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Cryptic species-level diversity in Dark-throated Oriole Oriolus xanthonotus

by Frank E. Rheindt, Meng Yue Wu, Nyanasengeran Movin & Knud A. Jønsson

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Summary.—Borneo is part of the Sundaland biodiversity hotspot, yet this large tropical island still harbours much unrecognised biodiversity. In this integrative study, we combine the results from phylogenomic, bioacoustic, biometric and morphological analyses, and show that the Sundaic species Dark-throated Oriole *Oriolus xanthonotus* comprises two species-level taxa, both of which occur on Borneo. The eastern species, here re-named Ventriloquial Oriole *O. consobrinus*, is characterised by plumage distinctions, most notably a more male-like coloration in females, and a characteristic ventriloquial song unique among the region's orioles. The precise contact zone of the two species on Borneo is incompletely mapped. Their populations on Borneo show pronounced character displacement whereby individuals of *O. xanthonotus* on other landmasses are significantly larger than Bornean individuals, presumably to prevent non-adaptive hybridisation with the larger Ventriloquial Oriole along their Bornean contact zone.

Borneo, the third largest island on Earth, forms part of the South-East Asian equatorial rainforest belt known as Sundaland (Fig. 1). The island boasts unusually high avian diversity and is well known ornithologically for its distinct montane endemic avifauna, featuring roughly 1–2 dozen endemic bird species depending on taxonomic treatment (MacKinnon & Phillipps 1993, Eaton $et\ al.\ 2021$). However, despite being $c.500\ km$ from the nearest part

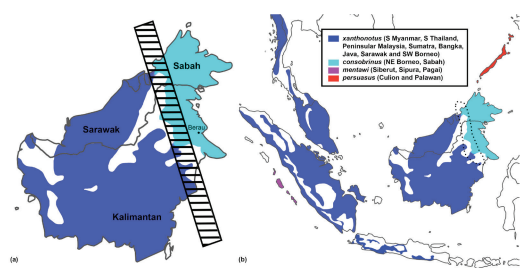


Figure 1. Distribution map of Dark-throated Oriole *Oriolus xanthonotus*, following Walther & Jones (2020). (a) Distribution of Bornean *O. x. xanthonotus* and *O. x. consobrinus*. Barred rectangle refers to possible contact zone between the two taxa. (b) Distribution map of all four taxa within the *O. xanthonotus* complex.

ISSN-2513-9894 (Online) of the Asian mainland in the Malay Peninsula, Borneo's endemic lowland avifauna has traditionally been considered rather limited. In the taxonomic traditions of the 20th century, the vast majority of Borneo's lowland rainforest birds were considered part of more widely distributed Sundaic species whose ranges often also include Sumatra, Java and Peninsular Malaysia (MacKinnon & Phillipps 1993, Sheldon *et al.* 2015). It has taken a steady stream of molecular and bioacoustic work to reveal that the populations of some Bornean lowland or hill forest birds are more distinct than previously anticipated, a few even at species level, thereby further boosting the island's number of endemic species (Moyle *et al.* 2005, Lim *et al.* 2010, 2017, Moyle *et al.* 2011, Ng *et al.* 2016, 2017a, Gwee *et al.* 2019, Cros *et al.* 2020).

Borneo's distribution of lowland endemism constitutes one of the world's most unusual biogeographic patterns, whose origin continues to require a satisfactory explanation. The greatest part of Borneo's lowland endemic birds—both among species traditionally recognised and those elevated to species via recent research—is restricted to the northeast of the island. Here labelled 'Sabah endemics' or 'Sabah specialists', these species are almost entirely confined to the state of Sabah, although some are known to spill over slightly into adjacent parts of North and East Kalimantan, Sarawak, and even Brunei. Most Sabah endemics are replaced over the rest of Borneo by a more widespread sister species that usually also occurs on Sumatra, Peninsular Malaysia, and sometimes even Java. This biogeographic division into Sabah and the remainder of Borneo along a suture zone, comprising multiple roughly parallel contact zones among avian taxa, exists against the backdrop of a conspicuous lack of geographic boundaries (Sheldon *et al.* 2009, Rheindt 2021), as the position of Borneo's largest rivers and mountains is incongruent with the divide.

The cataloguing of Borneo's avian endemism remains far from complete. Recent taxonomic treatments have suggested that splits between Sabah and the rest of Borneo may become necessary in approximately another dozen bird species (Eaton *et al.* 2021). In this study, we present evidence for one such taxonomic split in the Dark-throated Oriole *Oriolus xanthonotus* complex. This species is a typical Sundaic rainforest denizen, distributed from the southernmost tip of Myanmar in the Thai-Malay Peninsula through Sumatra, Java and Borneo to Palawan, in the south-west Philippines. Four subspecies have long been recognised, two of which occupy smaller marginal islands (*mentawi* on the three main Mentawai Islands of Siberut, Sipora and Pagi, off western Sumatra, and *persuasus* on Palawan and some smaller satellites; Fig. 1). Among the two more widespread subspecies, *consobrinus* is said to be restricted to northern, central and eastern Borneo (including Sabah), whilst nominate *xanthonotus* occurs in the remainder of Borneo, Sumatra, Java and the Thai-Malay Peninsula. The precise boundary between the two subspecies on Borneo remains to be elucidated.

Recent genomic work on Old World orioles (Jønsson *et al.* 2019) has uncovered unexpected relationships and divergences among Asian oriole species that possibly warrant taxonomic adjustments. Most of these refer to island taxa or populations otherwise isolated geographically. However, one complex that stands out is the Dark-throated Oriole, with two distinctive forms co-occurring on one landmass (Borneo) and presumably overlapping or abutting in range. Here we present parts of Jønsson *et al.*'s (2019) genomic data and add bioacoustic, biometric and morphological analyses to propose dividing Dark-throated Oriole into two species.

Materials and Methods

For genomic analyses and methods we refer to Jønsson et al. (2019). For phenotypic analyses, we measured 59 specimens of Dark-throated Orioles deposited in the Raffles



TABLE 1 List of specimens measured. All specimens are from the Raffles Collection in the Lee Kong Chian Museum of Natural History, Singapore.

		or reacting resort, on Superci			
Taxon	Landmass or state of collection	Locality	Collection date	Sex	Specimen registration number
xanthonotus	Sarawak	'Belingean' (probably Bukit Belingan south of Miri)	17/05/1917	Male	3.18788
xanthonotus	Sarawak	'Bukar', Samarahan (near Kuching)	25/10/1919	Male	3.18789
xanthonotus	Sarawak	'Bukar', Samarahan (near Kuching)	15/11/1919	Male	3.1879
xanthonotus	Sarawak	'Saribas'	01/03/1932	Male	3.18795
xanthonotus	Sarawak	'Lio Matu, Baram District' (now Lio Matoh, near Miri)	13/10/1920	Male	3.1878
xanthonotus	Sarawak	'Lio Matu, Baram District' (now Lio Matoh, near Miri)	23/10/1920	Male	3.18781
xanthonotus	Sarawak	'Long Musan', Baram, Sarawak	05/10/1920	Male	3.18782
xanthonotus	Sarawak	'Borneo, Baram'	05/03/1905	Male	3.18783
xanthonotus	Sarawak	'Sungei Lenian, Tenjar, Baram'	24/08/1919	Male	3.18786
xanthonotus	Sarawak	'Saribas'	01/01/1917	Male	3.18791
xanthonotus	Sarawak	'Saribas'	15/05/(illegible)	Male	3.18794
consobrinus	Sabah	'Bettotan' (now Betotan), Sandakan	11/08/1927	Male	3.18702
consobrinus	Sabah	'Bettotan' (now Betotan), Sandakan	14/08/1927	Male	3.18704
consobrinus	Sabah	'Bettotan' (now Betotan), Sandakan	22/08/1927	Male	3.18707
consobrinus	Sabah	'Bettotan' (now Betotan), Sandakan	22/08/1927	Male	3.18708
consobrinus	Sabah	Kudat	19/09/1927	Male	3.18713
consobrinus	Sabah	'Bettotan' (now Betotan), Sandakan	08/08/1927	Male	3.18698
consobrinus	Sabah	'Bettotan' (now Betotan), Sandakan	27/07/1927	Male	3.18697
consobrinus	Sabah	Rayoh	28/06/1928	Male	3.18718
consobrinus	Sabah	'Bettotan' (now Betotan), Sandakan	09/08/1927	Male	3.18700
consobrinus	Sabah	'Banguey Island' (now Banggi)	6/09/1927	Male	3.18694
consobrinus	Sabah	Kudat	17/09/1927	Male	3.18714
consobrinus	Sabah	'Malawalle Island' (now Malawali)	08/1927	Male	3.18716
consobrinus	Sabah	'Banguey Island' (now Banggi)	02/09/1927	Male	3.18693
consobrinus	Sabah	'Bettotan' (now Betotan), Sandakan	11/08/1927	Male	3.18701
consobrinus	Sabah	Rayoh	19/06/1928	Male	3.18717
xanthonotus	Sarawak	Baram	05/03/1905	Female	3.18784
xanthonotus	Sarawak	'South Paku, Paku Saribas'	28/10/1916	Female	3.18793
xanthonotus	Sarawak	Baram, Borneo	05/03/1905	Female	3.18785
xanthonotus	Sarawak	Wai Sa Baram, Borneo	27/10/1920	Female	3.18787
consobrinus	Sabah	'Bettotan' (now Betotan), Sandakan	08/1927	Female	3.18712
consobrinus	Sabah	Kudat	09/1927	Female	3.18715
consobrinus	Sabah	'Bettotan' (now Betotan), Sandakan	13/08/1927	Female	3.18703
consobrinus	Sabah	'Bettotan' (now Betotan), Sandakan	24/08/1927	Female	3.18711
consobrinus	Sabah	'Bettotan' (now Betotan), Sandakan	27/07/1927	Female	3.18696
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consobrinus	Sabah	'Banguey Island' (now Banggi)	07/09/1927	Female	3.18695
consobrinus	Sabah	'Bettotan' (now Betotan), Sandakan	09/08/1927	Female	3.18699
consobrinus	Sabah	'Bettotan' (now Betotan), Sandakan	19/8/1927	Female	3.18705
consobrinus	Sabah	'Bettotan' (now Betotan), Sandakan	21/08/1927	Female	3.18706
xanthonotus	Malay Peninsula	Chong Trang, North Malay Peninsula	01/12/1909	Male	3.18755
xanthonotus	Malay Peninsula	Temengoh now Temengor, Perak	02/08/1909	Female	3.18751
xanthonotus	Malay Peninsula	Cheras, Selangor	18/3/1908	Male	3.18769
xanthonotus	Malay Peninsula	Pelipak, Johor	29/3/1905	Male	3.18776
xanthonotus	Malay Peninsula	Pelipak, Johor	27/3/1905	Female	3.18775
xanthonotus	Malay Peninsula	Gunung Angsi, Negri Sembilan	4/1906	Female	3.18771
xanthonotus	Malay Peninsula	Klang, Selangor	14/1/1909	Male	3.18770
xanthonotus	Sumatra	Kota Pinang	4/6/1937	Male	3.18797
xanthonotus	Sumatra	Kota Pinang	5/7/1937	Male	3.18798
xanthonotus	Sumatra	Kota Pinang	7/8/1937	Male	3.18799
xanthonotus	Sumatra	Kerinci, Sumatra	28/11/1914	Female	3.18796
xanthonotus	Sumatra	Padang, Sumatra	14/11/1934	Male	3.18800
xanthonotus	Java	Wynkoops Bay now Pelabuan Ratu	13/3/1920	Male	3.18804
xanthonotus	Java	Wynkoops Bay	8/3/1920	Male	3.18803
xanthonotus	Java	Wynkoops Bay	8/3/1920	Male	3.18802
xanthonotus	Java	Wynkoops Bay	7/3/1920	Male	3.18801
xanthonotus	Java	Wynkoops Bay	19/3/1920	Male	3.18805
persuasus	Palawan (Philippines)	Puerto Princesa	17/09/1925	Female	3.18808
persuasus	Palawan (Philippines)	Puerto Princesa	08/08/1925	Female	3.18806
persuasus	Palawan (Philippines)	Puerto Princesa	13/08/1925	Male	3.18807
mentawi	Sumatra	Siberut Island	24/9/1924	Male	3.3968
mentawi	Sumatra	Sipora Island	20/10/1924	Female	3.3969
mentawi	Sumatra	Siberut Island	16/9/1924	Male	3.3967

Collection of the Lee Kong Chian Museum, Singapore, including three persuasus, three mentawi, 21 consobrinus and 32 nominate xanthonotus (Table 1). The latter taxon was represented by specimens from across multiple major landmasses (Table 1). Specimens were arranged according to taxon, landmass, collection date and sex, and were measured as follows: flattened wing length (from the carpal joint to the tip), maxilla length (from the tip to where the bill meets the skull), tail length (from the longest rectrix tip to the uropygial gland) and tarsus length (from tibiotarsus). Measurements of each parameter were performed by a single person (MYW: tail and wings; NM: maxilla and tarsus). All specimens were inspected for plumage coloration. Biometric comparisons of individuals from different landmasses and taxa were undertaken using principal component analysis (PCA) via the 'prcomp' function in R version 4.0.2 (R Core Team 2020), with PCA plots drawn using the 'ggplot2' package. We also carried out t-tests in R to identify any traits that might be significantly different between taxa.

Old World orioles are known for their fluty and variable songs. Song structure, including number of elements as well as their frequency and duration, varies greatly within and



among populations. Therefore, bioacoustic analyses that include sonogram measurements of individual call notes are not meaningful because of difficulties in assigning homology. Bioacoustic variability notwithstanding, Dark-throated Oriole calls can be assigned to particular call types within each population based on composite properties such as call structure, number of elements and frequency modulation. Some of these call types are specific to a particular taxon, whereas others are shared. Our bioacoustic analysis therefore centred on mapping the overlap of call types among different taxa. We gathered a total of 147 sound recordings from xeno-canto (www.xeno-canto.org), the Macaulay Library (https://macaulaylibrary.org) and private recordings (B. van Balen; available on request) for qualitative inspection. These vocalisations were categorised into 11 and ten call types for the eastern and western populations, respectively.

Results

Genomic analysis.—Jønsson *et al.* (2019) analysed two mitochondrial and two genomewide DNA data partitions across the genus *Oriolus*. We summarise and provide additional detail on the results of each of these four datasets with emphasis on the Dark-throated Oriole complex (Fig. 2).

Analysis of the single mitochondrial gene ND2 (1,041 base pairs [bp]) uniquely encompassed individuals of all four subspecies, including island representatives for Java, Sumatra, the Thai-Malay Peninsula, Borneo, Palawan and Mentawai. This analysis revealed

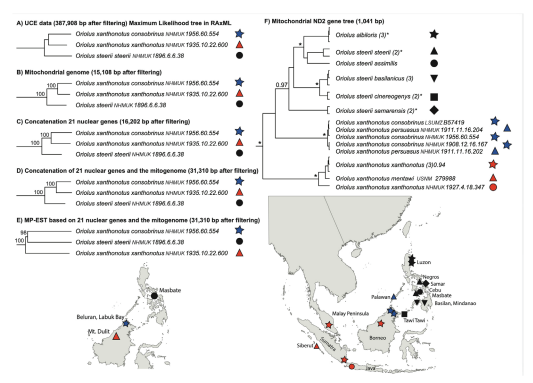


Figure 2. Phylogenetic trees based on four different data partitions adapted from Jønsson *et al.* (2019). A: nuclear ultraconserved element (UCE) loci; B: mitochondrial genomes; C: 21 independent nuclear sequence loci; F: mitochondrial NADH dehydrogenase 2 (ND2) gene sequences. D–E show trees based on a combination of the 21 nuclear sequence loci (C) and the mitochondrial genomes (B), constructed using the concatenation approach (D) and the species-tree approach as implemented in MP-EST (E). Maps show sampling localities for the phylogenetic trees placed above.

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ISSN-2513-9894 (Online) O. xanthonotus to be paraphyletic, with strong support for a closer relationship between the two eastern subspecies, O. x. consobrinus and O. x. persuasus, and the Philippine species White-lored Oriole O. albiloris and Philippine Oriole O. steerii, to the exclusion of the two western subspecies O. x. xanthonotus and O. x. mentawi (Fig. 2). Basal divergence between the 'western' dark-throated taxa (O. x. xanthonotus and O. x. mentawi) and the remaining orioles was estimated to have occurred c.7 million years ago, whereas divergence between 'eastern' dark-throated taxa (O. x. consobrinus and O. x. persuasus) and the two Philippine species was estimated to have occurred c.5 million years ago (fig. 1 in Jønsson et al. 2019).

For a smaller set of taxa representing all major Oriolus clades including an O. x. xanthonotus sample from Borneo, O. x. consobrinus and Philippine O. steerii, Jønsson et al. (2019) used shotgun sequencing to generate genome-wide sequences, resulting in the following four datasets:

In contrast to the single ND2 gene, analysis of more than 15,000 bp of the mitochondrial genome, containing ND2 and 15 other genes, supported monophyly of eastern and western representatives of Dark-throated Orioles (Fig. 2), with strong support for their being more closely related to each other than to Philippine Oriole (fig. 2B in Jønsson et al. 2019).

Analysis of a concatenation of 16 nuclear genes, amounting to >16,000 bp, agreed with the mitogenomic analysis in placing eastern and western Dark-throated Orioles closer to each other than to Philippine Oriole (Fig. 2; fig. 2C in Jønsson et al. 2019).

A concatenation of genome-wide ultra-conserved elements, amounting to almost 388,000 bp, and therefore surpassing the previous dataset almost 25-fold, had a more equivocal outcome: O. x. xanthonotus and O. x. consobrinus were placed as sister to each other, with Philippine Oriole as sister to these two, but with no strong branch support (Fig. 2). Moreover, the branch separating Philippine Oriole from the two dark-throated taxa was very short (fig. 2A in Jønsson et al. 2019).

In a species-tree analysis, as opposed to a concatenation analysis, using 21 nuclear genes and the entire mitogenome, O. x. consobrinus emerged as sister to Philippine Oriole, whilst nominate O. x. xanthonotus was more distant to these two, with fairly strong branch support (Fig. 2; bootstrap 98; see supplementary fig. S6 in Jønsson et al. 2019). This pattern was similar to the result of the ND2 analysis (Fig. 2). Species-tree analyses do not combine all loci into one super-alignment, but consider the phylogenetic information of each locus independently, and are widely regarded as the state of the art in phylogenetic analysis (e.g., Edwards et al. 2007).

Morphometric analysis.—The PCA plot for morphometric measurements clearly divided Sabah and Sarawak specimens along PC1, accounting for 38.8% of variation (Fig. 3). Generally, O. x. xanthonotus specimens from Sarawak grouped closer with O. x. xanthonotus individuals from the Thai-Malay Peninsula, Sumatra and O. x. mentawi from the West Sumatran Islands, whereas O. x. consobrinus from Sabah clustered more closely with O. x. persuasus from Palawan. Specimens of O. x. xanthonotus from Java were embedded with O. x. xanthonotus specimens from other landmasses (Fig. 3).

Morphometric measurements of Sabah vs. Sarawak specimens exhibited nearly nonoverlapping ranges and significant differences in all parameters except tail length (Fig. 4). Javan O. x. xanthonotus emerged as significantly different in wing and tarsus length compared to other O. x. xanthonotus populations. In O. x. mentawi from the West Sumatran Islands, only wing length was significantly different from the nearby mainland Sumatran O. x. xanthonotus population.

Plumage inspection.—The plumages of *O. x. xanthonotus* from the Thai-Malay Peninsula were similar to those of O. x. xanthonotus from eastern Borneo but differed considerably



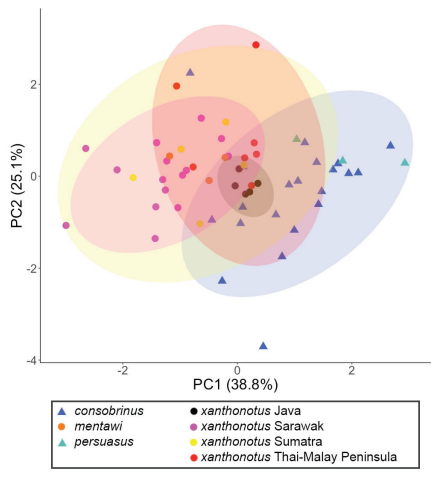


Figure 3. Principal component analysis plot for measurements across all four morphometric parameters. The 95% confidence interval ellipses are shown with the following colours: O. x. consobrinus (blue), O. x. *xanthonotus* Sarawak (pink), *O. x. xanthonotus* Java (grey) and the rest of the *O. x. xanthonotus* range (yellow). The percentage of total variation explained by each principal component (PC) is shown in parentheses.

from O. x. consobrinus in Sabah and O. x. persuasus on Palawan, which two showed similar plumage patterns.

Female O. x. xanthonotus has shorter, thinner and sparser streaking on the underparts, with a paler background coloration on the upper breast and a greyish-olive head that grades evenly into the upper mantle. This contrasts with female O. x. consobrinus and O. x. persuasus, which have longer, thicker and denser underparts streaking, a greyish background to the upper breast and a dark grey head that clearly contrasts with the upper mantle, giving their plumage a male-like character (Fig. 5).

Male O. x. xanthonotus also has shorter, thinner and sparser streaking on the underparts than both O. x. consobrinus and O. x. persuasus. Males of the latter show a diffuse border between the black throat and white upper breast, whereas O. x. consobrinus has a sharp divide between the black throat and white upper breast (Fig. 5).

Bioacoustic analysis.—We found two distinct vocal groups within the Dark-throated Oriole complex based on visual inspection of sonograms. Populations in the Thai-Malay Peninsula, Sumatra and Sarawak constitute one group separate from populations of Sabah and Palawan. Each of the two groups is characterised by sets of song and call types that

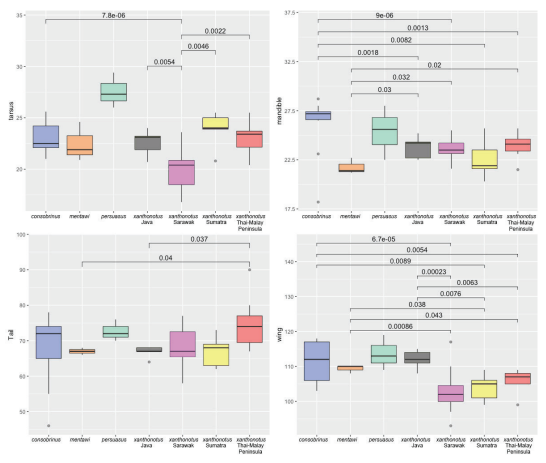


Figure 4. Box plots of morphometric measurements for tarsus (upper left), tail (lower left), maxilla (upper right) and wing length (lower right). Significant p values are shown above the plots for each corresponding pairwise comparison. Wing and tarsus comparisons involving O. x. persuasus overwhelmingly emerged as significant, and are not shown because of space constraints.

do not overlap (Figs. 6-7). At the same time, each group was found to emit call types of either inflected or deflected single notes which, however, differ greatly in duration between groups (Figs. 6–7: top row).

In contrast to the more conventional fluty multisyllabic oriole calls uttered by nominate O. x. xanthonotus (Fig. 6), the commonest calls in the Sabah and Palawan populations (O. x. persuasus and O. x. consobrinus) have a distinct ventriloquial quality (Fig. 7), more reminiscent of the sympatric Bornean Black Magpie Platysmurus aterrimus. Many of these ventriloquial calls exhibit a three-note structure, but with variations, sounding roughly like a monotonous du-du-duuuu with a variable ending, either inflected (e.g. XC 360969, XC 269344) or deflected (e.g. ML 202577).

Whilst O. x. xanthonotus from Java shares multiple call types with remaining nominate populations, we detected two unique call types (Fig. 6, blue frame) with a 'bulbul-like' (XC 618465) or 'plaintive cuckoo-like' (XC 393776) quality, quite unlike the typical fluty timbre that characterises the vocalisations of other populations. However, because of a low vocal sample size for Java, it is unclear whether these calls are of taxonomic importance or are merely rare vocal variants.



Figure 5. Photographs of female (left column) and male (right column) Dark-throated Oriole Oriolus xanthonotus specimens at the Lee Kong Chian Natural History Museum (Singapore). Dorsal (top row), ventral (middle row) and lateral views (bottom row). Specimens grouped by taxa, from left to right: Peninsular Malaysian O. x. xanthonotus, O. x. mentawi, O. x. consobrinus and O. x. persuasus (females); Peninsular Malaysian O. x. xanthonotus, Javan O. x. xanthonotus, O. x. mentawi, O. x. consobrinus and O. x. persuasus (males) (Movin Nyanasengeran)

Discussion

Character displacement on Borneo.—Steep mensural differences between the two species on Borneo suggest that ecological and evolutionary character displacement may be moulding their body size and shape. Character displacement—also termed reinforcement is an evolutionary phenomenon whereby populations of closely related species possess similar traits across most of their non-overlapping ranges, but adopt greater trait differences in areas where they co-occur thereby preventing competition (Brown & Wilson 1956, Hoskin et al. 2005). In South-East Asian birds, character displacement has been mapped for vocal traits in Ptilinopus fruit doves (Rheindt et al. 2011) and Rhipidura fantails (Ng et al. 2017b), but also for mensural characters in Pachycephala whistlers (Ashari et al. 2018) in a pattern reminiscent of Dark-throated Orioles on Borneo.

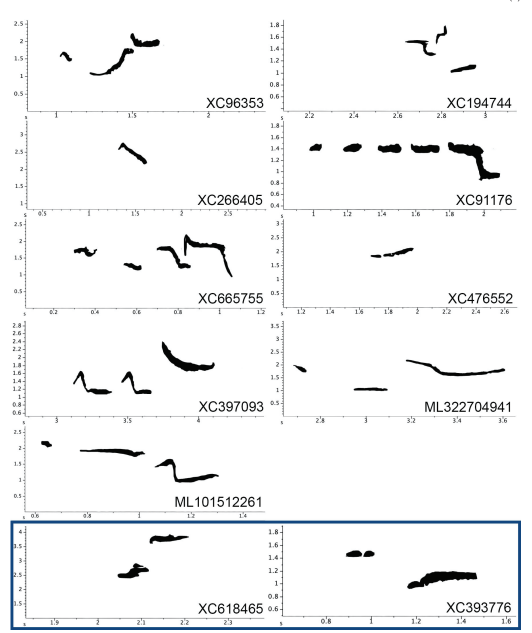


Figure 6. Sonograms of typical examples of the 11 song and call types identified across the Malay Peninsula, Sumatra, Java and Sarawak, reflecting the distribution of O. x. xanthonotus. Javan individuals were found to utter two unique call types that differ from all other recordings (blue frame). Frequency axis in kHz.

Measurements of tarsus length, wing length and bill length suggest that easterly O. x. consobrinus from Sabah is considerably larger than Bornean populations of nominate O. x. xanthonotus, with almost non-overlapping size ranges (Fig. 4). In a PCA combining all four morphometric characters, the 'clouds' of these two groups are nearly non-overlapping (Fig. 3). However, when Sabah O. x. consobrinus is compared with nominate O. x. xanthonotus outside Borneo, we find a large overlap in measurements, and differences are not significant (Figs. 3-4). Measurements of both Bornean populations (O. x. xanthonotus and O. x.

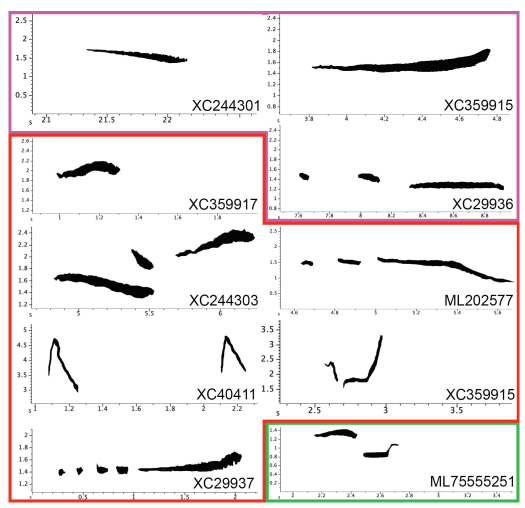


Figure 7. Sonograms of typical examples of the ten song and call types identified across Sabah and Palawan, reflecting the distribution of O. x. consobrinus and O. x. persuasus. The two populations shared three common call types (purple), whilst other call types may be distinct to each population, Sabah (red) and Palawan (green). Frequency axis in kHz.

consobrinus) may be driven apart to minimise the chances of reproductively suboptimal hybrid offspring in their contact zone. In biological terms, this has led to a Dark-throated Oriole (O. x. xanthonotus) on Borneo that is much smaller than populations throughout the rest of the range, and outsized by the O. x. consobrinus which it encounters on Borneo.

Mapping the contact zone in Borneo.—The precise extent of the ranges of various Sabah endemics or Sabah specialties on Borneo remains incompletely known, especially for taxa that have only recently been recognised at species level (e.g. Glissando Babbler Pellorneum saturatum vs. Leaflitter Babbler P. poliogene and others; Eaton et al. 2021, Garg et al. 2022). The same uncertainty applies to Dark-throated Orioles. Whilst the internet permits access to much photographic and sound material from Sabah, there is hardly any such material from the rest of Borneo. As of November 2021, sound recordings deposited on www.xeno-canto.org indicated that O. x. consobrinus extends south from Sabah at least to the Berau District of East Kalimantan (Fig. 1), where a recording by I. Woxvold has the vocal hallmarks of eastern O. x. consobrinus (XC 65382). Further north, the transition between the two appears to be close to the borders of Sabah, as recordings and personal observations from Brunei indicate the presence of the western (O. x. xanthonotus) fluty song and plumage types there (NM pers. obs.). Field ornithologists and hobby birdwatchers can contribute much to our understanding of Bornean ornithology by filling such knowledge gaps and thereby helping to achieve a more complete mapping of the distribution of the island's bird species.

Genomic evidence for a split.-Jønsson et al.'s (2019) extensive genomic analyses produced a surprisingly heterogeneous picture of the phylogenetic relationships of Darkthroated Oriole taxa and closely allied species in the Philippines. Analysis of a widely used mitochondrial gene unexpectedly pointed to a close relationship between the two eastern subspecies of Dark-throated Oriole (O. x. consobrinus and O. x. persuasus), Philippine Oriole and White-lored Oriole, whereas western dark-throated taxa (O. x. xanthonotus and O. x. mentawi) were phylogenetically more distant (Fig. 2). Importantly, this paraphyletic arrangement of the Dark-throated Oriole was backed by strong branch support and would have been widely interpreted as reflecting the true relationships in the complex c.10-20 years ago before the advent of modern next generation sequencing technology. After expanding the mitochondrial dataset to include 15 additional genes, however, Jønsson et al. (2019) found that the odd paraphyletic arrangement of Dark-throated Orioles was not corroborated (Fig. 2), and that they may -after all-form a monophyletic species.

Use of mitochondrial DNA in phylogenetics is known to generate artefactual relationships in cases when DNA introgression and instances of hybridisation may have confounded its gene tree (e.g. Rheindt & Edwards 2011, Andersen et al. 2021). A new era of phylogenomics based on genome-wide DNA has revolutionised the field and made routine the use of nuclear DNA loci. The most extensive nuclear dataset analysed by Jønsson et al. (2019) encompassed almost 388,000 bp from across the nuclear genome of these orioles, surpassing the volume of the mitogenomic dataset 25-fold. This expansive dataset provided only weak support for a monophyletic placement of eastern and western Dark-throated Orioles (Fig. 2). In addition, a species-tree analysis of 21 nuclear genes combined with the mitogenomes shifted support back to placing eastern O. x. consobrinus closer to Philippine Oriole rather than to western O. x. xanthonotus (Fig. 2).

In summary, the most likely explanation for the heterogeneous outcome of phylogenomic analyses—despite the massive DNA sequence volumes investigated—is a similar timing for the split among three lineages, one of which is the ancestor of White-lored and Philippine Orioles, whilst the other two represent the eastern and western subspecies groups of Darkthroated Oriole, respectively. The earlier timing of the split between eastern and western Dark-throated Orioles, preceding the diversification between this lineage of orioles in the Philippines, strongly supports elevation of the eastern genetic clade of Dark-throated Oriole to species level as O. consobrinus.

Biological evidence for a split.—Our morphometric and plumage analyses largely corroborate the work of early museum taxonomists who documented considerable differences in coloration especially in females. Eastern females are considerably more contrasting, saturated and male-like in appearance, and no cline in such coloration differences was evident in our sample of 59 specimens. Male differences, while less pronounced, are nevertheless readily discernible in the field and in specimens (Fig. 5). There is variation in the female plumage, with consobrinus exhibiting a more blackish crown similar to male plumage compared to xanthonotus on Borneo. This tendency for females to develop male-like plumage occurs elsewhere in the genus Oriolus but rarely in sympatry (e.g., Black-and-crimson Oriole O. cruentus; Kirwan et al. 2021). In the parapatric



populations of consobrinus and xanthonotus on Borneo, these differences in female plumage may present a prezygotic impediment to gene flow.

Furthermore, there are substantial bioacoustic differences between eastern and western Dark-throated Orioles, which may even outweigh the color differences in reproductive importance. The western subspecies group utters many versions of a typical multisyllabic fluty oriole song (Fig. 6) roughly reminiscent of other species in the region (e.g., Black-naped Oriole O. chinensis). This typical fluty song is unknown in the eastern subspecies group, whose primary vocalisations chiefly comprise a series of ventriloquial calls reminiscent of Bornean Black Magpie *Platysmurus aterrimus* (Fig. 7).

On the combined basis of phylogenomic, biometric, bioacoustic and plumage data, we propose that the two eastern subspecies of Dark-throated Oriole (O. x. consobrinus [northern and eastern Borneo] and O. x. persuasus [Palawan]) be recognised as a separate species, O. consobrinus. We propose 'Ventriloquial Oriole' as the vernacular name of this newly recognised species, reflecting its characteristic song, which is unique among the region's orioles. The western subspecies group—nominate O. x. xanthonotus and O. x. mentawi—still occupies the largest part of the distribution of this complex; therefore retention of the name 'Dark-throated Oriole' for the western group appears to be the best solution leading to the least nomenclatural disruption.

Taxonomic uncertainty of the Javan population.—'Oriolus Xanthonotus' [sic] was first described by Horsfield in 1821 with 'Java' as its type locality, rendering the taxonomic identity of the Javan population of special importance. In plumage and morphometrics, Javan specimens squarely fit within the range of variation displayed by nominate O. x. xanthonotus on other landmasses, especially Sumatra and the Thai-Malay Peninsula (Figs. 3–5). Therefore, use of the subspecies name xanthonotus as a moniker for the more westerly species of Dark-throated Oriole is not in doubt. In mtDNA, Javan populations are characterised by the deepest divergence within the O. x. xanthonotus cluster (Fig. 2), but still compatible with subspecies status. Vocal material from Java is currently limited, consistent with the defaunation that has occurred here during the last three decades (Eaton et al. 2015, Symes et al. 2018, Verma et al. 2020) leading to this population's decline and endangerment. Whilst the existing vocal material is suggestive of some potential differentiation of Java's nominotypical population (Fig. 6, blue frame), the possibility of unusual call variants cannot be eliminated, and more extensive sampling of Javan call notes is required.

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