliceral teeth two; one distal, one proximal, both very small. Endites and labium white to pale yellow; sternum with dusky pigmentation, small unpigmented patches circling sparse weak setae; coxae whitish; trochanters with proximal and distal pigmented patches; all legs yellow tan with broken dark annulations on femora and tibiae; prolateral proximal aspect of femur 1 with ~20 unpigmented weak setae; leg 1 > 20 × length of cephalothorax. Abdomen dorsally pale yellow- white with darker maculations over the entire surface, clothed with sparse hairs, with multiple transverse rows of small weak setae. Palpal tarsus (left) (0.875), palpal tibia short (1.875), thickened proximally (width 0.5), PTW/PTL = 0.267. Conductor length (0.55), conductor tip loosely whorled with very small distal apophysis in retrolateral view. Leg formula 1243; spination (only surfaces bearing spines listed): pedipalpal femur: none; tibia: many dorsal, many prolateral, few to none retrolateral; tarsus: setose with five closely appressed black spines on retrolateral surface of apical spur. Femur I-many prolateral/dorsal; legs II–IV one dorsal proximally. Trichobothrial distribution: all legs with one trichobothria distally on tibia and metatarsus.

**Female paratype (SDSU_TAC000659):** Total length 11.8, cephalothorax 3.8 long, 3.1 wide; clypeus 0.20. Eye diameters: AME 0.175, ALE 0.275, PME 0.225, PLE 0.20. Clypeal area, lateral aspects of head, and foveal area with dusty maculations. Pedipalp pale yellow-white, legs pale yellow-white with femora and tibiae of all legs with broken dark rings and conspicuous dark spots, first leg > 9 × length of cephalothorax. Chelicerae pale yellow, dusky on front proximal surface. Spermathecae with convoluted ducts and relatively large receptacula (e.g., Fig. 13F). Leg formula 1243; spination (only surfaces bearing spines listed): pedipalpal femur: few distal and dorsal; tibia: few dorsal and prolateral, few to none retrolateral; metatarsus: stronger and denser than other pedipalp elements. Femur I-many prolateral/dorsal; legs II–IV one dorsal proximally. Trichobothrial distribution: pedipalpal tibia with a series of dorsal trichobothria; all legs with one trichobothria distally on tibia and metatarsus.

**Other material examined.** Tule River drainage: California, Tulare County, Sequoia NF, Balch Park Road, Jenny Creek, 36.2843, -118.7335, coll. E. Ciaccio, 28 July 2017 (SDSU G2477–2481, 5F). Tulare County, Sequoia NF, Hwy 190 turnout, Belknap Creek, 36.1534, -118.5977, coll. E. Ciaccio, 23 Sept 2016 (SDSU G2296–G2300, 4F, M). Tulare County, Mountain Home State Forest, Hidden Falls campground, 36.2585, -118.6631, coll. E. Ciaccio, 28 July 2017 (SDSU G2472–G2476, 2F, 3I). Tulare County, Sequoia NF, Hwy 190, McIntyre Creek turnout, 36.1509, -118.5831, coll. E. Ciaccio, 27 July 2017 (SDSU G2467–2471, 3F, 2I). Tulare County, Sequoia NF, North Fork Middle Fork Tule River, 36.2082, -118.6488, coll. E. Ciaccio, 24 Sept 2016 (SDSU G2307–G2311, 4F, M). Tulare County, Sequoia NF, Road 208, North Fork Middle Fork Tule River, 36.1879, -118.6775, coll. E. Ciaccio, 23 Sept 2016 (SDSU G2289–2295, 1I, 1M, 5F). Tulare County, Sequoia NF, Forest Route 21S94, Windy Creek turnout, 36.0810, -118.6055, coll. E. Ciaccio, 27 July 2017 (SDSU G2462–2466, 4F, 1I). Cedar Creek drainage: Kern County, Sequoia NF, Alder Creek campground, north
Figure 13. Comparative female spermathecal morphology for California taxa. *H. bernardino* A Foresee Creek (SDSU_G2893); *H. petrunkevitchi* YOSE lineage B Yosemite (SDSU_G2564); *H. xomote* C Jenny Creek (SDSU_G2477) D Windy Creek (SDSU_G2465) E Belknap Creek (SDSU_G2296) F Alder Creek (SDSU_G2601); *H. petrunkevitchi* KAW lineage G Ladybug Trail (SDSU_G2275) H Mineral King Road (SDSU_G2485); *H. petrunkevitchi* KINGS lineage I Mill Creek (SDSU_G2543) J Providence Creek (SDSU_G2508); *H. petrunkevitchi* SAN lineage K Snowslide Creek (SDSU_G2557) L Balsam Creek (SDSU_G2415). Scale bars shown for select specimens. Detailed specimen information provided in Suppl. material 2.
Distribution and habitat. Known only from the upper Tule River and upper Cedar Creek drainages, at the southern end of the California Sierra Nevada mountains (Fig. 9). We hypothesize that higher elevation xeric ridges (Dennison Mountain ridge in particular) separate the distribution of this species from Kaweah River drainage populations of *H. petrunkevitchi*. Populations of *H. xomote* sp. nov. are predicted to be present in the White River drainage that lies between the Tule and Cedar Creek drainages (Fig. 9), although collecting efforts in this drainage have failed thus far. We
have collected these spiders on shaded granite boulders, in mineshafts, and in stream culverts, generally near water along rivers or streams, in conifer or mixed oak/conifer forests (Fig. 14; Suppl. material 2). See also Suppl. material 2 for locality (including elevation) and natural history information for specimens examined.

**Conservation.** Specimens are more abundant and populations appear more secure in the densely forested and higher elevation/higher latitude Tule River drainage. Specimens are less abundant and populations appear more fragmented in the lower elevation and more southerly Cedar Creek drainage. Recent large fires have occurred in both drainages.

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We dedicate this work to Dr. David Wake, a brilliant and generous herpetologist and evolutionary biologist from the Museum of Vertebrate Zoology and UC Berkeley, who passed away in early 2021. Dr. Wake’s work on California biodiversity and the speciation processes of this landscape, and his open-mindedness and appreciation for both vertebrate and invertebrates, have long served to inspire the senior author.

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Supplementary material 1

Appendix I
Authors: Erik Ciaccio, Andrew Debray, Marshal Hedin
Data type: xslx. file
Explanation note: Specimens used in phylogenomic analysis.
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Link: https://doi.org/10.3897/zookeys.1086.77190.suppl1

Supplementary material 2

Appendix II
Authors: Erik Ciaccio, Andrew Debray, Marshal Hedin
Data type: xslx. file
Explanation note: Specimens examined and used for study of morphology.
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