

Genomes of skipper butterflies reveal extensive convergence of wing patterns

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For centuries, biologists have used phenotypes to infer evolution. For decades, a handful of gene markers have given us a glimpse of the genotype to combine with phenotypic traits. Today, we can sequence entire genomes from hundreds of species and gain yet closer scrutiny. To illustrate the power of genomics, we have chosen skipper butterflies (Hesperiidae). The genomes of 250 representative species of skippers reveal rampant inconsistencies between their current classification and a genome-based phylogeny. We use a dated genomic tree to define tribes (six new) and subtribes (six new), to overhaul genera (nine new) and subgenera (three new), and to display convergence in wing patterns that fooled researchers for decades. We find that many skippers with similar appearance are distantly related, and several skippers with distinct morphology are close relatives. These conclusions are strongly supported by different genomic regions and are consistent with some morphological traits. Our work is a forerunner to genomic biology shaping biodiversity research.

genomics | taxonomy | higher classification | mimicry | Lepidoptera

It is the phenotype that is relevant when an animal interacts with its environment. Phenotypic traits have defined the place of an organism within its ecosystem and within a human-made classification system. Due to phenotypes being riddled by adaptive convergence, it is challenging to decipher phylogeny, and thus deduce phylogenetic classification from morphology. The phenotype is encoded by the genotype, which equally bears the footprint of evolution. DNA sequences are prime for evolutionary studies. Sanger-sequencing of select gene markers revolutionized phylogenetic research and refined morphology-based classification. However, frequent homoplasies are a serious obstacle to phylogeny reconstruction from small datasets. Since genomes are composed of millions of base pairs, the study of complete genotypes is expected to resolve many outstanding questions. While large-scale genomic studies are still scarce, they are most enlightening (1).

To exemplify a large-scale genomics project possible within a small laboratory, we use skipper butterflies (Hesperiidae). This family of about 4,000 species worldwide received less attention than other butterflies. Only a handful of studies have focused on skipper phylogeny (2–6). Pioneering work by Warren et al. (2, 3) revealed many surprising phylogenetic relationships compared with the morphological treatment (7–12). However, many questions remained unanswered. In particular, the subfamily Eudaminae was not analyzed in detail.

We obtained and analyzed genomic shotgun reads of 250 species of skippers covering all Eudaminae genera. We constructed a genome-level phylogeny of Hesperiidae and found that, due to convergence in wing patterns, many skippers have been placed in presumed evolutionary groups in which they do not belong. We also discovered a group of close relatives with disparate morphology. Furthermore, sequence data suggest possible genomic determinants of morphological traits.

Results and Discussion

Genomes of 250 Skippers. We selected 250 species of Hesperiidae from all subfamilies and tribes (*SI Appendix*, Table S1). Only 36

specimens were collected after 2012 and preserved in a DNA-friendly manner. Others were pinned in collections listed in *SI Appendix*, Table S1 (also see *Acknowledgments*) and included specimens gathered over a century ago. Nevertheless, sequence reads targeting $10\times$ coverage of the genome allowed us to assemble genomic regions by emphasizing protein-coding genes. We used 12,618 nuclear genes from the two reference genomes of Hesperiidae we have sequenced (13, 14). Aligned nuclear genomic regions covered $8,100,834\pm2,387,641$ positions in each specimen. Mitogenomes covered $10,663\pm674$ positions.

Genomic Tree of Skippers. We constructed genomic trees from concatenated alignments of coding regions (Fig. 1) and from coding regions with introns, and both trees had identical topologies. Furthermore, we constructed gene trees and combined them using the program ASTRAL (15), which produced the tree with the same topology. The trees were dated, i.e., transformed to predict absolute ages for its nodes (*SI Appendix, Methods*). The major phylogenetic groups in these trees agree with those defined by Warren et al. (2, 3) and follow-up publications that used larger sets of species and genes (4–6). Subfamily Coeliadinae (the awls) is sister to all other Hesperiidae (Fig. 1). Australian endemic *Euschemon*, as suggested by several studies (2, 3, 16), is unique and forms a

Significance

With decreasing cost, biodiversity genomics can be done by small laboratories, not large centers. Here, we overview the genomic landscape of an entire family of animals. By sequencing 250 representatives of skipper butterflies, we infer their phylogeny and search for genotypic determinants of phenotypic traits. We find that wing patterns are frequently convergent. This likely mimetic convergence is diversified, resulting in five distinct parallel patterns. Each of the five patterns occurs within at least two genera as well as in more distant relatives diverged more than 20 Mya. This work offers an example of how each of the nearly 6,000 families of animals will be studied in the future.

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Data deposition: The sequences reported in this paper have been deposited in the NCBI Sequence Read Archive (accession no. SRP147939) and the GenBank database (accession nos. MH357724–MH357835). This work was registered with ZooBank (www.zoobank.org) as SA8E82C4-AC8A-487B-8DF4-6BFF4BD48F3D, and registration numbers for 24 nomenclatural acts are given in Tables 1 and 2.

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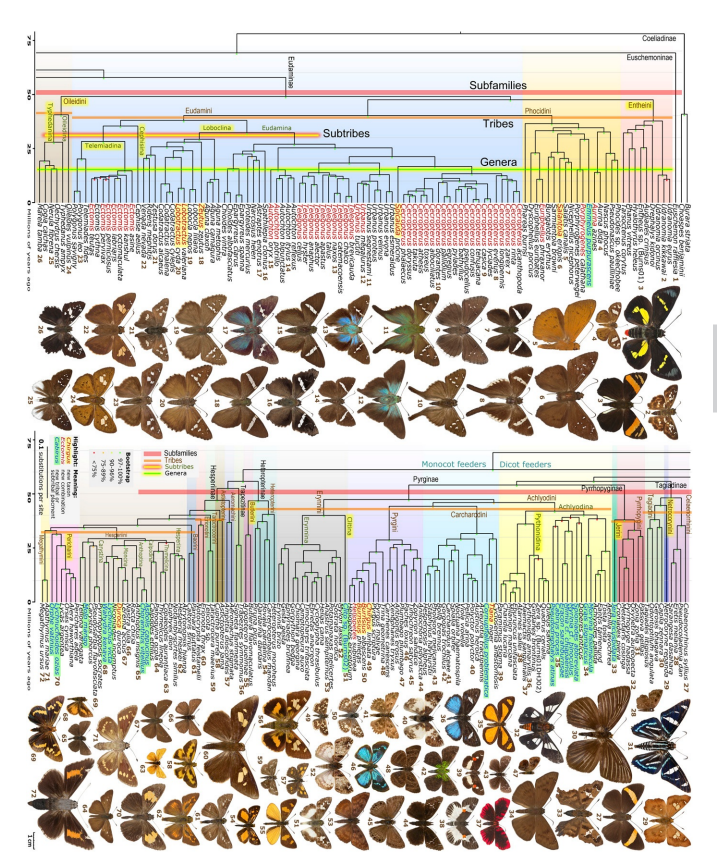


Fig. 1. Time-calibrated genomic tree of Hesperiidae constructed from nuclear gene coding regions of 250 species. The tree is cut in half: First and second segments are shown above and below. Colored horizontal lines delineate major taxonomic groups: red, subfamilies; orange (broken in several places), tribes; yellow shaded red (Eudaminae only), subtribes; and green shaded yellow (Eudaminae only), genera. New taxa are highlighted in yellow. New genera are in red font highlighted in yellow, and those used in different genus-species combinations than previously are in red font. Taxa placed in a different tribe or subtribe than previously are in blue font highlighted in green. Segments of the tree corresponding to different tribes are highlighted in different colors. Illustrated Hesperiidae species are associated with names in the tree by numbers in brown font. See SI Appendix for details.

subfamily of its own, sister to all other Hesperiidae except Coeliadinae. The next split in the genomic tree is different from that appearing in a number of DNA studies, but in agreement with the morphological view. The Spreadwing skippers (mostly dicot feeders) and the Grass skippers (mostly monocot feeders) are sisters. Both Spreadwing and Grass skippers have been previously divided into several subfamilies. Genomic data very strongly support (bootstrap above 99%) monophyly of these subfamilies. The two latest diverging subfamilies (Hesperiinae and Trapezitinae) have split off about 50 Mya (Fig. 1). Pyrginae, as defined by Warren et al. (3), diversified from their common ancestor before that time (>55 Mya). Moreover, while other subfamilies except Pyrginae (i.e., Eudaminae, Heteropterinae, and Hesperiinae) are well separated from each other, Pyrginae sensu Warren have split into three compact groups, within a short time and before divergence of Grass skippers into subfamilies. Thus, these three groups within Warren's Pyrginae are no less distinct than Hesperiidae subfamilies, and we treat them as such. Accordingly, we unify the sister tribes Tagiadini and Celaenorrhinini in a subfamily Tagiadinae, and reinstate the subfamily Pyrrhopyginae. The latter are strikingly distinct in appearance from other skippers, and have been traditionally considered a subfamily. They diverged from their common ancestor with Pyrginae about 55 Mya, before the divergence of Grass skippers. Thus, genomic data suggest that the family Hesperiidae consists of nine subfamilies (Fig. 1 and SI Appendix), and diversification into subfamilies occurred about 50 Mya.

Mitogenomes and Cytochrome Oxidase I Barcodes. In addition to a nuclear genome tree, we constructed a tree from mitogenomes. The resulting tree recapitulates major phylogenetic groups of the nuclear genome tree, but with weaker support. All of the subfamilies and tribes are composed of the same species in either mitogenome or nuclear genome phylogenies. Next, using mitogenomes of 250 species as a backbone, we increased taxonomic coverage by adding 290 species with cytochrome oxidase I (COI) barcodes only (SI Appendix, Fig. S1). These species grouped with mitogenomes of their expected relatives. Most of these barcode-only species were placed according to their current classification, with several exceptions discussed in SI *Appendix*. We used this mitogenome + barcode tree together with the nuclear genome tree (Fig. 1) as the basis for our proposed classification of Hesperiidae (SI Appendix, Taxonomic Appendix). Differences in barcodes and morphology suggested that 30 subspecies as defined by Evans (10–12) are more likely to be species, and such cases were analyzed in detail for wing pattern and genitalic differences (SI Appendix, Taxonomic Appendix).

Eudaminae Tribes and Subtribes. Previous studies refrained from defining tribes in the subfamily Eudaminae. We focused on this subfamily and attempted to delineate tribes consistently with the age of the tribes in Pyrginae (sensu stricto). The four Pyrginae tribes (Carcharodini, Achlyodini, Erynnini, and Pyrgini) outlined by Warren et al. (3) diverged around 42 Mya. Our genomic tree agrees with Warren's, except that it places Grais, Tosta, Morvina, Myrinia, Xispia, Pseudodrephalys, Mimia, Eracon, and Spioniades in Achlyodini; Cornuphallus in Carcharodini; Clito in Erynnini; and Jera in Pyrrhopyginae. Unexpectedly, Emmelus belongs to Eudaminae. Conversely, Cabirus is not Eudaminae (4), but Pyrginae: Achlyodini. All Pyrginae tribes received 100% bootstrap support and are conspicuous groups in the subfamily.

To achieve consistent classification, we cut the dated genomic tree around the time of Pyrginae divergence into tribes and find four Eudaminae phylogenetic groups supported by 100% bootstrap that we define as tribes. Two of the tribes that form best-separated groups are described in Table 1 as Oileidini and Entheini. The two others are closely related sisters Eudamini and Phocidini that diverged about 40 Mya, and are given a tribal rank due to their morphological distinction.

These four Eudaminae tribes correspond to groups with similar morphology (10, 11). For example, Entheini is largely the "B. Augiades group" of Evans. He defined it by the "divergent" third segment of palpi that stems from the outer edge of the second segment. Inconsistently, Evans included in this group three genera that have the central third segment: Phocides and Hypocryptothrix, that do not belong to Entheini, and Cabirus, that is not even in Eudaminae (Fig. 1). Interestingly,

Table 1. Description of new tribes and subtribes of Hesperiidae

New tribe or subtribe	Type genus	Diagnostic characters*	ZooBank ID [†]
Entheini Grishin, new tribe	Entheus Hübner, [1819]	276665.26:A192G; 85.28:C176T; 378.19:G1099C; 374.14:G1169T; keys to B.3a in ref. 10, but exclude B.9	303C1FD0-07CB-4919-900E-EA3D6347E5DD
Loboclina Grishin, new subtribe	Lobocla Moore, 1884	208.2:G145T, T146G; 18312.8:A619C, G620A; keys to B.4 in ref. 8 or C.5, C.10a, C.15.2 or C.18 in ref. 11	C606FC35-323D-4E55-AF5A-A86C6366BAFA
Cephisina Grishin, new subtribe	Cephise Evans, 1952	COI.bc:A44T, C84G, T479A; genitalia and palpi as described in ref. 17 (pp. 182–183) for Cephise are diagnostic	22B59811-F174-4FDF-A9D2-799897F4D44E
Telemiadina Grishin, new subtribe	Telemiades Hübner, [1819]	536.149:G1488C; 997.8:G514T; 860.7:A748G; 3001.3:C1773T; keys to B.2, C.3, C.7a (exclude C.7.6b), E.6a or E.9 in refs. 10 and 11	4AE0E59C-8B92-4C84-8651-E7A1C45C93C1
Oileidini Grishin, new tribe	Oileides Hübner, [1825]	1139.19:T562A; 851.8:C423A, G443A; 11945.11:G391A; 65.4:C330A; tuft of scales by anal fold from the base of hindwing, either above or below	CF9C3D29-523A-4D17-B140-9A69CFA98731
Typhedanina Grishin, new subtribe	Typhedanus Butler, 1870	1341.12:T25841C; 489.5:G307T; 3446.8:T2308A, C2309G, A2500C; tuft of scales by anal fold from the base of hindwing above, but not below	B4D56F93-67F9-476F-B69C-133D98BFBD58
Netrocorynini Grishin, new tribe	Netrocoryne C. & R. Felder, [1867]	2284.30:399A; 904.14:T439G; 275215.7:C925G; 998.8:G308A; 214.24:3520C; keys to B.1, C.1 or C.15 in ref. 8	DE61F048-02CF-4F8E-9392-D18A4618BABD
Jerini Grishin, new tribe	Jera Lindsey, 1925	103.23:796A; 420.27:G308A; 671.27:935C; 425.5:G1558T; keys to E.3 in ref. 11; forewing cell $>$ 3/5 of costa	AF3B5CEA-880A-4CB2-AF40-E6D87C39C040
Pythonidina Grishin, new subtribe	Pythonides Hübner, [1819]	274.29:G397A; 3478.6:T116C; 7985.5:G916A; 925.10:G199C; in ref. 11, keys to E.44a, E.49.1 or, if uncus undivided, then E.37a or 40d	CB890271-5483-4B5A-A7BC-27DBC5E23DE5
Clitina Grishin, new subtribe	Clito Evans, 1953	COI.bc:G29T, 81A, 169A, 266A, 302T, A353T, A521T; keys to E.52 or E.13.8 in ref. 11	971884E2-E5F7-46A3-B182-657729B6A778
Butleriini Grishin, new tribe	Butleria Kirby, 1871	2627.8:A1459T; 141.4:C104A; 37338.38:G133T, G134C; keys to H.4 & 5 in ref. 12	D621EF81-FA65-4858-9450-E0C041598D7A
Pericharini Grishin, new tribe	Perichares Scudder, 1872	596.8:C1601G; 144.41:G201C; 83.15:G8658A,T8657G; keys to K.27a in ref. 12	94B68BD2-7F83-4E58-80E1-7F5AC8C56511

ndix for the lists of genera included in each taxon and sequences of protein-coding regions with diagnostic characters

Notation 272.1:A192G means position 192 in the gene 1 on the scaffold 272 is G, changed from A in the ancestor; 169A, means position 169 is A, but the ancestral state is unclear; COI.bc is the COI barcode region [†] ZooBank registration URL given for each taxon should be preceded by http://www.zoobank.org/

Phareas that has a divergent third segment (and was included in the Augiades group) is not in Entheini. Its males possess, absent in Entheini, tufts of hair-like scales in the groove along the hindwing vein 1A+2A. Our tree places this unusual skipper in Phocidini, implying that its peculiar palpi are convergent.

Oileidini is sister to other Eudaminae (Fig. 1). Genera in this tribe were grouped with some Pyrginae genera by Evans (11), and the tribe may be intermediate in morphology between Pyrginae and Eudaminae. This is the smallest tribe (six genera) and is characterized by tufts of hair-like scales in the groove along the hindwing vein 1A+2A in males, either below (Oileides) or above (the others). Similar structures in *Phareas* (Phocidini) are found on both sides of the hindwing.

The sister tribes Eudamini and Phocidini are separated from each other by a short branch and could be one tribe. However, each is strongly monophyletic, and Phocidini skippers stand out morphologically and ecologically: Their forewing veins R4 and R5 originate near each other, hindwing tornus is usually expanded (not lobed), skippers hold wings spread flat when resting, many are crepuscular, and many species are sexually dimorphic. Our Phocidini is the "D. Celaenorrhinus group" of Evans (10) after adding Phocides, Phareas, and Emmelus and removing Cephise, which has lobed or tailed hindwing tornus, and Celaenorrhinus, whose males have a tuft of long scales on the hind tibiae that fits into a thoracic pouch (not found in Eudaminae). The genomic tree suggests that Oileides is polyphyletic: one species together with Aurina belong to Phocidini.

Eudamini is the largest and most diverse tribe. It encompasses more than half of the subfamily. The genomic tree reveals groupings within the tribe that are described here as subtribes (Table 1). One of them, subtribe Cephisina, is monotypic for the genus Cephise, which diverged from its sister tribe Telemiadina 35 Mya and is unique in its morphological features (17). Telemiadina includes three genera: Telemiades with its close sister Polygonus and Ectomis, into which we sink Hypocryptothrix, Heronia, Polythrix, Chrysoplectrum, and Speculum (see Uncanny Divergence Within a Genus). Along with Lobocla, Loboclina unifies genera with the arcuate antennal club from the "C. Urbanus group" of Evans plus Venada, Aguna, and a new genus, Zeutus (Table 2). Others belong to the "crown" group of Eudaminae. It includes an array of skippers that have been largely misclassified due to widespread and possibly mimetic convergence as detailed in Widespread Convergence in Wing Patterns and Shapes.

Eudaminae Genera and Subgenera. We define a major phylogenetic cluster of species with a common ancestor existing within a certain timeframe as being a genus (18). We cut a dated phylogenetic tree at a time point to maximize agreement with the current classification. This neither splits nor merges most genera that are well defined by morphological features, and we treat as genera the groups of species supported by the cut branches. The time 15 Mya corresponds to such point. It keeps well-known genera Aguna, Udranomia, and Urbanus proteus group unsplit. However, it separates traditional and morphologically distinct pairs of genera such as Epargyreus and Chioides. We attempt to reduce the number of monotypic genera, unless the genus is strongly distinct, because we wish to indicate relationships to other species by the name of a genus. As a result (Fig. 1 and SI Appendix), we delineated 50 Eudaminae genera, 4 of which are described as new in Table 2. The number of monotypic genera decreased from 10 to 4: Nicephellus, Spathilepia, and Zeutus (19), plus Emmelus transferred from Pyrginae. These four genera diverged from their sister taxa at least 18 Mya and are morphologically distinct.

Within some genera, we see groups of species that may be defined as subgenera (SI Appendix), and three new subgenera are described (Table 2). Some of the subgenera, such as Thorybes, have been used as genera for decades, but their genomic and morphological distinctness is smaller compared with most genera.

Widespread Convergence in Wing Patterns and Shapes. The most unexpected result of this study is the astounding number of misplacements of species into genera they do not belong to. The

Table 2. Description of new genera and subgenera of Hesperiidae

New genus or subgenus	Type species	Diagnostic characters*	Derivation of the name †	ZooBank ID [‡]
Tekliades Grishin, new genus	Thymele ramanatek Boisduval, 1833	68A, C90T, T145C, 412T, 553A, 583T; keys to I.1.9 in ref. 7	Masculine, a blend of [ramana] Tek and [Coe] liades	081564BA-DA0C-4C46-AEAB-6C00131AC8BD
Salantoia Grishin, new genus	Eudamus eriopis Hewitson, 1867	59C, A79T, T163A, 530T, 598A, T637A; D.3.2 or 3 in ref. 10; harpe flat, not hook-shaped	Feminine, a blend of Sala[tis] and [Sarmie]ntoia	3F82E9DE-A5A2-44B3-A13D-53CF8A673FAE
Spicauda Grishin, new genus	Goniurus procne Plötz, 1881	307T, T349A, 424(not T), G506A, T562A; keys to C.13.13c in ref. 10; harpe dorsally spiked	Feminine, a blend of spica and cauda	14D26B57-940C-407B-8E70-4E25203044B8
<i>Urbanoides</i> Grishin, new subgenus	Goniurus esmeraldus Butler, 1877	T49A, A85T or C, T212A, C542T, T544A, A607C or T, T619A or G; keys to C.13.6a in ref. 10	Masculine, means similar to Urbanus	20FAC3B6-F038-40A0-B182-3C7F32A40702
Zeutus Grishin, new genus	Cecropterus zeutus Möschler, 1879	A22T, C271A, T278A, A526T, T548C, A607T; genitalia as for <i>zeutus</i> in ref. 19 (p. 27)	Masculine, echoes the type species name	75715B9C-46AB-40F5-B738-420DABD56B63
Lobotractus Grishin, new genus	Eudamus valeriana Plötz, 1881	49A, T400A, 401T, A477G, 517T, C542T, T619A; as given for the "cyda group" in ref. 17 (p. 196)	Masculine, a blend of Lobo[cla] and [Coda]tractus	C6E5B5DF-1C74-4DBD-85C3-7285209F6F03
Caudatractus Grishin, new subgenus	Eudamus alcaeus Hewitson, 1867	355A, T556A, A592T; <i>Codatractus</i> (C.11 in ref. 10) with tailed hindwing	Masculine, includes tailed species of Codatractus	DF0F3C91-F56E-4B65-B86C-385A36F9D7FD
Asina Grishin, new subgenus	Eudamus asine Hewitson, 1867	T70A, T127A, T197C, 206T, 208A, A256T, T346A, 373A; keys to C.7.2a in ref. 10	Feminine, derived from the type species name	B3B7A6F6-A95C-4A2E-B9FB-80A7A8F86761
Tiana Grishin, new genus	Ebrietas niger Williams & Bell, 1940	T16C, A43T, G86A, T142C, A196G, T278A, 283C; keys to F.7.3 or 4 in ref. 11	Feminine, a blend of $T[osta]$ and [II] iana	B9382699-24FB-4466-B39B-94E6B544C425
Chirgus Grishin, new genus	Hesperia limbata Erschoff, 1876	85A, 205A, 223A, 241A, 263(not C), T277A, A415T, 479T, T574A; keys to G.1.2e or 9 in ref. 11	Masculine, a blend of <i>Chi</i> [lean] and [Py] <i>rgus</i>	7B1905F1-9471-4BBF-90BF-32360783AB1E
Burnsius Grishin, new genus	Syricthus communis Grote, 1872	205T, 223A, 241T, 263(not C), T277A, 479T; keys to G.1.5, 8, or 10a in ref. 11	Masculine, honors skipper taxonomist John M. Burns	48996B74-3AB1-4DEA-9A64-B8F112E62343
Duroca Grishin, new genus	Hesperia duroca Plötz, 1882	127T, 163C, 349C, T424C; keys to J.39.5a in ref. 12, harpe broad, hook-shaped	Feminine, echoes the type species name	476FE13C-5895-4139-BB11-44F835E21565

See SI Appendix for the lists of species included in each taxon

Sequence characters are for the COI barcode region and only their combination is diagnostic to distinguish from former genera of these species. Notation A79T means position 79 is T. changed from A in the ancestor: 59C means position 59 is C, but the ancestral state is unclear.

All names are treated as nouns in the nominative singular

[‡]ZooBank registration URL given for each taxon should be preceded by http://www.zoobank.org/

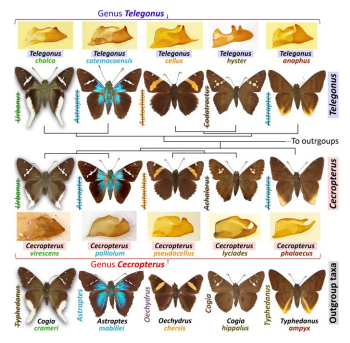


Fig. 2. Convergent wing patterns in *Telegonus, Cecropterus*, and others. Previously, they were placed in crossed out genera. Genitalic valvae are shown for the first two genera.

genera themselves, proposed over the years of classic entomological studies, mostly stood the test of genomic data: 55 genera were recognized before our work, and we revise them to be 50. We eliminated several monotypic genera for which visual morphological differences hindered close relationships with other species, and merged several phenotypically diverse but genotypically close genera (SI Appendix). Apparently, phenotypic distinction (e.g., wing shape, such as a tailed hindwing, or wing pattern, such as a pale stripe across the forewing) may be indicative of genetic differentiation. However, placement of a species into a genus by its dominant to human eye phenotypic feature is more problematic. For example, before our work, many tailed skippers were placed in the genus *Urbanus* based on the tail. Genomic data imply that half of them do not belong there, and we transferred them to three other genera. One genus is named here (Spicauda, Table 2), while the others have not previously included tailed skippers. Moreover, we transferred some skippers without long tails from Astraptes to *Urbanus*. They were formerly misclassified due to similar wing patterns consisting of shiny metallic-cyan wing bases and white forewing spots.

We found misclassifications to be widespread across Eudaminae and attribute them to convergence driven by the selection for large mimicry rings (20–22). This convergence is not confined to one or two basic patterns, but is more diverse. Some patterns are common in both the Old and New World, and, in addition to skippers, include butterflies from other families and even moths, flies, and beetles (20, 23). We find five different phenotypes (Fig. 2) that parallel each other in two genera (*Telegonus* and *Cecropterus*) and their more distant relatives (outgroups): (i) greenish bases of brown wings, white stripe on the forewing, and hindwing with a white tail and margins; (ii) metallic-blue wing bases, and forewing with white stripe (20); (iii) brown forewing with a yellow stripe across and apical white spots; (iv) cream-white, semitranslucent spots on the brown forewing; and (v) brown wings, and hindwing with yellow tornus.

At least four of these phenotypes are not ancestral and thus are convergent. Curiously, every species (10 out of 10) was placed by its appearance in a wrong genus before our study (crossed out genus in Fig. 2). In retrospect, assignment to a correct genus could have been possible through detailed comparison of male genitalia. In *Telegonus*, the dorsal side of the valva is concave in the middle and forms a mouth-like structure with two "kissing lips." In *Cecropterus*, the valva is dorsally and terminally convex, without a "kiss." These genitalic features agree with genomic phylogeny and reinforce our conclusions.

Uncanny Divergence Within a Genus. Before our work, each of the three genera *Ectomis*, *Hypocryptothrix*, and *Heronia* consisted of single species of unique appearance. No taxonomist had ever thought of them being a group. To our surprise, all phylogenetic trees we obtained (even from COI barcodes only) revealed only a slight divergence among these three and two other genera, *Polythrix* and *Chrysoplectrum*, suggesting that it is best to place all these skippers in a single genus, *Ectomis*. Moreover, their most divergent phylogenetic lineage was part of *Polythrix*, and is described here as a new subgenus *Asina* (Table 2). Barcode divergence among the subgenus *Ectomis* is within 10% and is less than within the genus of swallowtail butterflies *Pterourus* (24), which some researchers consider a subgenus of *Papilio*.

Despite limited genetic differentiation, the expanded *Ectomis* contains species of profound phenotypic divergence. All skippers in the former genus Polythrix (subsumed by Ectomis) are tailed. Others are not, although their hindwing is usually lobed at the tornus. While most *Ectomis* are brown skippers with a pale forewing band frequently divided into spots, some vary from solid dark brown to dark metallic green with a forewing central spot, or tawny with many white spots. Some are even part of the mimetic complex with brilliant blue thorax and wing bases above (20). Males of some species possess tufts of hairlike scales on the forewing below, while others have a double row of yellow spines on the hind tibiae. Male genitalia are as diverse as wing shapes and patterns, and the phylogenetic closeness is not apparent from genitalia. The valva varies from a simple curved plate without elaborations (in Ectomis cythna) to very complex with several processes (in "Heronia" labriaris). It would be of interest to investigate genetic mechanisms for such a rapid phenotypic divergence within Ectomis.

Connecting Genotype to Phenotype. Telegonus chalco (Fig. 2) and Telegonus brevicauda are sister species that do not look alike. T. brevicauda belongs to the "blue-wing-base" mimicry complex (Figs. 1–3) and is nearly indistinguishable from its more distant relative Urbanus tucuti (Fig. 1). We compared genomes of these three species to find rapidly diverging proteins between the two sisters (SI Appendix, Methods). Genes involved in the circadian clock system, transcription regulation, wing morphogenesis, fatty acid, and vitamin metabolism (SI Appendix, Tables S4 and S5) stood out. The differences in morphogenesis genes may be related to the hindwing tail development in T. chalco. Next, we looked for fixed mutations in the uniquely patterned T. chalco compared with the two similar species. Xanthine dehydrogenase

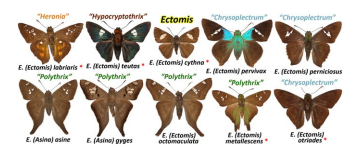


Fig. 3. Uncanny divergence within *Ectomis*. These skippers belonged to five genera listed above each image. Type species for these genera are marked with red asterisks.

had the largest number of such mutations (*SI Appendix*, Table S6). This enzyme is involved in pterin metabolism, and pterin derivatives were suggested to be responsible for wing coloration in pierid butterflies (25). The *T. chalco*-specific mutations may alter interactions between the enzyme and the transport proteins that deliver the enzyme to different pigment granules (26), and thus affect the wing color in *T. chalco*. Some of these mutations are present in other skipper species with extensive white scaling on hindwings.

Discussion: A Broad Picture. Today, and more so tomorrow, genome sequencing provides an efficient and cost-effective way to gain rapid insights about biodiversity. Given a taxonomic group, comparative analysis of genomic sequence along with morphology and ecology will be rich in discoveries. Even now, this can be done within a small laboratory. With the family Hesperiidae, we have shown how such a project can be accomplished and have reported some of the results that can be expected. Not only is this family interesting as a diverse group of butterflies, it also emphasizes several generalities. First, while the reference genomes required freshly collected specimens, the bulk of the project was done with old museum samples. Even specimens collected a century ago yielded usable genomic data. Second, we provided a genome-based phylogeny of the family and reclassified it taxonomically. We found that many species were classified differently than previously thought. Other families may display the same problem when examined genomically. Third, we encountered many examples of phenotypic convergence and divergence, and mined genomic data for the links between genotype and phenotype. With the ever-decreasing cost of sequencing, we expect that, soon, any phylogenetic project will start from sequencing of genomes.

Materials and Methods

For freshly collected specimens, DNA was extracted from tissue of a specimen (minus wings and genitalia that were kept in an envelope) field-stored

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in a vial with RNAlater. For pinned and dry specimens from museum collections, DNA was extracted from a whole abdomen or a leg. We have previously reported details of methods for DNA extraction, genomic library preparation, next-generation sequencing, and computational analysis of nuclear and mitochondrial genomes (13, 27). Phylogenetic trees were constructed with the programs RAxML (28) and ASTRAL (15), and were dated using the procedure described in *SI Appendix*, where further details of the methods are given. Genomic data obtained in the project have been deposited in the SRA database (accession SRP147939), and COI barcodes have GenBank accessions MH357724 through MH357835. ZooBank registration for this work, published on March 15, 2019, is 8A8E82C4AC8A-4B7B-8DF4-6BFF4BD48F3D.

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