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Phylogenetic relationships of the Helmeted Woodpecker (Dryocopus galeatus): A case of interspecific mimicry?

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ABSTRACT
Examples of phenotypic convergence in plumage coloration have been reported in a wide diversity of avian taxonomic groups, yet the underlying evolutionary mechanisms driving this phenomenon have received little scientific inquiry. We document a striking new case of plumage convergence in the Helmeted Woodpecker (Dryocopus galeatus) and explore the possibility of visual mimicry among Atlantic Forest woodpeckers. Our multilocus phylogenetic analyses unequivocally place D. galeatus within Celeus, indicating that the former has subsequently converged in appearance upon the distantly related and syntopic Dryocopus lineatus, to which it bears a remarkable resemblance in plumage coloration and pattern. Although details of the Helmeted Woodpecker’s ecology and natural history are only now beginning to emerge, its smaller size and submissive behavior are consistent with predictions derived from evolutionary game-theory models and the hypothesis of interspecific social-dominance mimicry (ISDM). Moreover, estimates of avian visual acuity suggest that size-related mimetic deception is plausible at distances ecologically relevant to Celeus and Dryocopus foraging behavior. In light of our results, we recommend taxonomic transfer of D. galeatus to Celeus and emphasize the need for detailed behavioral studies that examine the social costs and benefits of plumage convergence to explicitly test for ISDM and other forms of mimicry in these Atlantic Forest woodpecker communities. Future field studies examining potential cases of competitive mimicry should also take into account the mimic’s acoustic behavior, particularly in the presence of putative model species and other heterospecific competitors, as any discontinuity between morphological and behavioral mimicry would likely preclude the possibility of deception.

Keywords: Dryocopus galeatus, mimicry, molecular phylogenetics, plumage convergence

Relaciones filogenéticas de Dryocopus galeatus: ¿Un caso de mimetismo inter-específico?

RESUMEN
Se han reportado ejemplos de convergencia fenotípica en la coloración del plumaje en una amplia diversidad de grupos taxonómicos de aves. Sin embargo, los mecanismos evolutivos subyacentes que guían este fenómeno han sido pocos estudiados. Aquí documentamos un nuevo caso sorprendente de convergencia de plumaje en Dryocopus galeatus y exploramos la posibilidad de mimetismo visual entre los pájaros carpinteros del Bosque Atlántico. Nuestro análisis filogenético de múltiples loci ubicó inequívocamente a D. galeatus dentro de Celeus, indicando que el primero ha subsecuentemente convergido en apariencia a la especie distante y sintópica D. lineatus, a la cual se parece notablemente en cuanto a coloración y patrón del plumaje. Aunque solo recientemente han comenzado a emergir los detalles de la ecología y la historia natural de D. galeatus, su tamaño menor y su comportamiento sumiso son consistentes con las predicciones derivadas de los modelos de la teoría evolutiva de juegos y la hipótesis de mimetismo social dominante inter-específico (MSDI). Más aún, las estimaciones de agudeza visual de las aves sugieren que el engaño mimético relacionado con el tamaño es plausible a distancias ecológicamente relevantes de comportamiento de forrajeo de Celeus y Dryocopus. A la luz de nuestros resultados, recomendamos la transferencia taxonómica de D. galeatus a Celeus y enfatizamos la necesidad de estudios detallados de comportamiento que examinen los costos y beneficios sociales de la convergencia del plumaje para evaluar explícitamente el MSDI y otras formas de mimetismo en estas comunidades de pájaros carpinteros del Bosque Atlántico. Los futuros estudios de campo que examinen los casos potenciales de mimetismo competitivo deberían también considerar el comportamiento de mimetismo acústico, particularmente en presencia de especies modelo putativas y de otros competidores hetero-específicos, ya que cualquier discontinuidad entre el mimetismo morfológico y el comportamental probablemente excluiría la posibilidad de engaño.

Palabras clave: convergencia de plumaje, Dryocopus galeatus, filogenética molecular, mimetismo.
INTRODUCTION

Recent progress in assembling the avian tree of life has shed light on numerous instances of nonaposematic plumage convergence in disparate taxonomic groups (Weckstein 2005, Weibel and Moore 2005, Tello et al. 2009, Jonsson et al. 2010). The highest incidence, and perhaps the most comprehensive examples, of this phenomenon occur within the woodpeckers (Picidae), which exhibit convergent evolution of elaborate plumage patterns in ≥11 genera (Webb and Moore 2005, Weibel and Moore 2005, Benz et al. 2006, Moore et al. 2006, Fuchs et al. 2008). Although interspecific visual mimicry has long been suspected in various cases of avian plumage convergence, the adaptive significance and underlying socio-ecological mechanisms that promote phenotypic similarity in the absence of aposematism have received little attention and remain poorly understood compared with other forms of mimicry (Wallace 1869, Cody 1969, Diamond 1982, Ruxton et al. 2004, Rainey and Grether 2007).

Alfred Russel Wallace (1869) hypothesized that visual mimicry may explain the serial evolution of plumage convergence between Old World orioles (Oriolus) and Australasian friarbirds (Philemon) codistributed across Wallacea. Assuming a classic 3-player system comprising a model, a mimic, and a third-party observer, Wallace reasoned that the smaller, subordinate Oriolus species were mimicking the appearance of larger, highly aggressive friarbirds to avoid attack by hawks or other socially dominant nonmodel species. An alternative hypothesis was proposed over a century later by Diamond (1982), who argued that the subordinate Oriolus mimics were instead incurring social benefits by directly deceiving the Philemon models, thereby minimizing interspecific aggression at highly contested foraging sites and gaining access to nectar resources that would otherwise be unavailable. Diamond’s 2-player hypothesis was largely based on cursory field observations of social interactions between Philemon and Oriolus species across the Australo-Papuan region, and thus he remained unclear on whether third-party deception was also necessary to promote and maintain competitive mimicry.

Other researchers have invoked hypotheses of natural selection for enhanced interspecific signaling to explain convergent evolution of phenotypic similarities in birds. Moynihan (1968) theorized that convergence in plumage coloration may foster more efficient interspecific communication within mixed-species foraging flocks by co-opting adaptive signal–receiver biases. By contrast, Cody (1969) proposed that phenotypic similarities may actually enhance interspecific territoriality between ecological competitors by eliciting heightened aggression. He examined several cases of plumage convergence within woodpeckers (Dinopium–Chrysocolaptes, Meiglyptes–Hemicircus, Dryocopus–Campephilus, Micropternus–Blythipicus) as well as African bush-shrikes (Chlorophoneus–Malaconotus) and reasoned that visual mimicry would promote more efficient interspecific communication and exclusion of potential ecological competitors, given that these same social signals presumably form the basis for conspecific territorial interactions. However, this hypothesis has received criticism for its inconsistency with competitive exclusion theory and its failure to distinguish when convergent evolution should be favored over character displacement (Murray 1976, Prum 2014).

The primary theoretical deficiencies inherent in these previous works have recently been addressed by Prum and Samuelsion (2012), who developed an explicit evolutionary framework derived from game theory to examine the selection forces associated with nonaposematic visual mimicry. Elaborating upon the classic hawk–dove game, they used a well-documented case of plumage convergence between 2 North American woodpeckers (Picoides villosus and P. pubescens; Weibel and Moore 2005) to estimate the coevolutionary fitness dynamics between model and mimic, thereby establishing the conditions that promote evolution of interspecific social-dominance mimicry (ISDM). Prum and Samuelson (2012) defined ISDM as a type of social parasitism in which a smaller subordinate species uses visual deception to minimize competitive interference with a dominant model taxon and gain access to enhanced feeding opportunities. Several predictions with respect to the players’ ecology, behavior, and body size have emerged from these analyses that should further facilitate testing of ISDM in birds and other vertebrate groups. First, mimetic species are smaller and socially subordinate to their model counterparts, yet these size differences are constrained such that visual deception must be feasible at distances germane to the players’ behavioral ecology. Second, the costs of mimicry and the value of contested resources cannot be exceedingly high for mimic and model species to coexist through time. Third, shared similarities in appearance are not attributed to homologous traits, in that the mimic and model are not closely related sister species. And fourth, both mimic and model are under natural selection to maintain or evade visual deception, respectively. As such, coevolutionary radiations may emerge if the evolution of mimicry precedes diversification in the model species. Here, we examine the evidence for ISDM and alternative explanations of phenotypic convergence in the Helmeted Woodpecker (Dryocopus galeatus), a little-known Atlantic Forest endemic whose systematic affinities remain unclear given its enigmatic combination of morphological and behavioral characters.

Initially described as Picus galeatus (Temminck, 1822), the Helmeted Woodpecker was soon transferred to
Dryocopus by Gray (1845), where it has generally remained, though not without comment. Short (1982) was apparently the first to recognize that \textit{D. galeatus} shares morphological characters with both \textit{Dryocopus} and \textit{Celeus}, commenting that the species is “beautifully intermediate” and could be placed in either genus. Specifically, he noted the weaker curved bill, exposed nostrils, cinnamon wing linings, and white upper tail coverts that are characteristic of \textit{Celeus}, whereas the uniform dark dorsal plumage, ventral barring, whitish neck stripe, unbarred wings, and fully red crest are traits shared with Neotropical \textit{Dryocopus}. Despite these morphological similarities to \textit{Celeus}, Short concluded that \textit{D. galeatus} is most likely sister to the \textit{D. schulzi} + \textit{D. lineatus} clade, to which it bears strong phenotypic resemblance (Figure 1), and is narrowly sympatric (Figure 2) with the latter taxon (Short 1982, Winkler et al. 1995). Here, we build upon a recent molecular phylogenetic analysis of \textit{Celeus} by incorporating multilocus sequence data from \textit{D. galeatus} and putative congeners to resolve its systematic position and provide a phylogenetic basis for examining plumage evolution.

**METHODS**

**Taxon Sampling and Sequencing**

We obtained molecular sequence data from 44 woodpecker specimens representing 31 species in 13 genera (Table 1). Taxon sampling was concentrated within the clade Malarpicini, including 5 of 7 \textit{Dryocopus} species, all currently recognized \textit{Celeus} species (Benz and Robbins 2011), and ≥1 species from each of the remaining Neotropical genera (Winkler and Christie 2002). Six outgroup taxa were selected from the Megapicini and...
Dendropicini on the basis of previous molecular phylogenetic studies of the Picidae (Webb and Moore 2005, Benz et al. 2006, Fuchs et al. 2007). Whole genomic DNA was extracted from muscle tissue using proteinase K digestion under manufacturer's protocols (Qiagen DNeasy tissue kit). We employed standard PCR amplification and Sanger sequencing methods to generate sequence data from 4 mitochondrial genes (NADH dehydrogenase subunits 2 and 3 [ND2 1,041 bp; ND3 351 bp] and ATP synthase subunits 6 and 8 [ATP6, 684 bp; ATP8, 168 bp]) and 2 nuclear loci (intron 7 of the β-fibrinogen gene [β-FIBI7, 913 bp] and a segment of the nonhistone chromosomal protein HMG-17 gene including exon 2 and adjacent mRNAs [HMGN2, 709 bp]). Sequence data were obtained from ND2, ND3, and HMGN2 for all 44 specimens, whereas sequencing effort for β-FIBI7 was limited to contemporary samples (n = 41) and that of ATP6-8 to species in the genera Celeus and Dryocopus (n = 27).

### Table 1. taxa sampled for this study.

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*Tissue sources: KU = University of Kansas Biodiversity Institute; LSUMNS = Louisiana State University Museum of Natural Science; UNAM = Museo de Zoología, Universidad Nacional Autónoma de México; USNM = United States National Museum of Natural History; FMNH = Field Museum of Natural History; LACM = Los Angeles County Museum of Natural History; UWBM = Burke Museum, University of Washington. Asterisk indicates museum specimens sequenced from toepad samples.*
Ancient DNA sequencing techniques were used to obtain complete ND2, ND3, ATP6-8, and HMGN2 sequence data from 2 museum specimens of D. galeatus collected in 1959 from Tobunas, Argentina (Table 1). For comprehensive details of our laboratory protocols, see Benz and Robbins (2011). GenBank accession numbers for all sequence data generated prior to this investigation can be found in Benz and Robbins (2011) and Benz et al. (2006).

**Phylogenetic Analysis**

We used Akaike’s Information Criterion (AIC) implemented in jModelTest version 2 (Guindon and Gascuel 2003, Darriba et al. 2012) to determine the best-fitting models of evolution for individual nuclear loci and the concatenated mtDNA dataset partitioned by codon position. A series of Bayesian phylogenetic analyses was conducted for each of these datasets in MrBayes version 3.2.1 (Ronquist et al. 2012) to assess potential conflict in phylogenetic signal among individual gene trees. We used a flat default prior distribution for parameter estimation, and mitochondrial data partitions were permitted to vary independently by unlinking all parameters except topology and branch length. Two independent analyses were run per locus for \(2 \times 10^7\) generations and sampled every 100 generations, resulting in a total of \(2 \times 10^7\) samples. Stationarity for each analysis was assessed by examining average standard deviation of split frequencies, plotting model-parameter posterior-probability densities in Tracer version 1.5 (Rambaut and Drummond 2007) and examining clade posterior probabilities across runs using the compare and slide functions in AWTY (Nylander et al. 2008). Trees that were sampled before the analysis reached stationarity were discarded as burn-in. These same run parameters were then used for a combined data analysis with sequences partitioned by nuclear loci and mitochondrial codon position.

Maximum-likelihood analyses of the individual loci and 5-partition data matrix were conducted in GARLI version 2.0 (Zwickl 2006) to provide alternative estimates of topology and node support. A total of 30 runs were conducted under default parameters to ensure that the optimal \(-\ln L\) solution had been reached, and topologies were selected after 10,000 generations with no significant improvement in \(-\ln L\) (improvement values set at 0.01, with a total improvement <0.05 compared to the last topology recovered). Node support was assessed using 500 nonparametric bootstrap replicates that were run with the above default parameters.

**Plumage Analysis**

We examined multiple museum specimens of D. galeatus \((n = 8)\) and candidate model species D. lineatus \((n = 14)\) and C. robustus \((n = 5)\) to assess overall plumage similarities and the potential for visual deception between these codistributed taxa. Representatives of each of the 5 recognized D. lineatus subspecies were examined, including 4 individuals of D. lineatus erythrops, the southernmost taxon in the lineatus complex and the only subspecies codistributed with D. galeatus. To examine the distribution of key plumage traits more broadly within the Malarpicini, we examined multiple specimens of all Dryocopus, Celeus, Colaptes, and Piculus species to determine whether D. galeatus plumage traits are novel within Celeus and its sister group Colaptes + Piculus. Given that several of these plumage characters involve subtly different feather tracts that are potentially nonhomologous across the picinæ, we chose to map whole-phenotype illustrations on the combined-data maximum-likelihood topology to illustrate the distribution of convergent similarities in appearance among relevant taxa.

**RESULTS**

**Sequence Attributes**

The concatenated sequence alignment contained 3,855 characters, 1,275 of which were variable and 897 parsimony informative (Table 2). Mitochondrial sequences appeared to be of genuine origin, in that stop codons were not observed in open reading frames, base composition was homogeneous across samples, and codon-specific substitution rates were consistent with known biases. Third-position substitutions accounted for 58.7% (527 bp)
of the total informative sequence variation and for 69.2% (392 bp) of the informative sites recovered within *Celeus*. By comparison, β-FIBI7 and HMGN2 sequences exhibited little genetic variation, containing just 60 (6.6%) and 85 (9.5%) informative substitutions within the full data matrix, respectively, of which 17 (1.9%) and 18 (2.0%) were recovered within *Celeus*. Informative indels were also rare within nuclear sequences; however, a 2-bp deletion shared by all members of *Celeus* and *D. galeatus* corroborate the phylogenetic results presented below (Figure 3). Both specimens of *D. galeatus* yielded nearly identical sequences with no conflict among overlapping amplicons, which suggests an absence of contaminant DNA. All new sequences generated for this investigation have been deposited in GenBank (accession nos. KT204492–KT204537).

**Phylogenetic Analysis**

Analyses of AIC values generated in jModelTest indicated that the general time-reversible substitution model GTR+I+Γ was most appropriate for the first and third codon positions, whereas the model HKY+Γ was selected for the more conservative second codon position, and a transversion model of evolution (TVM+Γ) was best suited for the HMGN2 and β-FIBI7 nuclear loci (Table 2). Phylogenetic analyses of the individual nuclear loci and the combined 4-gene mitochondrial dataset recovered similar results, differing primarily in the degree of intragenic resolution, which reflects the large disparity in rates of evolution and informative variation among these marker sets (Figure 3). Because few conflicts were observed among individual gene trees and none were statistically significant, our primary focus here is on the combined-data phylogenetic analyses, which form the basis of our discussion; however, we emphasize that both the mitochondrial and HMGN2 analyses strongly rejected the monophyly of *Celeus* as currently defined (Figure 3). Phylogenetic results of the β-FIBI7 analyses are provided in Figure 4 (*D. galeatus* was not sequenced for this locus).

Maximum-likelihood and Bayesian analyses of the 6-gene, 5-partition data matrix recovered concordant topologies with strong bootstrap and posterior probability support across all but 3 nodes among ingroup taxa (Figure 5). Monophyly of *Celeus* was unequivocally rejected in all analyses, with *D. galeatus* placed between the basal *C. torquatus* ‑ *C. loricatus* lineage (Clade III) and the remainder of *Celeus* diversity. Moderate levels of pairwise sequence divergence recovered between *D. galeatus* and other *Celeus* species (8.5–10.2%; ND2 uncorrected) suggest that the former has no close relatives and likely represents an early split within the genus. Although the phylogenetic position of *D. galeatus* received strong
statistical support in these analyses, weak support between Clade I and Clade II indicate uncertainty in this arrangement, which the nuclear gene tree analyses were also unable to resolve because of a lack of informative variation (Figures 3 and 4). This uncertainty had no effect on the position of Clade III, which was consistently recovered as the basal lineage within Celeus. The Colaptes + Piculus clade was placed sister to Celeus with moderate to strong support, followed by a paraphyletic Dryocopus lineage in which both Old World members of the genus are more closely related to Mulleripicus than to New World Dryocopus (Figure 5). Phylogenetic relationships among outgroup taxa were consistent with previous multilocus investigations (Benz et al. 2006, Fuchs et al. 2007), so these species were omitted from the final topology to facilitate phenotype mapping within the Malarpicini.

**Plumage Analysis**

Similarities in appearance between D. galeatus, D. lineatus, and C. robustus are primarily attributed to the fully red crest, pale venter with narrow black banding, and uniform black dorsum (assuming an at-rest posture) shared by each of these taxa. Examination of 128 representative study
skins encompassing 48 species in 5 genera confirmed that these traits are absent in all other Celeus species, as well as in the sister group Piculus + Colaptes, indicating a convergent evolutionary origin of these traits within D. galeatus (Figure 5). Although both candidate model species exhibit fine-scale differences in plumage pattern and coloration, D. galeatus more closely resembles D. l. erythrops in several key aspects. The white lateral neck stripes in D. l. erythrops narrow above the malar and transition to pale orange-brown as they approach the nares, a pattern that is mirrored in D. galeatus but with little or no stripe definition above the malar and darker cinnamon at the nares (Figure 1). These traits are absent in male C. robustus, which, with the exception of white and black ear coverts, have a fully red head, throat, and neck. Although female C. robustus exhibit a more extensive white and black streak that extends from the ear to the base of the bill, these prominent plumage differences

FIGURE 5. Phylogenetic relationships within the woodpecker clade Malarpicini, inferred from the combined sequence alignment (ND2, ND3, ATP6, ATP8, HMGN2, and β-FIB17). Bayesian posterior probabilities and maximum-likelihood bootstrap support values are indicated above and below each node, respectively. Representative images from each of the 3 primary Celeus clades illustrate the phenotypic disparity between Dryocopus galeatus and its true congeners. Illustrations a–k are reproduced with the permission of Lynx Edicions.
strongly contrast with the head and neck plumage of *D. galeatus* and *D. l. erythrops*. Extensive white scapular patches are characteristic of *D. lineatus* populations from northern Mexico to southern Brazil, yet most populations of *D. l. erythrops* have fully black scapulars, which is also the case in *D. galeatus*. Notable phenotypic differences between *D. galeatus* and *D. l. erythrops* include a pale chin with fine black striping that transitions to an extensive and uniform black upper chest in the latter, which clearly contrasts with the cinnamon chin and small black patch restricted to the throat in the former. Nonetheless, these differences are not readily apparent from lateral profiles (Figure 1). Further distinctions include the ivory bill, dark brown irides, and pale rump, which are characteristic of *D. galeatus* and contrast with the dark gray maxilla, white irides, and black rump in *D. l. erythrops*.

**DISCUSSION**

In his review of avian evolution and homodynamy in morphologically uniform groups, Bock (1963) examined 3 pairs of woodpecker genera (*Dryocopus*–*Campephilus*, *Gecinulus*–*Blythipicus*, and *Dinopium*–*Chrysocolaptes*) that share highly similar plumage characteristics but exhibit key morphological differences in the foot, tail, and bill—adaptive features that are closely linked to taxon-specific foraging strategies. He concluded that these plumage similarities were evidence of recent shared ancestry, in that “the complexity of these color patterns precludes any reasonable possibility of their arising independently in three pairs of genera; no known selection forces could explain such a pattern of convergence.” Over the past decade, molecular phylogenetic analyses have confirmed that the inverse is the case, documenting convergent evolution of phenotypic traits in 11 picid genera, and now our data provide yet another example in which plumage convergence in the Helmeted Woodpecker has confounded phylogenetic relationships in the Malopigini (Webb and Moore 2005, Weibel and Moore 2005, Benz et al. 2006, Moore et al. 2006, Fuchs et al. 2008). Although some uncertainty remains with respect to branching patterns among primary *Celeus* clades, our mitochondrial and nuclear data strongly support inclusion of *D. galeatus* within the genus (Figures 3 and 5). These results greatly expand the phenotypic diversity of *Celeus* and demonstrate the independent evolutionary origin of plumage similarities between *D. galeatus* and its larger, socially dominant ecological competitors *D. lineatus* and *C. robustus*, both of which are sympatric throughout much of the Helmeted Woodpecker’s distribution (Figure 2). By all other accounts, the Helmeted Woodpecker is morphologically highly similar to other *Celeus* species. Its culmen is noticeably curved (similar in shape to that of *C. flavescens*), whereas all true *Dryocopus* exhibit distinctly straight and chisel-shaped bills that are more robust (wider at the base) and typically darker in coloration. As in all other *Celeus* species, the nares of *D. galeatus* are fully exposed and lack the stiff feathers that partially or fully cover the nares of *Dryocopus* species. The dark brown irides of the Helmeted Woodpecker are also characteristic of *Celeus*, whereas all but 1 species of *Dryocopus* (*D. schulzi*) have pale whites irides. An absence of fluid-preserved anatomical specimens of *D. galeatus* precludes, at this time, further morphological comparison and diagnosis of the internal synapomorphies detailed by Goode (1972); nonetheless, it appears that previous workers have failed to see the true evolutionary relationships of *D. galeatus*, largely because of the historical bias toward using labile plumage traits to infer phylogenetic relationships while disregarding other more conservative and potentially informative traits. We suspect that this striking case of plumage convergence likely constitutes a form of interspecific mimicry whereby *D. galeatus* receives social advantages, including reduced competitive interference and enhanced access to foraging sites, by visually deceiving one or more of its ecological competitors. Although extensive field studies will be required to confirm whether deception is indeed taking place and to what extent ISDM or alternative evolutionary mechanisms are driving these convergent similarities in appearance, the limited behavioral evidence presently available appears to be consistent with the ISDM hypothesis, which we discuss below.

**Nonmimetic Evolutionary Convergence**

Environmental adaptation based on principles of natural selection is frequently invoked to explain general phenotypic trends and geographic variation in plumage coloration within birds (Mayr 1963, Zink and Remsen 1986, Hill and McGraw 2006). By contrast, examples of environmental selective pressures driving comprehensive phenotypic convergence of elaborate plumage patterns are relatively uncommon and are typically poorly substantiated. Perhaps the best example of the latter occurs between *Macronyx croceus* (Motacillidae) of open African savannas and the meadowlark (Icteridae) species complex *Sturnella* spp., which inhabit grassland environments throughout much of the New World. The brown-streaked dorsal plumage, bright yellow venter, and black pectoral band shared by these allopatic species suggest that there is potential for wholesale phenotypic convergence among taxa exposed to similar environmental selective pressures. Further examples of plumage convergence frequently attributed to environmental adaptation are seen in several oceanic birds, including *Alle alle* and various *Pelecanoides* species, whose black dorsal plumage and white venter are likely related to similar selective pressures associated with foraging in open-ocean environments. The proposition.
that environmental adaptation may be driving the present case of plumage convergence among Atlantic Forest woodpeckers is clearly falsified by the fact that *D. lineatus* ranges from northern Mexico to eastern Argentina (Figure 2), occupying a wide diversity of habitats, including mangroves, thorn-scrub, open-gallery forest, dense rainforest, and pine–oak environments, which collectively range from 0 to 2,100 m a.s.l. (Short 1982). That any aspect of this species’ visual appearance could be under similar environmental selective pressures in such diverse habitat types is highly unlikely. Moreover, visually similar species in both *Dryocopus* and *Campephilus* further extend the distribution of these phenotypic traits.

Natural and sexual selection processes acting on labile or highly modular phenotypic traits may also foster nonmimetic convergent similarities in appearance (Endler and Théry 1996, Hill and McGraw 2006). Analyses of plumage evolution in New World orioles (*Icterus*) revealed evidence of convergence or reversals in 42 of 44 plumage characters, with repeated evolutionary origins of broad phenotypic similarities in 3 distinct clades (Omland and Lanyon 2000). Although several instances of plumage convergence within *Icterus* involve largely allopatric species, thereby precluding mimicry or socio-ecological explanations for similarities in appearance, such examples of parallel selection in allopatry appear to be uncommon in birds. Furthermore, examples of convergent plumage evolution among allopatric species typically involve rather coarse plumage details and lack the fine-scale, comprehensive phenotypic similarities necessary for successful visual mimicry. If similar natural or sexual selection processes are responsible for convergent plumage similarities in the Helmeted Woodpecker, why are these traits wholly absent in other *Celeus* species as well as *Colapates* and *Piculus*? Likewise, these hypotheses do not explain why *D. galeatus* more closely resembles sympatric populations of *D. l. erythrops* rather than other congenerics or allopatric members of the *lineatus* complex.

**Interspecific Visual Mimicry**

Aposematic forms of mimicry (e.g., Batesian and Müllerian) arise via natural selection processes that typically involve a third-party observer in addition to the model and mimetic taxa (Ruxton et al. 2004). In the absence of aposematic signaling for predation avoidance, interspecific visual mimicry is thought to confer social advantages to species in close ecological competition by enabling mimetic taxa to avoid aggression through deception and increase their foraging opportunities (Wallace 1869, Diamond 1982, Rainey and Grether 2007, Prum and Samuelson 2012). Diamond (1982) hypothesized that competitive mimicry reduces the frequency of attack by larger model species, thereby altering the social hierarchy at highly contested foraging sites, countering size-mediated social dynamics; however, he remained skeptical of whether selection pressures associated with the risks of aggression were sufficient to prevent models attacking the smaller mimetic species. As such, Diamond did not rule out the possible role of a third-party observer in promoting and maintaining competitive mimicry between *Philemon* and *Oriolus* species.

More recently, Prum and Samuelson (2012) used evolutionary game-theory modeling to confirm that intraspecific attacks are indeed constrained by the costs of aggression, thereby fostering social opportunities for interspecific competitive mimicry purely within the context of a 2-party system. Fitness dynamics derived from these game-theory models predict the evolution of ISDM when associated costs of mimicry are low, the background fitness of the mimetic species is greater than that of the model, and the values of contested resources are neither exceedingly high nor low. The ISDM hypothesis also predicts that selection on the dominant model species will favor evolution of divergent phenotypic traits to reduce the efficacy of mimicry and its associated costs. In time, these counterselective forces may lead to coevolutionary radiations between model and mimetic taxa, as seen in the *Philemon* and *Oriolus* example or perhaps between *Dryocopus* and *Campephilus* (Diamond 1982, Prum 2014). Lastly, size differences between mimic and model are constrained in ISDM such that visual deception must be feasible at distances relevant to their behavioral ecology. The latter prediction is readily testable and should provide critical insight into the functional significance of phenotypic convergence in birds, given that neither environmentally based nonmimetic hypotheses nor the earlier mimicry hypotheses of Moyáñan (1968) and Cody (1969) predict explicit size relationships between model and mimic.

In his recent review of avian visual mimicry, Prum (2014) proposed 50 phylogenetically independent examples of ISDM from 30 families. The average mimic body mass was 55.7% of the putative model counterpart, and a linear regression of body mass between subordinate mimic and dominant model revealed a strong positive correlation with a slope of 0.5684 and *R*² value of 0.83. This close association between asymmetry in body size and similarity in appearance strongly suggest that these convergent signals are evolving in the context of competitive mimicry to facilitate interspecific deception of both the mimic’s identity and its body size. By simply appearing to be large rather than physically evolving larger body size to legitimately dominate an ecological competitor, the mimic species may incur advantages in physiological efficiencies while simultaneously having access to a wider diversity of ecological resources, which translates to greater adaptability and evolutionary persistence. The selective pressures for such advantages may be especially
acute in groups that exhibit highly specialized foraging strategies, which may explain the high prevalence of phenotypic convergence in the picinae.

**Visual Mimicry in the Helmeted Woodpecker**

Interspecific mimicry in *D. galeatus* was first proposed by Willis (1989), who identified the larger Robust Woodpecker (*Campephilus robustus*) as a possible dominant model (Figure 1), given that the two occasionally forage together in mixed-species flocks. In light of our phylogenetic results, it's plausible that *galeatus* is a mimic of both *C. robustus* and *D. lineatus*; however, the latter bears greater similarity to *galeatus*, given its conspicuous white neck stripes and darker lores. Moreover, we underscore the fact that *galeatus* is sympatric with southern populations of *D. l. erythrops*, both of which lack the white scapular patches that are otherwise characteristic of the broadly distributed *lineatus* complex (Winkler and Christie 2002). At ~28 cm in length, the Helmeted Woodpecker is 22–25% smaller than *D. l. erythrops* (36 cm) or *C. robustus* (37 cm), respectively, but weighs less than half of either model taxon. This disparity in size and weight undoubtedly confers a substantial physical advantage to the larger *Dryocopus* and *Campephilus* models, given that even small differences (<10%) in body mass can lead to greater success in competitive interference for numerous avian groups (Ford 1979, Maurer 1984, Millikan et al. 1985, Alatalo and Moreno 1987, Robinson and Terborgh 1995).

Critically, the size differences observed between *galeatus* and either model species appear to be consistent with the ecological and psychophysical constraints required for visual deception as outlined by Prum (2014). Given that the difference in distance between 2 objects sharing the same visual angle scales linearly with difference in size, mistaking a mimic species for a conspecific dominant model would require overestimating the true distance by just 32%, which is well within the ecological context in which these species regularly encounter one another. Although avian visual acuity varies substantially among taxonomic groups, ophthalmological research and psycho-physical data suggest that in nonraptorial birds, visual deception of this nature is possible at distances ≥3 m (Hodos 1993, Prum 2014).

Knowledge of the Helmeted Woodpecker’s ecology and foraging behavior remains extremely limited in comparison with that of other Neotropical picids; however, *galeatus* appears to be an ant specialist (*Crematogaster* sp.) that regularly consumes small fruits such as *Alchornea sidifolia* berries (Santos 2008, Lammertink et al. 2012, K. J. Zimmer personal observation). This secretive species primarily forages at middle levels on interior branches, quietly probing rotting wood, which is consistent with *Celeus* foraging behavior (Short 1982, Winkler and Christie 2002). All 12 species currently recognized within *Celeus* are documented ant or termite specialists that regularly consume fruits and rarely exhibit strong excavating or bark-scaling behavior characteristic of *Dryocopus* and *Campephilus*. Both *C. robustus* and *D. lineatus* consume fruits, and the latter regularly forages on ants, including *Crematogaster*, *Azteca*, and *Camponotus* species. The diet of these larger picids differs from that of *galeatus* in that both *Dryocopus* and *Campephilus* species use their powerful bills to excavate beetles and their larvae from deep within rotten to semirotten substrates. Given that rotting trees of appropriate age and decomposition are often in limited supply within forest environments, competition for suitable foraging substrates, rather than direct competition for a particular food species, may be responsible for the evolution of competitive mimicry between *galeatus* and either dominant model species. Although southern populations of *D. l. erythrops* appear to forage within higher strata than either *galeatus* or *C. robustus*, the competition for foraging sites may encompass an entire tree, given that most *Dryocopus* and *Campephilus* species generally do not tolerate unfamiliar conspecifics in the vicinity of an active feeding site. Despite an absence of detailed knowledge about the socio-ecological interactions within this trio of woodpecker taxa, evidence of interspecific mimicry is most consistent with the ISDM hypothesis presented by Prum and Samuelson (2012), given that (1) woodpeckers are not known to sequester toxins in their skin or feathers and (2) Cody’s (1969) suggestion that phenotypic convergence promotes enhanced interspecific territoriality is disproved by the fact that both *galeatus* and *C. robustus* occasionally attend mixed-species flocks (Willis 1989). Moreover, Cody’s hypothesis does not take into account the strong asymmetry in body size associated with cases of competitive mimicry, casting doubt on the evolutionary stability of mutual exclusion. Moynihan’s (1968) hypothesis is not applicable in the present case because neither mimic nor model features facultative mixed-flock attendants. Likewise, little support for this hypothesis is seen in other proposed examples of avian competitive mimicry (Diamond 1982, Prum 2014).

Behavioral investigations examining the socio-ecological circumstances associated with cases of avian phenotypic convergence will be required to confirm the prevalence of ISDM versus traditional 3-party mechanisms of visual mimicry. Although Prum and Samuleson (2012) demonstrated that the evolution and persistence of competitive mimicry is possible exclusively within a 2-party system, it seems plausible that ISDM may operate synergistically with third-party deception mechanisms, because the potential pool of nonmodel ecological competitors is much larger. Given that most birds communicate a broad array of intraspecific and interspecific information through vocalizations, future field studies examining potential cases
of competitive mimicry must also take into account the mimic’s vocal behavior, particularly in the presence of models and other heterospecific competitors, because any discontinuity between morphological and behavioral mimicry would likely preclude the possibility of deception. As such, we predict that mimic taxa will vocalize less in the presence of ecological competitors than their nonmimetic congeners. Although little is known of the Helmeted Woodpecker’s vocal behavior, it appears to call less frequently than other members of Celeus, which may account for the dearth of visual sightings and the possibility of extinction reported by Short (1982).

Conservation Status and Taxonomic Implications
The Helmeted Woodpecker inhabits semideciduous and mixed-forest environments from São Paulo, Paraná, and Santa Catarina in southeastern Brazil, west to eastern Paraguay, and south to Misiones in extreme northeastern Argentina (Short 1982, Collar et al. 1992, Hayes 1995, Winkler and Christie 2002). This little-known species has undergone dramatic population declines and vanished from much of its former distribution in the second half of the 20th century as a result of extensive regional deforestation (Short 1982, Galindo-Leal and de Gusmão Câmera 2003, Santos 2008). Consequently, it is currently listed as “vulnerable” by BirdLife International and considered to be among the rarest of Neotropical woodpeckers (Lammertink et al. 2012, BirdLife International 2014). Although new details of its ecology and natural history are slowly emerging, the conservation status of D. galeatus remains unclear and deserves careful examination, given regional trends in habitat loss.

In light of our phylogenetic data, described morphological differences (Short 1982), distinct vocalizations (galeatus has vocalizations similar to those of Celeus torquatus and C. flavus; XC61093, Xeno-Canto, http://www.xeno-canto.org), and mechanical sound production (drumming of galeatus is Celeus-like; XC24502, XC17067, Xeno-Canto), the Helmeted Woodpecker clearly requires reclassification as Celeus galeatus.

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