Origin of elevational replacements in a clade of nearly flightless birds – most diversity in tropical mountains accumulates via secondary contact following allopatric speciation

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Abstract

Tropical mountains are biodiversity hotspots. In particular, mountains in the Neotropics exhibit remarkable beta diversity reflecting species turnover along gradients of elevation. Elevational replacements of species are known since early surveys of the tropics, but data on how such replacements arise are scarce, limiting our understanding of mechanisms underlying patterns of diversity. We employed a phylogenetic framework to evaluate hypotheses accounting for the origin of elevational replacements in the genus *Scytalopus* (Rhinocryptidae), a speciose clade of passerine birds with limited dispersal abilities occurring broadly in the Neotropical montane region. We found that species of *Scytalopus* have relatively narrow elevational ranges, closely related species resemble each other in elevational distributions, and most species replacing each other along elevational gradients are distantly related to each other. Although we cannot reject the hypothesis that a few elevational replacements may reflect parapatric speciation along mountain slopes, we conclude that speciation in *Scytalopus* occurs predominantly in allopatry within elevational zones, with most elevational replacements resulting from secondary contact of formerly allopatric species. Our work suggests that accumulation of species diversity in montane environments reflects colonization processes even in dispersal-limited animals.

Introduction

Species turnover along gradients of elevation is a salient pattern in tropical biogeography. Ever since pioneering work by Francisco José de Caldas and Alexander von Humboldt on plant geography, naturalists noted that many species occur over narrow ranges of elevation and replace each other along mountain slopes (Nieto, 2006; von Humboldt & Bonpland, 2009). Elevational replacements of closely related species are prevalent in the tropics (Terborgh, 1971; Diamond, 1973; Wake & Lynch, 1976), where organisms likely have narrower physiological tolerances than in temperate zones (Janzen, 1967; McCain, 2009). Marked changes in species assemblages with elevation (e.g. of plants, invertebrates and vertebrates; Patterson et al., 1998; Kessler, 2001; Jankowski et al., 2013b; García-Robledo et al., 2016; Gill et al., 2016; Badgley et al., 2018) thus result in tropical mountains being hotspots of beta diversity (Melo et al., 2009; Fjeldså et al., 2012). Therefore, knowledge about evolutionary and ecological mechanisms involved in the origin of elevational replacements is central to understanding major patterns in the distribution of life (Janzen, 1967; Huey, 1978).

Abutting species distributions along elevational gradients may reflect either (1) parapatric ecological speciation leading to divergence of a formerly widespread species into two or more daughter species with restricted ranges, or (2) secondary contact following range expansions of species originating in allopatry (Endler, 1982; Hua, 2016). Although one should exercise caution when making inferences about the geographic context of speciation based on current geographic distributions (Losos & Glor, 2003), these alternative hypotheses are, in principle, amenable to testing by means of phylogenetic analyses: parapatric divergence predicts that species replacing each other with elevation are sister to each other, whereas secondary contact predicts they are not (Patton & Smith, 1992; Moritz et al., 2000). Evidence from a few studies on animals indicates that most speciation in the montane Neotropics occurs in allopatry and that species replacing each other along elevational gradients are not each other's closest relatives (Patton & Smith, 1992; Cadena et al., 2012; Caro et al., 2013). Therefore, elevational replacements may primarily reflect secondary contact but more studies are necessary to confirm this pattern.

As evidenced by some of the first large-scale surveys of the geographic and ecological distributions of species (Chapman, 1917; Todd & Carriker, 1922; Chapman, 1926), most birds living in montane areas of the Neotropics have narrow elevational ranges (Jankowski *et al.*, 2013a; but see Gadek *et al.*, 2018 for exceptions). For example, median ranges of

species across their geographic distributions span only ca. 1100 m in three families of Neotropical birds (Figure 1). Narrow elevational ranges of individual species are often coupled with spatial segregation with ecologically similar species along mountain slopes. For instance, a series of landmark studies in the Peruvian Andes documented multiple cases of pairs of congeneric species of birds replacing each other with elevation as well as scenarios where up to 4-5 congeners occur successively along mountain slopes with minimum overlap (Terborgh, 1971; Terborgh & Weske, 1975; Terborgh, 1977).

Tapaculos in the genus *Scytalopus* (Rhinocryptidae) are small passerine birds with poor dispersal abilities ranging broadly in the Neotropical montane region. *Scytalopus* are mouse-like birds which forage walking or hopping on or near the ground in dense vegetation; they are unable to engage in long, powered flights because they have small and rounded wings and unfused clavicles (Figure 2; Krabbe & Schulenberg, 2003; Maurício *et al.*, 2008). *Scytalopus* tapaculos avoid highly lit open areas and are rarely found far from vegetation cover except in barren high-elevation environments. Local diversity of *Scytalopus* is typically low, yet multiple species may be found in different habitats in a given landscape, being prime examples of patterns of elevational replacement (Krabbe & Schulenberg, 1997, 2003). For example, on a single morning walking trails upslope in forests along the Cerro de Montezuma on the Pacific-facing slope of the Andes of Colombia, birdwatchers may successively encounter *S. chocoensis*, *S. alvarezlopezi*, *S. vicinior*, *S. spillmanni*, and *S. latrans*; if they visit paramos at higher elevations in the region, then they may also find *S. canus* (Stiles *et al.*, 2017; see below). Elevational replacements in the genus are typically sharp. Species seldom co-occur at the same elevations and when they do so, they often segregate by habitat. Current taxonomy recognizes 44 species of *Scytalopus*, but this is most certainly an underestimate given marked genetic structure within species, geographic variation in vocalizations, and the potential for discovery of new taxa in unexplored regions (Cadena *et al.*, 2019b).

Elevational replacements in Scytalopus are somewhat paradoxical given hypotheses posed to account for such patterns. First, given morphological and life-history traits making these birds notably poor dispersers one would expect that longdistance dispersal or range expansions would hardly bring multiple species together in a given slope, especially because mountain regions are separated by geographic barriers associated with population isolation and diversification in other birds (Hazzi et al., 2018). Alternatively, given what we know about the geographic context of speciation in birds in general (Phillimore et al., 2008; Price, 2008) and in the Andes in particular (García-Moreno & Fieldså, 2000; Caro et al., 2013), parapatric divergence of multiple species along elevational gradients in several geographic regions would also appear unlikely. A preliminary analysis of diversification in Scytalopus focusing on few taxa from eastern Ecuador and Peru suggested secondary contact is a more likely explanation for elevational replacements (Arctander & Fieldså, 1994; Roy et al., 1997). However, because the phylogeny employed for analyses may have been problematic (Arctander, 1995), taxonomic and geographic coverage were limited, and our understanding of geographic ranges, species limits and diversity in the group has changed substantially (Krabbe & Schulenberg, 1997; Cadena et al., 2019b), revisiting the questions is warranted. We used a comprehensive molecular phylogeny and data on geographic and elevational distributions of species to describe the mode of speciation and thereby examine the origin of elevational replacements in Scytalopus. Understanding ecological and physiological mechanisms maintaining patterns of elevational segregation has been the focus of multiple studies (reviewed by Jankowski et al., 2013a), yet analyses of the origins of elevational replacements remain scarce (Cadena, 2007; Freeman, 2015; Cadena et al., 2019a).

Methods

A recent study reconstructed molecular phylogenies of Scytalopus employing sequences of mitochondrial DNA (mtDNA), and of 80 exons and > 1800 regions flanking ultraconserved elements in the nuclear genome (Cadena et~al., 2019b). We used the mtDNA ND2 data set from that study, which had denser taxonomic and geographic sampling, as the basis for our analyses; phylogenies inferred with this data set were largely congruent with those based on nuclear data. We inferred a gene tree using Beast 2.4.8 (Bouckaert et~al., 2014) for an alignment of 90 sequences, including 83 Scytalopus and seven outgroups. We applied a relaxed uncorrelated clock (mean=0.0125, SD=0.15; Smith & Klicka, 2010) and a Yule speciation tree prior. We ran chains for 100 million generations and discarded the initial 50% as burn-in.

We were able to gather elevational distribution data for 57 taxa of Scytalopus; these represent all named species as well as several distinct populations, which may be undescribed species given genetic divergence, vocal variation, or geographic distributions (see Cadena et al., 2019b). We obtained information on elevational ranges from taxonomic descriptions (Whitney, 1994; Krabbe & Schulenberg, 1997; Cuervo et al., 2005; Krabbe et al., 2005; Maurício, 2005; Bornschein et al., 2007; Donegan & Avendaño-C., 2008; Krabbe & Cadena, 2010; Whitney et al., 2010; Donegan et al., 2013; Hosner et al., 2013; Maurício et al., 2014; Avendaño et al., 2015; Avendaño & Donegan, 2015; Stiles et al., 2017), regional handbooks (Fjeldså & Krabbe, 1990; Ridgely & Greenfield, 2001; Hilty, 2003; Schulenberg et al., 2007; Herzog et al., 2016), and the Handbook of the Birds of the World (del Hoyo et al., 2018). Additionally, for 11 taxa we defined elevational distributions based on expert knowledge (A M. Cuervo, N. K. Krabbe, D. F. Lane, and T. S. Schulenberg, unpubl. data); this was particularly useful for unnamed populations differing phenotypically or genetically from others. We performed all analyses described below using all 57 taxa and separately for a data set including only 47 of them which represent all currently recognized species (American Ornithologists Union 1998; Stiles et al., 2017; Remsen et al., 2018) plus three unnamed species from the Andes of Peru (an unnamed form referred to S. altirostris mentioned by Cadena et al., 2019b was not considered). To conduct analyses, we trimmed the phylogeny constructed with the complete data set (i.e., 90 terminals) to include only the 57 or 47 taxa considered.

To describe changes in elevational distributions over the history of Scytalopus, we mapped the midpoint of the elevation range of each taxon in the phylogeny using the contMap function of phytools, which estimates character states at nodes and along branches using a maximum-likelihood approach (Revell, 2012). Additionally, we calculated Pagel's \(\lambda\) (Pagel, 1999) as a measure of phylogenetic signal of midpoint elevation and accounted for phylogenetic uncertainty by calculating this statistic across the final 1,001 trees in the posterior distribution using phytools. Using likelihood-ratio tests, we also tested whether Pagel's λ in each tree was significantly different from 0 (i.e. no phylogenetic signal) and 1 (i.e. the value expected under pure Brownian motion). Phylogenetic signal is often interpreted in terms of the degree of conservatism or lability of a trait (e.g. Blomberg et al., 2003), but one must exercise caution when making inferences about factors underlying patterns because different evolutionary processes may produce similar values of phylogenetic signal (Revell et al., 2008). Nonetheless, high phylogenetic signal can be interpreted as a strong tendency of closely related species to resemble each other in a given trait (Revell et al., 2008). Therefore, high phylogenetic signal in elevational ranges could indicate close resemblance between close relatives in such ranges. In addition to examining the midpoint of elevational ranges of species, we also conducted analyses based on minimum and maximum elevation, obtaining qualitatively similar results. Although elevational ranges are not organismal traits per se, they do reflect heritable phenotypes allowing organisms to persist over a given range of conditions, which justifies our study of the evolution of such ranges in a phylogenetic framework (Cadena, 2007).

To evaluate whether sister taxa have similar or contrasting elevational distributions as predicted by allopatric and parapatric speciation, respectively (Patton & Smith, 1992), we identified sister taxa across the 1,001 final trees in the posterior distribution. We then calculated the elevational overlap of pairs of sister taxa by dividing the amount of overlap by the elevational range of the taxon with the narrowest range (Kozak & Wiens, 2007; Cadena et al., 2012). A value of 1 indicates that either ranges are exactly the same or that the narrower range is entirely contained in the wider range; a value of 0 indicates that elevational distributions do not overlap. The number of pairs of sister taxa employed for analyses varied across trees in the posterior from 18 to 23 (median = 20.3, total across trees = 51 pairs) in the 57-tip data set, and from 12 to 18 (median = 15.1, total across trees = 52 pairs) in the 47-tip data set. These analyses were restricted to sister taxa representing terminal branches in the phylogeny (i.e., we did not employ ancestral state reconstructions to compare elevational ranges in cases when a species representing a long terminal branch was sister to a clade formed by \geq 2 species).

Finally, we graphically examined the phylogenetic affinities of species of *Scytalopus* replacing each other along elevational gradients in four regions of South America to examine whether such replacements more likely reflect secondary contact or parapatric speciation along mountain slopes. These regions were: (A) the Sierra Nevada de Santa Marta in northern Colombia, (B) the Pacific slope of the Western Cordillera of Colombia, (C) Zamora-Chinchipe Province on the Amazonian slope of the Andes of Ecuador, and (D) the Río Satipo Valley in Junín Department, eastern Andean slope of Peru. All analyses were conducted and figures plotted in the R programming environment (R Core Team 2018).

Results

Scytalopus tapaculos jointly occupy a wide range of elevations in the Neotropics, from sea level up to 4,600 m in the high Andes (Table 1, Figure 3). Elevational ranges vary substantially among species, from very narrow (200 m) to quite broad (3,500 m), yet most species occupy only a relatively small fraction of the elevational gradients in which they occur: mean ranges were 1046.4 m (SD= 614.6 m) in the 57-taxa data set and 1124.3 m (SD= 633.5 m) in the 47-taxa data set. Given the geographic setting where species of Scytalopus occur, where mountains reach very high altitudes and habitats for birds may extend over several thousand meters (e.g., Figure 1), the elevational ranges of species are generally rather narrow. Overall, taxa within main clades of Scytalopus have roughly similar elevational distributions. For example, most species in the Brazilian clade (Figure 3, clade A) occur at low to mid elevations. In contrast, most species in its sister Andean clade (Figure 3, clade B) occur exclusively at high elevations; exceptions include S. fuscus, found in lowlands, and S. magellanicus, with the widest range in the genus (0 to 3,500 m). Species from the tropical Andes and Central America (Figure 3, clade C) show wide variation in elevational distributions, but species within subclades in the region tend to have similar ranges. For example, species in clades D, G and I all occur at high elevations in the tropical Andes except for S. femoralis, S. micropterus and S. caracae, which inhabit lower elevations than their close relatives (Figure 3). All taxa in clades E, F and H occur at mid elevations, with some ranging to lower elevations (Figure 3).

Phylogenetic signal for the midpoint of the elevational range of species measured by Pagel's λ , which differed significantly from 0 across all trees and for both data sets (p<0.001), was relatively high (mean=0.89 for the 57-taxon data set and 0.86 for the 47-taxon data set; Figure 3). However, in most trees (75% in the 57-taxon data set and 61% in the 47-taxon data set) phylogenetic signal was also significantly different from 1 (p<0.05), implying that elevational ranges are more divergent than expected given evolution under pure Brownian motion. These results indicate that closely related Scytalopus tend to strongly resemble each other in the midpoint of their elevation ranges, but differences among species cannot be fully accounted for by time since their divergence. The minimum and maximum elevations in the ranges of species also had significant phylogenetic signal; estimates of Pagel's λ for maximum elevation were very similar to those we obtained for midpoint elevation, while those for minimum elevation were slightly lower yet qualitatively similar (data not shown).

Most sister taxa in Scytalopus have similar elevational distributions (Figure 4). Mean overlap in elevational ranges between sister taxa was slightly higher in the 57-taxon data set (pooled mean across the 1,001 trees=0.72, SD= 0.34) than in the 47-taxon data set (mean=0.63, SD=0.37). Regardless of the data set employed for analyses, half or more pairs of sister taxa overlapped substantially in elevational ranges (overlap >0.8) while less than a quarter of pairs of sister taxa had distinct elevational ranges (overlap <0.2; Figure 4). Furthermore, the majority of sister taxa identified across the 1,001 trees (7 pairs out of 8, or 13 pairs out of 14 depending on the data set) showing overlap <0.2 do not occur on the same gradient, i.e. they are allopatric. These results suggest speciation in Scytalopus occurs predominantly within elevational zones and not in parapatry along mountain slopes. The only possible exception to this pattern is divergence between S. acutirostris and S. femoralis, which may or may not be sister species, but are close relatives replacing each other along the Amazonian slope of the Central Andes of Peru (see below). We note that our analyses included data for at most 57 taxa yet the true number of species in Scytalopus is likely higher. Because most of the lineages which we did not consider are closely allied to other lineages with similar elevational ranges (e.g. groups within S. atratus or S. parvirostris; Cadena et al. 2019b) we believe that if there is any bias in our results it would be in the direction of underestimating the true overlap in elevational ranges of close relatives. In other words, greater taxonomic coverage would likely reinforce our conclusion that sister taxa have similar elevational distributions.

The majority of Scytalopus replacing each other along the elevation gradients we examined are distant relatives (Figure 5). The two species endemic to the Sierra Nevada de Santa Marta belong to distinct clades last sharing a common ancestor ca. 8 million years ago (clades D and H in Figure 2; Figure 5). Likewise, the four species occurring in forests on the Pacific slope of the Western Andes of Colombia belong to four different clades (clades D, E, F and I in Figure 3), whereas a fifth species (S. canus) occurring in paramo habitats above treeline belongs to yet another clade (clade B). The species found above and below S. alvarezlopezi in the region (S. vicinior and S. chocoensis, respectively; Figure 5) belong to clade E, but they are not sister to each other. The five species replacing each other along the Amazonian

slope of the Andes in Zamora-Chinchipe, Ecuador, belong to four distinct clades (B, D, H and I in Figure 3; Figure 5). Two of the species with parapatric distributions in this gradient (S. latrans and S. micropterus) are closely related, but the posterior probability of the hypothesis that they are sisters is only 0.43. Finally, in contrast to patterns observed in other regions, four of the six species replacing each other with elevation the Río Satipo Valley in Junín, Peru, belong to clade I (Figure 3; Figure 5). Moreover, three of these species (S. femoralis, S. gettyae and S. acutirostris) belong to a group nested within clade I including several closely allied taxa with shallow divergence in mtDNA (Figure 3; Figure 5). The remaining two species found in this gradient belong to clade B (an unidentified taxon; Hosner et al., 2013) and clade H (S. atratus). Unpublished evidence indicates a seventh species (S. aff. parvirostris) belonging to clade G also occurs in the Río Satipo Valley (N. Krabbe, pers. comm.) but we did not include it in Figure 5 because information on its elevational range in the area is lacking. The most recent common ancestor of species occurring in elevational gradients in western Colombia, in eastern Ecuador, and in eastern Peru is the most recent common ancestor of all Scytalopus, which existed ca. 9.8 million years ago (Figure 5). Taken together, the above data indicate that most species in the genus replacing each other along gradients of elevation in South America did not evolve in parapatry in situ, but rather met in each gradient following divergence in allopatry. However, the possibility of parapatric speciation remains plausible for some of the species replacing each other with elevation on the eastern slope of the Andes.

Discussion

Many species of birds and other organisms have restricted elevational distributions, particularly in the tropics. This results in biodiversity patterns observable globally (e.g., tropical mountains are hotspots of species turnover in space because species replace each other with elevation; Fjeldså et al., 2012) and regionally (e.g., diversity may peak at mid elevations or decline with elevation in a given mountain; Quintero & Jetz, 2018). We probed into evolutionary processes resulting in replacements of species along mountain slopes by examining the elevational ranges of species in the context of a phylogeny of Scytalopus tapaculos, a speciose clade of Neotropical montane birds in which elevational replacements of species are commonplace. We found that (1) elevational ranges of species of Scytalopus are relatively narrow given the high mountains where they live, (2) closely related species in the genus usually resemble each other in elevational distributions, (3) most pairs of sister taxa have largely overlapping elevational ranges, and (4) species coexisting regionally with elevational segregation on mountains are very often –but not always– distantly related to each other. Our study thus suggests that speciation in Scytalopus occurs predominantly in allopatry within elevational zones, and that elevational replacements typically result from secondary contact of formerly allopatric species rather than from primary divergence in parapatry (see also Arctander & Fjeldså, 1994). However, the latter possibility cannot be entirely excluded for some species and regions.

In agreement with our results, previous work on birds (García-Moreno & Fjeldså, 2000; Caro et al., 2013) and other animals (Patton & Smith, 1992; Lynch, 1999) indicates that species replacing each other with elevation in the Neotropics are often not sister to each other. Furthermore, sister species in several vertebrate clades overlap considerably in their elevational ranges in Neotropical mountains (Cadena et al., 2012), suggesting that speciation occurs most often in allopatry within elevational zones and thus that elevational replacements result predominantly from secondary contact (but see Kozak & Wiens, 2007). Work on this topic in other tropical regions has been more limited, yet evidence from Africa (Fuchs et al., 2011) and southeast Asia (Moyle et al., 2017) also indicates secondary contact is the most likely explanation for elevational replacements (but see Bryja et al., 2018; Eldridge et al., 2018). Likewise, assembly of biotas in other mountain systems often results more from colonization by lineages from other regions than from diversification within mountains (Johansson et al., 2007; Merckx et al., 2015). Therefore, considering processes influencing the dynamics of geographic ranges which lead to secondary sympatry following divergence in allopatry is central to understanding how diversity accumulates in montane regions.

Those unfamiliar with *Scytalopus* tapaculos might be unsurprised by our finding that species originating in distinct mountains may come together into regional sympatry with elevational segregation in a given mountain. After all, tapaculos are birds and birds fly around. We, however, find this result quite striking because, unlike many birds, *Scytalopus* are notably poor dispersers (Krabbe & Schulenberg, 2003). Birds in the genus walk and hop much more than what they fly, have tiny and rounded wings which preclude long-distance powered flight, and have even lost fused

clavicles, one of the most exquisite putative adaptations of birds in general to their flighted life style. The behavior of *Scytalopus* also makes them highly reluctant to disperse: most species very rarely venture far from forest cover, having been described as agoraphobic (Krabbe & Schulenberg, 1997) or photophobic (Sick, 1993). Even during storms in the high Andes, tapaculos tend to stay put: rather than moving downslope to avoid inclement weather, individuals maintain their territories and forage in tunnel systems under the snow (Fjeldså, 1991).

How did such undispersive birds manage to get around, colonizing an isolated mountain system like the Sierra Nevada de Santa Marta twice or the two slopes of the northern Andes multiple times? Tapaculos are not alone in achieving such feats. Phylogeographic analyses of Henicorhina wood-wrens (Troglodytidae), another group of poorly dispersive birds, also revealed that elevational replacements result from secondary contact of formerly allopatric lineages (Caro et al., 2013; Cadena et al., 2019a). An explanation for the apparent paradox of poor dispersers repeatedly coming into contact from disjunct areas, even in highly isolated mountains, is that individual birds did not disperse over large distances crossing barriers now appearing unsurmountable. Rather, populations likely tracked the dynamics of their favored environments, gradually expanding their geographic distributions in concert with climatic change. During cool periods in Earth history, montane environments in tropical mountains were displaced downslope, which increased opportunities for formerly isolated areas to become connected by vegetation; in turn, isolation among such areas likely increased during warmer periods when vegetation zones retreated upslope (Hooghiemstra & Van der Hammen, 2004; Bush et al., 2011). Repeated cycles of disconnection and connection of montane areas (Ramírez-Barahona & Eguiarte, 2013) may thus have spurred cycles of allopatric speciation and subsequent secondary contact, thereby enabling the regional accumulation of diversity (Roy et al., 1997). Owing to the expected narrow thermal tolerance of tropical montane organisms (Janzen, 1967), one would expect this mechanism of divergence and accumulation of diversity in mountains to be especially prevalent in the tropics (Ghalambor et al., 2006; Kozak & Wiens, 2007; Cadena et al., 2012). An alternative, nonexclusive explanation for cycles of allopatric speciation followed by establishment of secondary sympatry, is that species may go through phases of expansion and contraction of their geographic ranges even in the absence of marked changes in the physical environment (Cadena et al., 2019a). This may occur owing to evolution of phenotypic traits influencing dispersal (Hosner et al., 2017), or to changes in ecological specialization and interactions with other species (Ricklefs, 2010).

Although adaptation to divergent selective pressures along gradients of elevation may seem like a prime precursor to the origin of new species (e.g. Funk et al., 2016; Hua, 2016), parapatric speciation along mountain slopes appears to be rare. In contrast to data discussed above, several studies do suggest that species replacing each other along elevational gradients may be closely related (Bates & Zink, 1994; Hall, 2005; DuBay & Witt, 2012), yet evidence that these replacements do not reflect separate colonization events of elevation belts or lowland-highland vicariance resulting from uplift processes (Brumfield & Edwards, 2007; Ribas et al., 2007; Santos et al., 2009) is lacking. To our knowledge, the clearest example of parapatric speciation in mountains involves sister species in the plant genus Senecio occurring on Mount Etna, Italy, which differ strikingly in ecology and phenotype despite experiencing extensive gene flow (Chapman et al., 2013; Osborne et al., 2013; Chapman et al., 2016). A promising additional case is that of Syma kingfishers in New Guinea, in which two distinct species co-occurring with elevational segregation have experienced gene flow yet maintain divergence in regions of the genome likely involved in adaptation and, presumably, mate choice (Linck et al., 2019). In contrast to the Senecio and Syma examples, avian taxa replacing each other with elevation in Neotropical mountains seldom show evidence of gene flow, with the only documented cases of interbreeding between elevational replacements in the region we are aware of being those of Anairetes tit-tyrants in Peru (Dubay & Witt, 2014) and Ramphocelus tanagers in Colombia (Morales-Rozo et al., 2017). This apparent paucity of hybridization between birds replacing each other with elevation in the Neotropics further suggests that elevational replacements did not originate through primary divergence in parapatry in the absence of barriers to gene flow, but instead via secondary contact of reproductively isolated populations. However, our inference of lack of hybridization between elevational replacements in the Andes is largely based on patterns of phenotypic variation; in most cases it remains to be seen whether genetic data reveal cryptic gene flow (Weir et al., 2015).

While we cannot reject the hypothesis that closely allied species of *Scytalopus* replacing each other along the eastern slope of the Andes colonized such region independently, our analyses indicate that parapatric speciation along the elevational gradient may have occurred there. Several of the species replacing each other with elevation in the Satipo Valley of Peru are closely related to each other, belonging to a clade of relatively recent origin in which mtDNA

divergence is shallow and rates of speciation appear faster than in the rest of the genus (Cadena et al., 2019b). Two members of this clade, S. latrans and S. micropterus, also replace each other with elevation in eastern Ecuador and Colombia. Because shallow divergence in putatively neutral loci and high rates of speciation may reflect rapid divergence mediated by adaptation in the face of gene flow, future studies should explicitly test predictions of ecological speciation (Smith et al., 2005) to determine whether elevational replacements on the eastern Andean slope may indeed be uniquely explained by parapatric divergence. The same is true for the western slope of the northern Andes, where phylogeographic patterns suggest parapatric speciation may have occurred in amphibians and reptiles (Arteaga et al., 2016; Guayasamin et al., 2017).

Beyond examining patterns of relationships among species and populations, studies of the mechanisms underlying adaptation and of how adaptive evolution in the face of gene flow may lead to speciation along elevational gradients are needed (see Hua, 2016 for a theoretical perspective). In birds, for example, putatively adaptive variation with elevation has been documented in various traits influencing functions such as respiratory physiology (Scott, 2011; Dawson et al., 2016; York et al., 2017), thermoregulation (Scott et al., 2008; Symonds & Tattersall, 2010), foraging (Kleindorfer et al., 2006; McCormack & Smith, 2008), locomotion (Altshuler et al., 2004; Milá et al., 2009), and vocal signalling (Dingle et al., 2008; Kirschel et al., 2009). Whether any of such selective pressures may account for speciation by directly or pleiotropically influencing mating patterns in Scytalopus in the eastern Andean slope and in other groups is essentially unknown.

Having understood that elevational replacements more often reflect secondary contact than parapatric divergence, a central question involving the origin of non-overlapping ranges characterizing many species assemblages from tropical mountains remains open. Do contrasting elevational distributions of species originate in allopatry or upon secondary contact? In a classic study, Diamond (1973) reasoned that elevational parapatry reflects the outcome of competitive interactions, whereby interspecific competition between formerly isolated species favors divergence of elevational ranges when they come into contact. Alternatively, contrasting elevational ranges may arise in allopatry, with the nonoverlapping ranges of species one observes reflecting sorting processes, such that only species differing in elevational ranges a priori may successfully attain regional sympatry with segregation along mountain slopes (Cadena, 2007; McEntee et al., 2018). A recent analysis revealed that sympatric sister species of birds in the tropics have more different elevational ranges than allopatric sister species regardless of their age, which was interpreted as evidence in favor of the hypothesis that elevational divergence is driven by competition upon secondary contact (Freeman, 2015). While abutting elevational ranges may indeed be maintained by competition in some cases (Cadena & Loiselle, 2007; Jankowski et al., 2010; Freeman & Montgomery, 2016), other biotic and abiotic forces may also mediate species turnover with elevation (Elsen et al., 2017). Moreover, because most cases of elevational replacements do not involve sister species, more work is necessary to determine the geographic context in which contrasting elevational ranges arise. Analyses incorporating species interactions into models of trait evolution (Nuismer & Harmon, 2015) while jointly considering the potential for such interactions to occur given geographic distributions of species (Drury et al., 2016; Clarke et al., 2017) would be a fruitful avenue for future studies on the topic (Quintero & Landis, 2019). Other mechanisms through which elevational distributions of species may change including tectonic processes of uplift or subsidence which may displace organisms vertically in passive fashion also merit consideration (Heads, 1989, 2005; Ribas et al., 2007). Scytalopus tapaculos are well suited for additional studies on the dynamics of elevational ranges integrating ecology, evolution and Earth history.

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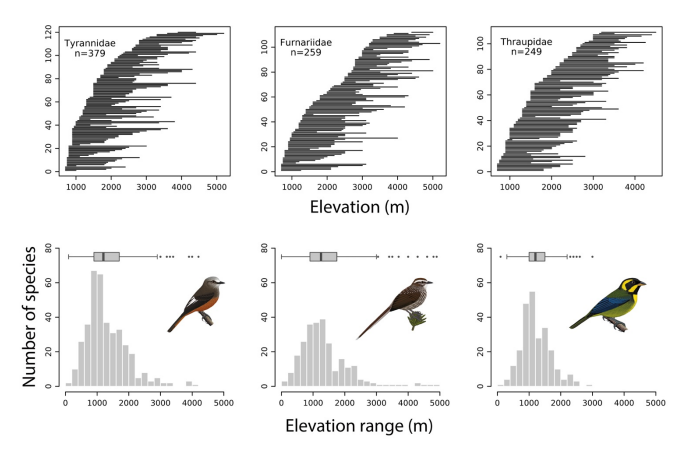


Figure 1. Restricted elevational distributions of species in three diverse families of Neotropical birds: ovenbirds (Furnariidae), tyrant flycatchers (Tyrannidae), and tanagers (Thraupidae). The panels in the top row depict the range of elevations occupied by species with lower elevational limits at or above 700 m ordered from lowest to highest along the vertical axis, showing turnover of species with elevation. Panels in the bottom row summarize the data above with histograms and boxplots, indicating that most species inhabit only a fraction of the elevational ranges existing on mountains like the Andes; across the three families, median elevational ranges span only ca. 1100 m and very few species have ranges greater than 2500 m. Data are from Parker et al. (1996) and illustrations from Ayerbe-Quiñones (Ayerbe-Quiñones, 2018).

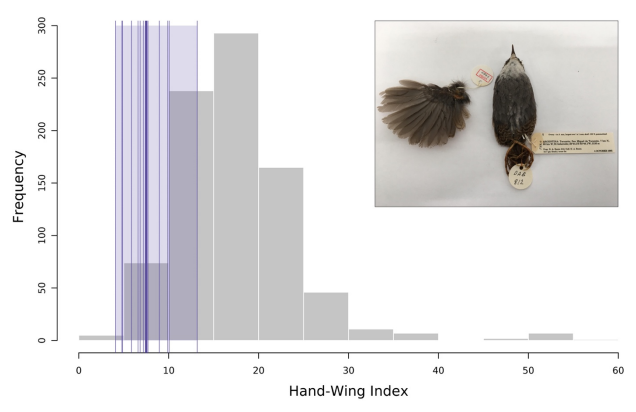


Figure 2. The frequency distribution of the hand-wing index, a proxy for dispersal abilities in birds, shows that *Scytalopus* tapaculos exhibit reduced potential for flighted dispersal relative to other birds. The dimensionless hand-wing index describes wing size and shape as a function of wing length (measured from the carpal joint to the longest primary feather) and secondary length (measured from the carpal joint to the tip of the first secondary feather), with larger values indicating greater dispersal abilities (Claramunt *et al.*, 2012). The histogram in grey depicts the distribution of the hand-wing index across a sample of 851 species of New World passerine birds (measurements from Claramunt *et al.*, 2012; P. Montoya, G. Bravo, and E. Tenorio, unpubl. data). Data for *Scytalopus* are in purple, showing the median (vertical thick bar) and the range (shaded area) of the hand-wing index across 12 species (thin bars are mean values per species). The inset illustrates the small and rounded wing of a specimen of *S. superciliaris* from Argentina housed at the Burke Museum of Natural History and Culture at the University of Washington (photograph by Cooper French).

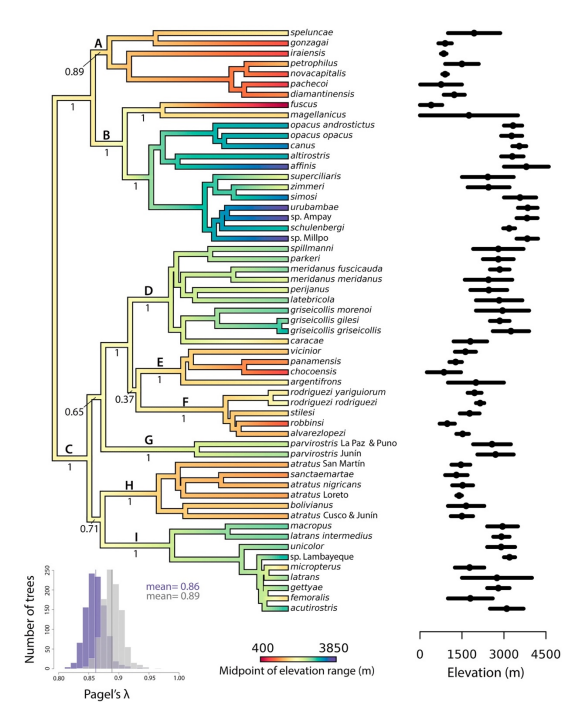


Figure 3. Elevational ranges of species are broadly similar within main clades of Scytalopus tapaculos and there is relatively high phylogenetic signal in midpoint of the elevational range. The phylogeny is the maximum clade credibility tree with midpoint of elevation mapped on branches using maximum-likelihood; posterior probabilities are shown only for major clades. The elevation range and midpoint elevation of each taxon are shown in front of each tip. The histogram in the bottom left depicts the distribution of phylogenetic signal in midpoint elevation measured by Pagel's λ estimated across 1,001 posterior trees (purple for the 47-taxa data set, gray for the 57-taxa data set).

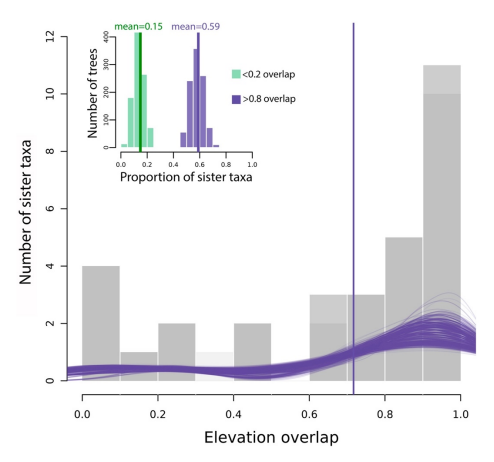


Figure 4. Most sister species of Scytalopus have similar elevational distributions. The main figure shows an overlay of 1,001 histograms, each corresponding to a phylogenetic tree in the posterior, showing the distribution of values of elevational overlap between sister taxa; purple lines are density plots and the vertical line signals the pooled mean across trees. Histograms in the inset show the distribution of the proportion of sister taxa in the 57-taxon data set with little (<0.2) and high overlap (>0.8) in elevational distributions across the 1,001 trees. We obtained qualitatively similar results with the 47-taxon data set (see text).

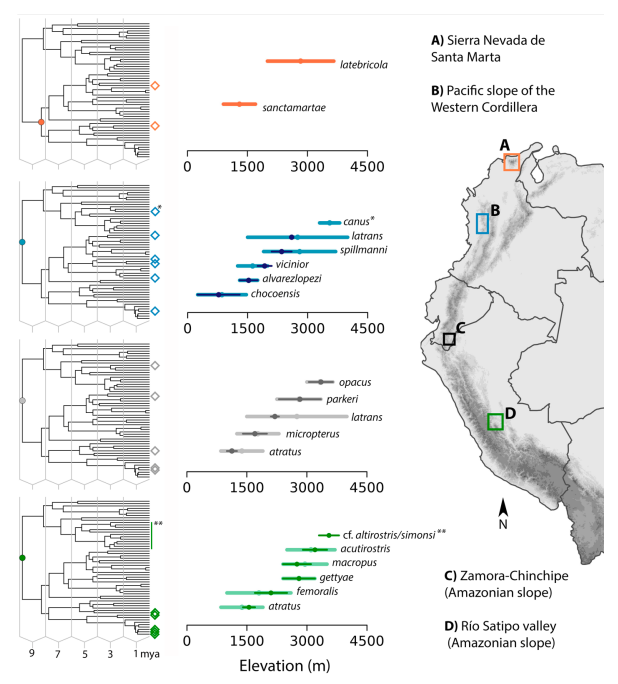


Figure 5. Scytalopus species replacing each other with elevation in four regions of South America are often, but not always, distantly related. Rhombuses at the tips of the phylogeny highlight species that replace each other along each elevational gradient, and the dot denotes the most recent common ancestor of these species. Horizontal bars depict the elevational range of species, where the lighter color represents the elevation range for the species across its distribution and the darker color the elevational range reported for each specific gradient (B: Stiles et al., 2017; C: Krabbe & Schulenberg, 1997; D: Hosner et al., 2013); a single bar is shown for the species from the Sierra Nevada de Santa Marta because both are endemic and for S. canus because the species does not occur in the specific gradient we highlight (i.e., Cerro Montezuma). The range of S. latrans in the Cerro de Montezuma is shown as a dot because it is only known from the very highest elevations in the area. Although the taxonomic identity of the species occurring at the highest elevations in the Satipo Valley gradient is uncertain, it very likely belongs to the clade depicted with the vertical line on the tree (Hosner et al., 2013).

Table 1. Elevational distributions of *Scytalopus* tapaculos considered in analyses. For each taxon, we provide the minimum, maximum and midpoint of the elevation range (in m), as well as sources for these data. Taxa with asterisks were those excluded from analyses involving only the 47 species recognized (or soon to be recognized) by taxonomists (see text).

Taxon	Min.	Max.	Mid.	Source
Scytalopus magellanicus	0	3500	1750	del Hoyo et al., 2018
Scytalopus altirostris	2900	3700	3300	Schulenberg et al., 2007
Scytalopus affinis	3000	4600	3800	Schulenberg et al., 2007
$Scytalopus\ urubambae$	3500	4200	3850	Schulenberg et al., 2007
$Scytalopus\ simosi$	3000	4150	3575	Herzog et al., 2016
Scytalopus zimmeri	1700	3200	2450	del Hoyo et al., 2018
Scytalopus superciliaris	1500	3350	2425	del Hoyo et al., 2018
Scytalopus fuscus	0	800	400	del Hoyo et al., 2018
Scytalopus canus	3300	3800	3550	A. Cuervo
Scytalopus opacus opacus	2900	3650	3275	A. Cuervo
$Scytalopus\ opacus\ and rostictus\ *$	3000	3650	3325	Krabbe & Cadena, 2010
$Scytalopus\ schulenbergi$	2975	3400	3187.5	Whitney, 1994
Scytalopus sp. Ampay	3450	4200	3825	N. Krabbe
Scytalopus sp. Millpo	3450	4220	3835	N. Krabbe
$Scytalopus\ speluncae$	1000	2870	1935	del Hoyo et al., 2018
Scytalopus gonzagai	660	1140	900	Maurício et al., 2014
$Scytalopus\ petrophilus$	900	2100	1500	Whitney et al., 2010
$Scytalopus\ diamantinensis$	850	1600	1225	Bornschein et al., 2007
$Scytalopus\ pachecoi$	10	1500	755	Maurício, 2005
Scytalopus iraiensis	750	950	850	del Hoyo et al., 2018
$Scytalopus\ nova capitalis$	800	1000	900	del Hoyo et al., 2018
Scytalopus parvirostris (La Paz & Puno)	1900	3250	2575	N. Krabbe
$Scytalopus\ parvirostris\ (Junı́n)\ *$	2050	3350	2700	N. Krabbe
Scytalopus panamensis	1050	1500	1275	del Hoyo et al., 2018
Scytalopus chocoensis	250	1465	857.5	Krabbe & Schulenberg, 1997
$Scytalopus\ rodriguezi\ rodriguezi$	2000	2300	2150	Krabbe et al., 2005
$Scytalopus\ rodriguezi\ yariguiorum\ *$	1700	2200	1950	Donegan et al, 2013
$Scytalopus\ stilesi$	1420	2130	1775	Cuervo et al., 2005
$Scytalopus\ robbinsi$	700	1250	975	Krabbe & Schulenberg, 1997
$Scytalopus\ vicinior$	1250	2000	1625	Ridgely & Greenfield, 2001
$Scytalopus\ latebricola$	2000	3660	2830	del Hoyo et al., 2018
$Scytalopus\ meridanus\ meridanus$	1600	3300	2450	del Hoyo et al., 2018
$Scytalopus\ meridanus\ fuscicauda\ *$	2500	3200	2850	Krabbe & Schulenberg, 1997
$Scytalopus \ argentifrons$	1000	3000	2000	del Hoyo et al., 2018
Scytalopus caracae	1200	2400	1800	del Hoyo et al., 2018

$Scytalopus\ spillmanni$	1900	3700	2800	del Hoyo et al., 2018
$Scytalopus\ parkeri$	2250	3350	2800	Krabbe & Schulenberg, 1997
Scytalopus griseicollis griseicollis	2600	3900	3250	Krabbe & Schulenberg, 1997
Scytalopus griseicollis gilesi	2500	3200	2850	Donegan & Avendaño, 2008
$Scytalopus\ griseicollis\ morenoi\ *$	2000	3900	2950	Avendaño & Donegan, 2015
$Scytalopus\ alvarezlopezi$	1300	1750	1525	Stiles et al., 2017
Scytalopus perijanus	1800	3120	2460	Avendaño et al., 2015
$Scytalopus\ sanctae martae$	900	1700	1300	del Hoyo et al., 2018
Scytalopus atratus nigricans (Tamá) *	1150	1900	1525	Hilty, 2003
Scytalopus atratus (Loreto)	1300	1500	1400	T. Schulenberg & D. Lane
Scytalopus atratus (SanMartín) *	1125	1800	1462.5	T. Schulenberg & D. Lane
Scytalopus atratus (Cusco & Junín) *	1100	1900	1500	T. Schulenberg & D. Lane
$Scytalopus\ bolivianus$	1000	2300	1650	del Hoyo et al., 2018
Scytalopus latrans	1500	4000	2750	del Hoyo et al., 2018
$Scytalopus\ latrans\ intermedius\ *$	2620	3200	2910	N. Krabbe
Scytalopus unicolor	2400	3400	2900	Schulenberg et al., 2007
$Scytalopus \ macropus$	2400	3500	2950	Schulenberg $et~al.,~2007;~{\rm Fjeldså~\&~Krabbe},~1990$
$Scytalopus\ micropterus$	1250	2300	1775	del Hoyo et al., 2018
$Scytalopus\ femoralis$	1000	2600	1800	Schulenberg et al., 2007
$Scytalopus\ acutirostris$	2500	3700	3100	Schulenberg et al., 2007
$Scytalopus\ gettyae$	2400	3200	2800	Hosner et al, 2013
Scytalopus sp. Lambayeque	3000	3400	3200	D. Lane