1	Distribution of iridescent colours in hummingbird communities results
2	from the interplay between selection for camouflage and communication
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Abstract

Identification errors between closely related, co-occurring, species may lead to misdirected social interactions such 12 as costly interbreeding or misdirected aggression. This selects for divergence in traits involved in species identification 13 among co-occurring species, resulting from character displacement. On the other hand, predation may select for crypsis, 14 potentially leading co-occurring species that share the same environment and predators to have a similar appearance. 15 However, few studies have explored how these antagonistic processes influence colour at the community level. Here, 16 we assess colour clustering and overdispersion in multiple hummingbird communities across Ecuador and identify the 17 processes at stake by controlling for species phylogenetic relatedness. In hummingbirds, most colours are iridescent 18 structural colours, defined as colours that change with the illumination or observation angle. Because small variations 19 in the underlying structures can have dramatic effects on the resulting colours and because iridescent structures can 20 produce virtually any hue and brightness, we expect iridescent colours to respond finely to selective pressures. Moreover, 21 we predict that hue angular dependence – a specific aspect of iridescent colours – may be used as an additional channel 22 for species recognition. In our hummingbird assemblages in Ecuador, we find support for colour overdispersion in 23 specific body patches at the community level even after controlling for the phylogeny, especially on iridescence-related 24 traits, suggesting character displacement among co-occurring species. We also find colour clustering at the community 25 level on dorsal patches, suspected to be involved in camouflage, suggesting that the same cryptic colours are selected 26 among co-occurring species. 27

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²⁹ Angle-Dependent Colouration, Community structure, Ecuador

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Colour is a complex communication channel widespread among various taxa and involved in many ecological and 31 evolutionary processes [1]. It can be described by multiple variables, including hue (colour in its common sense, such as 32 red, green, blue, etc.) and brightness (average level of grey of a colour, i.e. whether the object is light or dark). Colours 33 can be produced by two non-mutually exclusive means: pigmentary colours are produced by the selective absorption of 34 incoming light by pigments, while structural colours are produced by the interaction of incoming light with nanostructures, 35 causing diffraction, interferences or scattering [2]. Among structural colours, iridescent colours are characterised by a shift in hue with changes in illumination or observation angle [3]. Iridescent colours are found in many bird families such as 37 Anatidae (ducks) Phasianidae (fowls), Sturnidae (starlings), or Trochilidae (hummingbirds), and thought to be involved 38 in numerous adaptations [4]. But evolution of iridescent colours at the community level remains poorly understood. Yet, 39 they may display evolutionary patterns that differ from non-iridescent colours. Indeed, as opposed to other types of 40 colours, iridescent colours can produce virtually any hue and are expected to respond more readily and finely to selection, 41 because large changes of hue can be achieved by small changes in the underlying structures [5]. They can also result in 42 directional colours only seen at specific angles, as well as highly reflective colours [6]. 43

Because colours are involved in many different ecological processes, they are subject to multiple selection pressures, 44 often with opposite effects [7]. Colour may indeed increase or decrease detectability of an animal depending on the colour 45 constrast with its surroundings. In particular, colour can reduce predation risk via crypsis or aposematism or serve as a 46 means of species identification. In this case, two opposite evolutionary forces act on colours: (i) On the one hand, species 47 living in the same environment are likely experiencing similar selective pressures, such as predation. The environment 48 is characterised by ambient light and vegetation, which both influence greatly which colours are poorly detectable and 49 which colours are highly detectable [8, 9]. We thus expect co-occurring species to harbour the same, poorly detectable, 50 colours as this would decrease the risk of being detected by predators, thereby causing a clustering pattern in colouration 51 at the community level, all else being equal. This colour clustering can result from convergence between sympatric species 52 evolutionary process), from environmental filtering (ecological process), i.e. species sorting locally according to the traits 53 they harbour, or a mixture of the two (detailed in table 1). (ii) On the other hand, sympatric closely-related species are more likely to face problems of species recognition, eventually resulting in reproductive interference - a phenomenon 55 where an individual courts or mates with individuals of another species, producing no offspring or low fertility hybrids, leading to costly interbreeding [10]. Species misidentification can also lead to misdirected aggression and costly fighting 57 when individuals compete over resources or territories. Hence, any feature that would enhance species recognition is expected to be selected for. In this context, closely related species living in sympatry should be under strong selective 59 pressure to diverge in traits involved in communication, if divergence enhances species recognition. Divergence can result 60 from a process called character displacement (RCD for reproductive character displacement, ACD for agonistic character 61 displacement; evolutionary process) [11–13] or from species sorting (ecological process). For ACD, it is worth noting that 62

traits are expected to diverge only in case of moderate ecological competition, whereas they should converge in case of high competition [13, 14]. Multiple empirical studies have shown character displacement for songs (e.g. Gerhardt [15] in frogs and Grant and Grant [16] in birds), or olfactory signals [17]. However, fewer studies have looked at divergence in colour patterns (but see Sætre et al. [18], Naisbit et al. [19], Lukhtanov et al. [20], Martin et al. [21], Doutrelant et al. [22], and Hemingson et al. [23]). Almost all these studies were at the species level, and at best involved comparison between closely related species. Many of them also did not use objective spectrometry measurements and instead relied on human vision, which did not allow them to analyse colours as perceived by the intended receiver, in the case of this study: birds [24-27].

In birds, it has been showed that colouration is under different selective pressures depending on the body patch 71 location: dorsal patches, which are exposed to aerial predators, are mainly involved in camouflage while ventral and facial 72 patches are mainly involved in communication [7, 28]. In this study, we test this hypothesis for iridescent colours at 73 the community level by looking at phenotypic structure in hummingbird local assemblages across different body parts. 74 Hummingbirds are an interesting study system to test this hypothesis as various published accounts of sexual displays and 75 aggressive encounters among hummingbirds have made clear that certain feather patches such as the crown and throat are 76 consistently used during these displays [29–32]. On the other hand, colours displayed on the dorsal side of hummingbirds 77 tend to resemble background colours and thus have been suggested to be cryptic [33]. Accordingly, we predict that co-78 occurring hummingbird species should display similar hues on dorsal patches, leading to phenotypic clustering of hues 79 (i.e. co-occurring species are more similar than expected by chance, prediction 1) and different hues on ventral patches, 80 resulting in a phenotypic overdispersion pattern (i.e. co-occurring species are more dissimilar than expected by chance, 81 prediction 2). For brightness, we can formulate two alternative predictions: on the one hand, it might evolve in the same 82 way as hue, also because of reproductive character displacement and selection for camouflage, leading to the same outcome 83 as for hue (prediction 3, equivalent to predictions 1 and 2 but for brightness). On the other hand, because brightness level positively correlates with signal conspicuousness, poorly detectable signals have similar brightness, and highly detectable 85 signals have similar brightness. Hence, we may instead expect that species co-occurring should converge for brightness on all patches (prediction 3bis) if the same patches are involved in the same ecological process (communication or camouflage). 87 Compared to other types of colouration, iridescent colours might enable species recognition on another dimension in the sensory space. Two species can have the same hue or brightness at a given angle but can differ at another angle, 89 via an additional variable we call "hue shift". Because hue shift cannot be seen at long distances, it may allow species 90 to diverge without interfering with camouflage against predators [4, 34]. Accordingly, we predict overdispersion for hue 91 shift not only on ventral patches, but also on dorsal patches (prediction 4). However, hue shift is often highly correlated 92 with hue due to the optics underlying iridescence (Dakin and Montgomerie [35] for example reported $R^2 \ge 0.95$ for the 93 correlation between hue and hue shift). We test this correlation with the data from this article and discuss how it may impact our results.

At the community level, we predict that community colour volume (also known as functional richness FRic in functional ecology [36]) and brightness range increase with species richness more than expected in a random species assemblage (null ⁹⁸ model) because co-occurring species would use different colours (hue or brightness) (prediction 5).

Here we test our five predictions by quantifying both iridescent and non-iridescent colours of 189 hummingbird assemblages in Ecuador that include 112 species and span a large variety of habitats, and by assessing the phenotypic structure (clustering, random distribution, overdispersion of colours) and investigate the underlying processes by taking into account species phylogenetic relatedness within these assemblages. Comparing the uncorrected and the phylogenetically-corrected phenotypic structure of hummingbird communities will allow us to identify which mechanisms (character displacement, species sorting with mutual exclusion of similar species, environmental filtering; as detailed in table 1) underlie the community structure of iridescent colours in hummingbirds.

¹⁰⁶ Materials and methods

All scripts and data used to produce the results and figures from this article are available at https://doi.org/10.5281/ zenodo.3355444

109 Community data

Hummingbirds are particularly suited as a study system to explore the possible effect of reproductive character displace-110 ment on iridescent colours because (i) they display a large variety of hues [37] and all species harbour some iridescent 111 patches, many of which have a very strong angular dependence, rapidly shifting from e.g. pink to green or black [38, 112 39] (but note that many hummingbirds species also have non-iridescent, pigmentary, patches), (ii) they belong to a very 113 speciose family whose phylogeny is well established and readily available [40, 41], (iii) they live only in the Americas, 114 especially in the tropics where numerous species can coexist locally [37] (iv) there is an extensive documentation of hy-115 bridisation between co-occurring species (see for example [42, 43] for our region of interest), which creates the perfect 116 opportunity to study reproductive interference and (v) almost all species are available in museum collections and their 117 colour can be objectively measured using spectrometric measurements [44]. 118

Presence/absence data for hummingbird assemblages at 189 sites in Ecuador (see map in fig. S3) were compiled from data in peer-reviewed papers and reports from environmental organisations [45]. These sites cover a large variety of elevation ranges (fig. S3) and habitats [45, 46]. This dataset was previously thoroughly reviewed by comparing the observations with the known elevational and geographical ranges of each species [46] and includes observations of 112 of the 132 hummingbirds species found in Ecuador [47].

¹²⁴ Colour measurements and analyses

For each one of the 112 species, we borrowed one adult male in good condition from either the Museum National d'Histoire Naturelle (MNHN) in Paris or the Musée des Confluences, in Lyon (full list in Online Supplementary Information). We ensured that the specimen colouration was representative of the other specimens available in the collections to the human eye. When multiple subspecies were living in the area where presence was recorded, we randomly picked one of them. We

consistently took spectral reflectance measurements on the 8 following patches (described in fig. S1): crown, back, rump, tail, throat, breast, belly, wing. We also made additional measurements on patches that visually differed in colouration from these 8 main ones, as in Gomez and Théry [7] and Doutrelant et al. [22].

We measured reflectance using a setup similar to Meadows et al. [48], relying on the use of two separate optical fibres. 132 Light was conducted from an Oceanoptics DH-2000 lamp emitting over the 300-700 nm range of wavelengths to which 133 birds are sensitive [49] to the sample through an illuminating FC-UV200-2-1.5 x 100 optical fibre (named illumination 134 fibre). Light reflected by the sample was then collected by a second identical optical fibre (named collection fibre) and 135 conducted toward an Oceanoptics USB4000 spectrophotometer (used with the SpectraSuite 2.0.162 software). This setup 136 allows for a precise independent rotation of the illumination and the collection fibres, necessary for the measurements of 137 iridescent colours [6]. For more details about the measurement conditions as recommended in White et al. [50], see the 138 supplementary materials (ESM). 139

For every patch, we recorded a first reflectance spectrum at the position of the fibres which maximised total reflectance. 140 To measure hue angle dependency (iridescence), we then moved both fibres 10° away from the previous position and 141 recorded a second spectrum, as in Meadows et al. [51]. More recent measurement methods revealed that it would be more 142 accurate to keep the angular span between the illumination and collection fibres constant [52]. We however confirmed 143 that this did not impact our results by running our analyses once with all data and once with only data at a given angular 144 span (which represented 94% of the total data). All measurements were performed in a dark room with temperature 145 control. Recorded spectra were normalised by an Avantes WS-1 white standard and a measurement with the lamp shut 146 down (dark reference) and integration times were determined for each sample as to maximise the intensity of the signal 147 without saturating the spectrometer. 148

Final values were averaged over 5 consecutive measurements and spectra were smoothed using a loess algorithm and interpolated every 1 nm and negative values were set to zero using the R package pavo [53].

We analysed spectra using Endler and Mielke [54] model with relative quantum catches Q_i (without Fechner's law). 151 All birds are tetrachromats and can see light with wavelengths from 300 to 700 nm, which includes ultra-violet light (UV) 152 [55]. But different bird species vary in their sensitivity [56]: some are UV-sensitive (UVS) while others are violet-sensitive 153 VS). Literature on colour vision in hummingbirds suggests that both types are found within the family (see Chen and 154 Goldsmith [49] and Herrera et al. [57] for UVS species and Ödeen and Håstad [58] for VS species). Because we did not 155 have enough information to compute ancestral states and vision type for all species in our study and because it was 156 found to have little influence in previous studies [7, 28], we ran our analyses as if all species were VS, using the spectral 157 sensitivities of a typical VS bird, Puffinus pacificus [59], whose photoreceptor absorbances match closely those reported for 158 hummingbirds [58]. We used different illuminants defined in Endler [8], depending on the habitat of the species described 159 in Stotz et al. [60] (detailed in SI): "large gaps" illumination was used for species living in the canopy while "forest shade" 160 was used for species living in the understory. Hue was a tridimensional variable defined by the position (x, y and z) of the 161 reflectance spectrum in the tetrahedron representing bird colour vision space [54] and brightness was defined as in Endler 162 and Mielke [54] (perceived intensity of colour, also sometimes referred to as luminance). We ensured that all indices were 163

repeatable (table S1) by measuring twice the same individual and patch on 20 patches and computing the intra-class 164 coefficient (ICC) with the rptR R package [61]. We add another variable to describe iridescence: hue shift, defined as 165 the difference between hue at maximum reflectance and hue at 10° away from maximum reflectance, in a similar fashion 166 to Dakin and Montgomerie [35]. Because it is the difference of two tridimensional variables (hue at the position where 167 reflectance was maximum and hue at 10° away), hue shift is tridimensional as well. Dakin and Montgomerie [35] found a 168 high correlation between hue and hue shift at the intraspecific level in the peacock Pavo cristatus, we also report a high 169 correlation at the interspecific level in humming birds by performing a linear regression in \mathbb{R}^3 between hue and hue shift 170 $(R^2 = 0.51, F(3; 1372) = 469.7, p < 0.0001)$. New measurement methods have since been developed and propose a new 171 definition for hue shift which is not correlated to hue but they were not available at the time of this study [52]. 172

We analysed the colour volume for each species by measuring the convex hull volume of all colour patches on the bird, as suggested in Stoddard and Prum [62]. We compared the relationship between the colour volume of a community and the number of species within this community relative to a null model (prediction 5) obtained by creating random assemblages from a species pool containing all species from all communities. In other words, actual assemblages are compared to fictional assemblages with exactly the same number of species but no abiotic or biotic constraints on the species composition.

However, the colour volume does not take into account the patch location on the bird body, raising several concerns. First, two species could use the same colour but at different places on their body. They would then look different to an observer but not identified as such in this analysis. Additionally, we expect different evolutionary signals on different patches, that could even each other out, and blur the outcome at the bird level. For these reasons, we also performed our analyses separately for each one of the following eight patches: crown, back, rump, tail, throat, breast, belly, wing (locations shown in fig. S1).

¹⁸⁵ Trochilidae phylogeny and comparative analyses

A distribution of 100 phylogenetic trees of the Trochilidae family was downloaded from birdtree.org [40] to take into account phylogenetic uncertainty in the comparative analyses [63]. The 112 species included in this study constitute a fairly even sampling of the hummingbird phylogeny (fig. S2).

We used the method developed by Hardy and Senterre [64] and Baraloto et al. [65] to analyse respectively the phyloge-189 netic (Π_{ST}) and phenotypic (τ_{ST}) structures of the hummingbird communities of Ecuador (clustering or overdispersion). 190 This method relies on computing indices inspired by the Simpson index and the fixation index F_{ST} , comparing the ob-191 served diversity within and between the communities. For phylogeny, Π_{ST} can reveal phylogenetic clustering ($\Pi_{ST} > 0$) 192 or phylogenetic overdispersion ($\Pi_{ST} < 0$) within communities. Likewise, for phenotypic traits, τ_{ST} can reveal phenotypic 193 clustering $(\tau_{ST} > 0)$ or phenotypic overdispersion $(\tau_{ST} < 0)$ within communities. Statistical significance of overdispersion 194 or clustering is obtained from comparing the observed value to that obtained from 1000 random communities (created by 195 drawing from the total species pool, using algorithm 1s from Hardy [66], which keeps the local species richness per site 196 constant). This approach compares the phenotypic structure to what would be expected by chance. 197

To disentangle the relative effect of ecological (species sorting) and evolutionary mechanisms (selection), we also perform 198 our analyses by taking into account the phylogenetic relationships between species. If the species in the community are 199 more clustered or overdispersed than expected given their phylogenetic relationships, this is taken as evidence that the 200 trait has not evolved in a Brownian fashion (detailed in table 1). To this end, we used the decouple function [67], which 201 returns phylogenetically predicted and residual trait values by performing a linear regression of individual trait values 202 explained by the phylogeny. We computed the value of τ_{ST} on trait values decoupled from the phylogeny. This value is 203 hereafter denoted $dc\tau_{ST}$. Similarly to the classical τ_{ST} , the sign of $dc\tau_{ST}$ indicates phenotypic clustering ($dc\tau_{ST} > 0$) or 204 overdispersion $(dc\tau_{ST} < 0)$ once the effect of the phylogenetic structure of the communities has been decoupled. 205

	$ au_{ST} < 0$ Phenotypic overdispersion	$\tau_{ST} = 0$ No community structure	$ au_{ST} > 0$ Phenotypic clustering
$dc\tau_{ST} < 0$ Character displacement (diver- gence): co-occurring species are more dissimilar than expected given their phylogenetic relationships, which means they evolved towards dissimilarity in their colours.	Co-occurring species are less similar than expected by chance because of character displacement.	Co-occurring species are nor more neither less similar than expected by chance despite character displace- ment because closely related species co-occur more often than expected at random (phylogenetic clustering; $\Pi_{ST} > 0$).	Co-occurring species are more sim- ilar than expected by chance de- spite character displacement because closely related species co-occur more often than expected at random (phy- logenetic clustering; $\Pi_{ST} > 0$).
$dc\tau_{ST} = 0$ Brownian trait evolution	Competitive exclusion : co- occurring species are more dissimilar than expected by chance because distantly-related (and therefore dissimilar) species co-occur more often than expected at random (phy- logenetic overdispersion; $\Pi_{ST} < 0$).	Co-occurring species are not more similar nor more different than ex- pected by change or than predicted given their phylogenetic relation- ships.	Environmental filtering : co- occurring species are more similar than expected by chance because closely-related (and therefore simi- lar) species co-occur more often than expected at random (phylogenetic clustering: $Pi_{ST} > 0$).
$dc\tau_{ST} > 0$ Evolutionary convergence : co- occurring species are more similar than expected given their phylo- genetic relationships, which means they evolved towards similarity in their colours.	Co-occurring species are less sim- ilar than expected by chance de- spite evolutionary convergence be- cause distantly-related species co- occur more often than expected at random (phylogenetic overdisper- sion; $\Pi_{ST} < 0$).	Co-occurring species are neither more nor less similar than expected by chance despite evolutionary be- cause distantly-related species co- occur more often than expected at random (phylogenetic overdisper- sion; $\Pi_{ST} < 0$).	Co-occurring species are more simi- lar than expected by chance because of evolutionary convergence.

Analyses performed on a tree distribution (Π_{ST} and $dc\tau_{ST}$) with *n* trees return a distribution of *n* statistics values and *n* p-values p_i . We summarised this information by computing the median of the statistics and the overall p-value *p* by using Jost's formula [68]:

$$p = k \sum_{i=0}^{n-1} \frac{(-\ln(k))^i}{i!} \qquad \text{where } k = \prod_{i=1}^n p_i$$
(1)

209 Results

We find a strong phylogenetic clustering within communities ($\Pi_{ST} = 0.062 > 0, p < 0.0001$), indicating that co-occurring species are more closely related than expected by chance.

²¹² Phenotypic structure of the communities (predictions 1 - 4)

When looking at the bird entire body (when all patches are included simultaneously) by computing the overlap of the colour volumes, we did not find any phenotypic structure.

When the different major patches (crown, back, rump, tail, throat, breast, belly and wing) are examined separately (table 2 and table S2), we find clustering ($\tau_{ST} > 0$) in hue and hue shift on the back, rump, tail, belly and wing. Once we decouple the effect of the shared evolutionary history, we find clustering on the crown and the back ($dc\tau_{ST} > 0$) but overdispersion on the belly for both hue and hue shift ($dc\tau_{ST} < 0$). Hue shift is also overdispersed on the rump and the tail ($dc\tau_{ST} < 0$). There is no phenotypic structure on the throat, breast or wing for hue and hue shift nor on the rump or the tail for hue.

We find no phenotypic structure (neither clustering nor overdispersion) for brightness on any patches before phylogenetic correction. After phylogenetic correction, brightness values for the throat, breast and belly are clustered among co-occurring species ($dc\tau_{ST} > 0$) but show no phenotypic structure for the crown, the back, the wing and the tail.

²²⁴ Effect of community species richness on colour characteristics (prediction 5)

We found that the brightness range within a community increased in the same way as a null model built from random species assemblages (fig. 1b). For colour volume, we find some outliers with a higher colour volume than expected for community with the same number of species (fig. 1a).

228 Discussion

Our findings suggest that colour structure within hummingbird communities likely results from the interplay between two selective pressures, acting in opposite directions: selection by the local environment (e.g. camouflage from predators, leading to phenotypic clustering on dorsal patches, and selection for species recognition, leading to phenotypic overdispersion on ventral and facial patches.

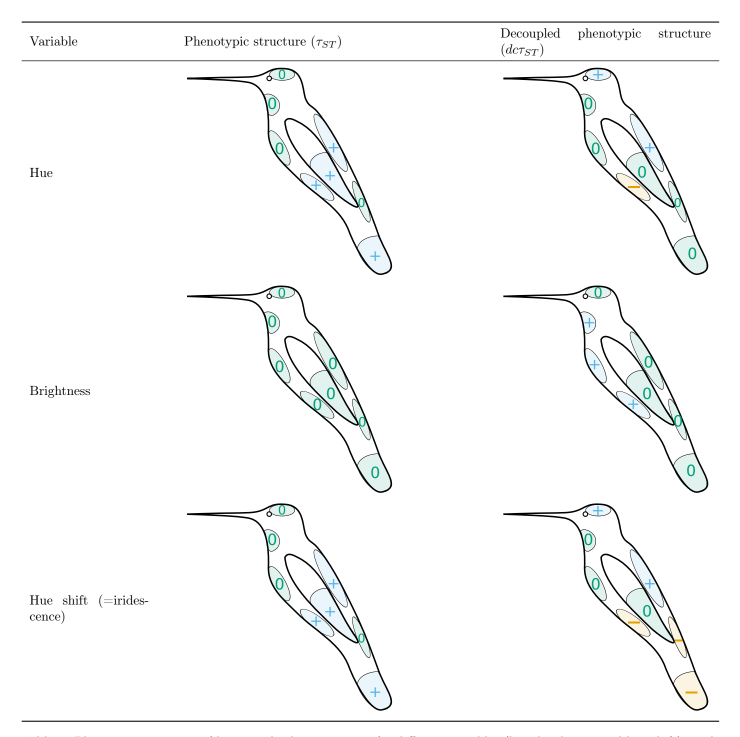


Table 2: Phenotypic structure of hummingbird communities for different variables (hue, brightness and hue shift) on the patches studied (crown, back, rump, tail, throat, breast, belly, wing; names and locations illustrated in fig. S1). Hue is a tridimensional variable defined by the reflectance spectrum position x, y and z in the tetrahedron representing avian colour space. Blue plus signs + indicate significant phenotypic clustering (τ_{ST} or $dc\tau_{ST} > 0$), orange minus signs - indicate significant phenotypic overdispersion (τ_{ST} or $dc\tau_{ST} < 0$), and green zeros 0 represent the absence of phenotypic structure. The left column shows the raw phenotypic structure of the community (columns in table 1), which may be influenced by the phylogenetic structure while the right column shows the phenotypic structure of the community, decoupled from all effects caused by the phylogeny (rows in table 1). By comparing the values of τ_{ST} and $dc\tau_{ST}$ for each trait colour variable (hue, brightness and hue shift), we can assume a probable evolutionary scenario for each patches, based on the explanation in table 1. Exact values for the statistics are available in table S2.

²³³ Evidence for different evolutionary scenarios depending on patch location

At the entire bird level (i.e. when pooling together all patches), we did not find any phenotypic structure. But as mentioned earlier, this was expected since different locations on the birds are expected to be under different selection regimes [7, 28].

In accordance with our prediction 5, community colour volume (as estimated by the convex hull of hue and brightness range within a community) increases slightly faster with the number of species in the community than predicted by a null model. This suggests that co-occurring species in these communities tend to use more similar colours than expected by chance. However, this is not the cause for the majority of communities, where co-occurring species do not use more nor less similar colours than expected by chance. This is further confirmed by the absence of phenotypic structure on the colour volume and the brightness when the effect of the phylogeny is not decoupled.

This could be the consequence of similar selective pressures between the communities we studied, leading colours in all assemblages to be randomly determined. This is however not very likely because the communities we studied differ a lot in both their vegetation background and therefore in the pressure for crypsis [45] and in their species composition. A more likely hypothesis is that co-occurring species tend to use the same colours but not necessarily on the same patches, which would also explain the absence of phenotypic structure when we pool all patches without taking into account their location. This is confirmed by our analysis patch by patch, where we find either clustering or overdispersion depending on the location of the patch.

²⁵⁰ Selection for convergence and phenotypic clustering

In accordance with our predictions, co-occurring hummingbird species tend to have similar hues on patches more likely 251 dedicated to camouflage (back, rump, tail, wing; prediction 1) but not on patches more likely used in communication 252 (crown, throat, breast; prediction 2), as shown in table 2 and table S2. This new result for iridescence colours matches 253 what has been previously described for non-iridescent colours [7, 28]. The phenotypic clustering observed for hue on the 254 rump, the tail and the wing vanishes after decoupling the clustering effect due to phylogenetic structure. This means that 255 phenotypic clustering of hue on the rump, the tail and the wing is not caused by convergent evolution of co-occurring 256 species but by environmental filtering, leading related, similar-looking species to live in the same area (as explained in 257 table 1). This is confirmed by the high value of phylogenetic clustering. This sign of phylogenetic clustering completes 258 the results from Graham et al. [45] on the same dataset. We showed that intra-community species relatedness is high 259 compared to inter-community species relatedness (Π_{ST}), while they showed that intra-community species relatedness (Net 260 Relatedness Index) is higher than expected from random assemblages in 71 % of the cases [45]. This phylogenetic clustering 261 may be caused by a strong niche conservatism but our study cannot discriminate whether such niche conservatism involves 262 colour or other ecological traits. However, hummingbirds' costly hovering flight at high elevation due to weaker lift caused 263 by the decreasing atmospheric pressure [69–71] and high foraging specialisation [72] likely contribute to this pattern. 264 Alternatively, phylogenetic clustering could also be caused by a very low dispersal ability of hummingbirds, but this 265 remains quite unlikely as the rare studies on this topic have shown that different hummingbird species display a wide

variation in their dispersal ability [73, 74].

Contrary to our prediction 2, we also find clustering of hue on the belly before the use of the decouple function. However, the fact that it turns into overdispersion after the use of the decouple function, and not simply into a random phenotypic structure (as opposed to the rump, the tail and the wing mentioned just before), suggests this initial clustering (right column in table 1) is mainly caused by environmental filtering on another trait but that hue on the belly is still under selection for divergence (first row in table 1). This other trait may be the colour of another patch or other ecological traits, as we explained previously.

We found a significant clustering of brightness on the throat, breast and belly after controlling for the phylogeny, 274 indicating that brightness on those patches is more similar than expected given the phylogeny among co-occurring species 275 (prediction 3bis). This suggests that the same patches have been selected to be involved either in communication or 276 in camouflage among species living in the same environment. This is seen after controlling for the phylogeny and it 277 is therefore not caused by the phylogenetic relatedness of co-occurring species. This is not surprising as many studies 278 showed the paramount importance of the throat in the courtship display of many hummingbird species [29–32, 75] Two 279 main hypotheses can explain why co-occurring species tend to communicate (or camouflage themselves) using the same 280 patches: (i) There may be selective pressures for the use of specific patches in camouflage in a given environment (e. g., 281 patches that are more exposed to predators' sight). (ii) Convergence in patches used in communication may be selected 282 because it improves competitor identification in the case of a strong ecological niche overlap (convergence by agonistic 283 character displacement as shown in Grether et al. [13] and Tobias et al. [76]). 284

All those results suggest a strong effect of the environment in the evolution of colour in agreement with McNaught and Owens [77] who found that bird plumage colour was due to the light environment and not to reproductive character displacement in Australian birds. However, we do not find clustering on all patches, which means that the effect of habitat pressure is somehow limited or counterbalanced by reproductive or agonistic character displacement. On the contrary, for some patches, we found patterns that are likely the result of character displacement.

²⁹⁰ Character displacement and phenotypic overdispersion

In agreement with our prediction 2, after decoupling the effect of the phylogeny, there is overdispersion of hue on the belly, likely caused by character displacement (table 1). At a completely different taxonomic scale, focusing on a single hummingbird genus (*Coeligena*) with 11 species, Parra [33] also found that the belly was always involved in the difference in hue between subspecies. It was sometimes even the only patch causing those differences, as for example between *Coeligena torquata fulgidigula* and *Coeligena torquata torquata*. This suggests that the interspecific divergence we found on the belly at the community level on the whole Trochilidae family can be observed at different geographic and taxonomic scales, and even between subspecies of the same species.

As predicted, we also find more phenotypic overdispersion for hue shift than hue after decoupling the effect of the phylogeny, for example, on the rump and on the tail (prediction 4). It is possible that hue shift is less sensitive to selection for convergence because it may vary without disturbing camouflage efficacy. However, we did not find the expected

relaxing of clustering on hue shift on patches such as the back. This is likely caused by the fact that hue shift is highly correlated with hue, as found in this study and in Dakin and Montgomerie [35], who used the same indices to quantify iridescence. This correlation is due to the optics controlling iridescence, meaning that species that display similar hues should also display the same hue shift if they use the same underlying multilayer structures. The fact that the correlation is not perfect and that we nonetheless get different phenotypic patterns for hue and hue shift on some patches suggests that co-occurring species use different multilayer structures (as recently confirmed by [78]), which can produce different iridescent effects while displaying the same hue (functional convergence on hue).

Against our prediction 2, we did not find phenotypic overdispersion on any of the colour variables on patches such as 308 the throat or the crown, that are thought to be sexually selected and often used in courtship displays [29, 79]. Several 309 hypotheses can explain this fact: (i) The overdispersion on some patches (hue on the belly and hue shift on the rump and 310 tail) is sufficient to enable species recognition. (ii) The current phenotypic structure, which is neither overdispersed nor 311 clustered, on those patches is sufficient to enable species recognition. Indeed, the absence of phenotypic overdispersion 312 does not mean that species look the same. It simply means that colour differences between species living in the same 313 community and species in different communities occur in similar ranges. This difference may be sufficient to relax the 314 selective pressure towards reproductive character displacement. (iii) The pressure towards overdispersion is balanced by 315 habitat filtering (for both ventral and dorsal patches), resulting in no apparent phenotypic structure. The latter hypothesis 316 was also a candidate explanation of the pattern found by Martin et al. [21], where sympatric closely related species are 317 more divergent than allopatric ones, but only when the range overlap is limited. They suggested that local adaptation 318 could hinder divergence when species ranges was exactly the same.(iv) Species recognition is achieved by additional means 319 and divergence occurs on others traits, such as modified feathers [80], song [81, 82] or non-vocal noises [83–85] and size. 320 Notably, different species of hummingbirds can have very different courtship behaviour: leks for hermits [86, 87], dives 321 and shuttle displays for bees [31, 84, 88], for instance. 322

Taken together, our results suggest that hummingbird iridescent colours are determined by different evolutionary 323 mechanisms depending on their location. Within a community, co-occurring hummingbird species tend to display the 324 same hues on dorsal patches probably because of selective pressures related to the local environment, such as selection 325 for crypsis by predators, causing phenotypic clustering at the community level. This phenotypic clustering does not seem 326 to be caused by adaptive convergence on colours but rather by environmental filtering perhaps linked to other ecological 327 traits such as elevation tolerance or flight ability. In spite of such environmental filtering, character displacement leads 328 to overdispersion for hue on the belly and hue shift on the rump and the tail. Iridescence may therefore enable species 329 recognition without affecting camouflage efficacy of birds, by opening up a new dimension in the sensory space: hue shift. 330

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³³⁵ Conflict of interest disclosure

- ³³⁶ The authors of this preprint declare that they have no financial conflict of interest with the content of this article. Marianne
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338 References

- JW Bradbury and SL Vehrencamp. Principles of Animal Communication. 2. ed. OCLC: 759797180. Sunderland,
 Mass: Sinauer Associates, 2011. 697 pp. ISBN: 978-0-87893-045-6.
- [2] AR Parker. "515 Million Years of Structural Colour". In: Journal of Optics A: Pure and Applied Optics 2.6 (2000),
 R15–R28. DOI: 10.1088/1464-4258/2/6/201.
- 343 [3] P Vukusic. "Natural Photonics". In: Physics World 17.2 (2004), p. 35. DOI: 10.1088/2058-7058/17/2/34.
- [4] SM Doucet and MG Meadows. "Iridescence: A Functional Perspective". In: Journal of The Royal Society Interface
 6 (Suppl 2 2009), S115–S132. DOI: 10.1098/rsif.2008.0395.focus. pmid: 19336344.
- [5] RO Prum. "Anatomy, Physics, and Evolution of Structural Colors". In: *Bird Coloration, Volume 1: Mechanisms and Measurements.* Ed. by GE Hill and KJ McGraw. Vol. 1. 2 vols. Bird Coloration. Harvard University Press, 2006,
 p. 640. ISBN: 978-0-674-01893-8.
- [6] DC Osorio and AD Ham. "Spectral Reflectance and Directional Properties of Structural Coloration in Bird Plumage".
 In: Journal of Experimental Biology 205.14 (2002), pp. 2017–2027. pmid: 12089207.
- ³⁵¹ [7] D Gomez and M Théry. "Simultaneous Crypsis and Conspicuousness in Color Patterns: Comparative Analysis of
 a Neotropical Rainforest Bird Community". In: *The American Naturalist* 169.s1 (2007), S42–S61. DOI: 10.1086/
 ³⁵³ 510138.
- JA Endler. "The Color of Light in Forests and Its Implications". In: *Ecological Monographs* 63.1 (1993), pp. 1–27.
 DOI: 10.2307/2937121.
- ³⁵⁶ [9] D Gomez and M Théry. "Influence of Ambient Light on the Evolution of Colour Signals: Comparative Analysis of
 a Neotropical Rainforest Bird Community". In: *Ecology Letters* 7.4 (2004), pp. 279–284. DOI: 10.1111/j.1461 0248.2004.00584.x.
- J Gröning and A Hochkirch. "Reproductive Interference Between Animal Species". In: The Quarterly Review of
 Biology 83.3 (2008), pp. 257–282. DOI: 10.1086/590510.
- [11] WL Brown and EO Wilson. "Character Displacement". In: Systematic Biology 5.2 (1956), pp. 49–64. DOI: 10.2307/
 2411924.
- R Butlin. "Speciation by Reinforcement". In: Trends in Ecology & Evolution 2.1 (1987), pp. 8–13. DOI: 10.1016/
 0169-5347(87)90193-5.

- GF Grether, N Losin, CN Anderson, and K Okamoto. "The Role of Interspecific Interference Competition in Char acter Displacement and the Evolution of Competitor Recognition". In: *Biological Reviews* 84.4 (2009), pp. 617–635.
 DOI: 10.1111/j.1469-185X.2009.00089.x. pmid: 19681844.
- JA Tobias and N Seddon. "Signal Design and Perception in *Hypochemis* Antbirds: Evidence for Convergent Evolution
 via Social Selection". In: *Evolution* 63.12 (2009), pp. 3168–3189. DOI: 10.1111/j.1558-5646.2009.00795.x. pmid:
 19659594.
- In: Animal Behaviour 47.4 (1994), pp. 959–969. DOI: 10.1006/anbe.1994.1127.
- BR Grant and PR Grant. "Songs of Darwin's Finches Diverge When a New Species Enters the Community". In:
 Proceedings of the National Academy of Sciences 107.47 (2010), pp. 20156–20163. DOI: 10.1073/pnas.1015115107.
 pmid: 21048082.
- PMB Bacquet, O Brattström, HL Wang, CE Allen, C Löfstedt, et al. "Selection on Male Sex Pheromone Composition
 Contributes to Butterfly Reproductive Isolation". In: *Proceedings of the Royal Society B: Biological Sciences* 282.1804
 (2015), p. 20142734. DOI: 10.1098/rspb.2014.2734. pmid: 25740889.
- [18] GP Sætre, T Moum, S Bureš, M Král, M Adamjan, et al. "A Sexually Selected Character Displacement in Flycatchers
 Reinforces Premating Isolation". In: *Nature* 387.6633 (1997), pp. 589–592. DOI: 10.1038/42451. pmid: 847.
- RE Naisbit, CD Jiggins, and J Mallet. "Disruptive Sexual Selection against Hybrids Contributes to Speciation
 between Heliconius Cydno and Heliconius Melpomene". In: Proceedings of the Royal Society of London B: Biological
 Sciences 268.1478 (2001), pp. 1849–1854. DOI: 10.1098/rspb.2001.1753. pmid: 11522205.
- VA Lukhtanov, NP Kandul, JB Plotkin, AV Dantchenko, D Haig, et al. "Reinforcement of Pre-Zygotic Isolation
 and Karyotype Evolution in Agrodiaetus Butterflies". In: Nature 436.7049 (2005), pp. 385–389. DOI: 10.1038/
 nature03704.
- PR Martin, R Montgomerie, and SC Lougheed. "Color Patterns of Closely Related Bird Species Are More Divergent
 at Intermediate Levels of Breeding-Range Sympatry". In: *The American Naturalist* 185.4 (2015), pp. 443–451. DOI:
 10.1086/680206.
- ³⁹⁰ [22] C Doutrelant, M Paquet, JP Renoult, A Grégoire, PA Crochet, et al. "Worldwide Patterns of Bird Colouration on
 ³⁹¹ Islands". In: *Ecology Letters* 19.5 (2016), pp. 537–545. DOI: 10.1111/ele.12588.
- [23] CR Hemingson, PF Cowman, JR Hodge, and DR Bellwood. "Colour Pattern Divergence in Reef Fish Species Is
 Rapid and Driven by Both Range Overlap and Symmetry". In: *Ecology Letters* 22.1 (2019), pp. 190–199. DOI:
 10.1111/ele.13180.
- ATD Bennett, IC Cuthill, and KJ Norris. "Sexual Selection and the Mismeasure of Color". In: *The American Naturalist* 144.5 (1994), pp. 848–860. DOI: 10.1086/285711.
- 397 [25] IC Cuthill, ATD Bennett, JC Partridge, and EJ Maier. "Plumage Reflectance and the Objective Assessment of Avian
- Sexual Dichromatism". In: The American Naturalist 153.2 (1999), pp. 183–200. DOI: 10.1086/303160.

- ³⁹⁹ [26] MD Eaton. "Human Vision Fails to Distinguish Widespread Sexual Dichromatism among Sexually "Monochromatic"
- Birds". In: Proceedings of the National Academy of Sciences 102.31 (2005), pp. 10942–10946. DOI: 10.1073/pnas. 0501891102. pmid: 16033870.
- [27] R Montgomerie. "Analyzing Colors". In: Bird Coloration, Volume 1: Mechanisms and Measurements. Ed. by GE Hill
 and KJ McGraw. Vol. 1. 2 vols. Bird Coloration. Harvard University Press, 2006, p. 640. ISBN: 978-0-674-01893-8.
- K Delhey. "Revealing The Colourful Side of Birds: Spatial Distribution of Conspicuous Plumage Colours on The
 Body of Australian Birds". In: *bioRxiv* (2019), p. 647727. DOI: 10.1101/647727.
- FG Stiles. "Aggressive and Courtship Displays of the Male Anna's Hummingbird". In: *The Condor* 84.2 (1982),
 pp. 208–225. DOI: 10.2307/1367674. JSTOR: 1367674.
- [30] RK Simpson and KJ McGraw. "Experimental Trait Mis-Matches Uncover Specificity of Evolutionary Links between
 Multiple Signaling Traits and Their Interactions in Hummingbirds". In: *Evolution* (2018). DOI: 10.1111/evo.13662.
- [31] RK Simpson and KJ McGraw. "Two Ways to Display: Male Hummingbirds Show Different Color-Display Tactics
 Based on Sun Orientation". In: *Behavioral Ecology* 29.3 (2018), pp. 637–648. DOI: 10.1093/beheco/ary016.
- [32] BG Hogan and MC Stoddard. "Synchronization of Speed, Sound and Iridescent Color in a Hummingbird Aerial
 Courtship Dive". In: Nature Communications 9.1 (2018), p. 5260. DOI: 10.1038/s41467-018-07562-7.
- [33] JL Parra. "Color Evolution in the Hummingbird Genus Coeligena". In: Evolution 64.2 (2010), pp. 324–335. DOI:
 10.1111/j.1558-5646.2009.00827.x. pmid: 19703221.
- [34] BD Wilts, K Michielsen, J Kuipers, H De Raedt, and DG Stavenga. "Brilliant Camouflage: Photonic Crystals in the
 Diamond Weevil, *Entimus Imperialis*". In: *Proceedings of the Royal Society B: Biological Sciences* 279.1738 (2012),
 pp. 2524–2530. DOI: 10.1098/rspb.2011.2651.
- [35] R Dakin and R Montgomerie. "Eye for an Eyespot: How Iridescent Plumage Ocelli Influence Peacock Mating Success".
 In: Behavioral Ecology 24.5 (2013), pp. 1048–1057. DOI: 10.1093/beheco/art045.
- [36] S Villéger, NWH Mason, and D Mouillot. "New Multidimensional Functional Diversity Indices for a Multifaceted
 Framework in Functional Ecology". In: *Ecology* 89.8 (2008), pp. 2290–2301. DOI: 10.1890/07-1206.1.
- [37] J Del Hoyo, A Elliott, J Sargatal, DA Christie, and E de Juana. Handbook of the Birds of the World Alive. 2017.
 URL: hbw.com.
- 425 [38] J Dorst. "Recherches sur la structure des plumes des trochilidés". OCLC: 14220401. Paris: Université de Paris, 1951.
 426 260 pp.
- 427 [39] H Dürrer. "Schillerfarben der Vogelfeder als Evolutionsproblem". Medizinischen Fakultät der Universität Basel, 1975.
- [40] W Jetz, GH Thomas, JB Joy, K Hartmann, and AO Mooers. "The Global Diversity of Birds in Space and Time".
 In: Nature 491.7424 (2012), pp. 444–448. DOI: 10.1038/nature11631. pmid: 23123857.
- 430 [41] JA McGuire, CC Witt, JVJ Remsen, A Corl, DL Rabosky, et al. "Molecular Phylogenetics and the Diversification of
- Hummingbirds". In: Current Biology 24.8 (2014), pp. 910–916. DOI: 10.1016/j.cub.2014.03.016. pmid: 24704078.

- [42] GR Graves and RL Zusi. "An Intergeneric Hybrid Hummingbird (Heliodoxa Leadbeateri Heliangelus Amethysticollis) from Northern Colombia". In: *The Condor* 92.3 (1990), pp. 754–760. DOI: 10.2307/1368695.
- FG Stiles and JO Cortés-Herrera. "Diagnosis and Observations of a Hybrid Hummingbird (Metallura Tyrianthina x
 Aglaiocercus Kingi) in the Eastern Andes of Colombia". In: *Revista de la Academia Colombiana de Ciencias Exactas*,
 Físicas y Naturales 39.153 (2015), pp. 481–490. DOI: 10.18257/raccefyn.260.
- [44] SM Doucet and GE Hill. "Do Museum Specimens Accurately Represent Wild Birds? A Case Study of Carotenoid,
 Melanin, and Structural Colours in Long-Tailed Manakins Chiroxiphia Linearis". In: Journal of Avian Biology 40.2
 (2009), pp. 146–156. DOI: 10.1111/j.1600-048X.2009.03763.x.
- [45] CH Graham, JL Parra, C Rahbek, and JA McGuire. "Phylogenetic Structure in Tropical Hummingbird Communities". In: *Proceedings of the National Academy of Sciences* 106 (Supplement 2 2009), pp. 19673–19678. DOI: 10.1073/pnas.0901649106. pmid: 19805042.
- [46] JL Parra, JA McGuire, and CH Graham. "Incorporating Clade Identity in Analyses of Phylogenetic Community
 Structure: An Example with Hummingbirds." In: *The American Naturalist* 176.5 (2010), pp. 573–587. DOI: 10.
 1086/656619. pmid: 20849270.
- [47] RS Ridgely and PJ Greenfield. The Birds of Ecuador: Status, Distribution and Taxonomy. Ithaca, NY: Cornell
 University Press, 2001. 880 pp. ISBN: 978-0-8014-8720-0.
- [48] MG Meadows, NI Morehouse, RL Rutowski, JM Douglas, and KJ McGraw. "Quantifying Iridescent Coloration in
 Animals: A Method for Improving Repeatability". In: *Behavioral Ecology and Sociobiology* 65.6 (2011), pp. 1317–
 1327. DOI: 10.1007/s00265-010-1135-5. pmid: 876.
- [49] DM Chen and TH Goldsmith. "Four Spectral Classes of Cone in the Retinas of Birds". In: Journal of Comparative
 Physiology A 159.4 (1986), pp. 473–479. DOI: 10.1007/BF00604167.
- TE White, RL Dalrymple, DW Noble, JC O'Hanlon, DB Zurek, et al. "Reproducible Research in the Study of
 Biological Coloration". In: Animal Behaviour 106 (2015), pp. 51–57. DOI: 10.1016/j.anbehav.2015.05.007.
- ⁴⁵⁵ [51] MG Meadows, TE Roudybush, and KJ McGraw. "Dietary Protein Level Affects Iridescent Coloration in Anna's
 ⁴⁵⁶ Hummingbirds, *Calypte Anna*". In: *Journal of Experimental Biology* 215.16 (2012), pp. 2742–2750. DOI: 10.1242/
 ⁴⁵⁷ jeb.069351. pmid: 22837446.
- ⁴⁵⁵ [52] H Gruson, C Andraud, W Daney de Marcillac, S Berthier, M Elias, et al. "Quantitative Characterization of Iridescent
 ⁴⁵⁹ Colours in Biological Studies: A Novel Method Using Optical Theory". In: Interface Focus 9.1 (2019), p. 20180049.
 ⁴⁶⁰ DOI: 10.1098/rsfs.2018.0049.
- [53] R Maia, H Gruson, JA Endler, and TE White. "Pavo 2: New Tools for the Spectral and Spatial Analysis of Colour
 in R". In: *Methods in Ecology and Evolution* 10.7 (2019), pp. 1097–1107. DOI: 10.1111/2041-210X.13174.
- Image: JA Endler and PW Mielke. "Comparing Entire Colour Patterns as Birds See Them". In: Biological Journal of the
 Linnean Society 86.4 (2005), pp. 405–431. DOI: 10.1111/j.1095-8312.2005.00540.x.

- ⁴⁶⁵ [55] DC Osorio and M Vorobyev. "A Review of the Evolution of Animal Colour Vision and Visual Communication
 ⁴⁶⁶ Signals". In: *Vision Research*. Vision Research Reviews 48.20 (2008), pp. 2042–2051. DOI: 10.1016/j.visres.2008.
 ⁴⁶⁷ 06.018. pmid: 18627773.
- ⁴⁶⁸ [56] A Ödeen and O Håstad. "Complex Distribution of Avian Color Vision Systems Revealed by Sequencing the SWS1
 ⁴⁶⁹ Opsin from Total DNA". In: *Molecular Biology and Evolution* 20.6 (2003), pp. 855–861. DOI: 10.1093/molbev/
- 470 msg108. pmid: 12716987.
- ⁴⁷¹ [57] G Herrera, JC Zagal, M Diaz, MJ Fernández, A Vielma, et al. "Spectral Sensitivities of Photoreceptors and Their Role
 ⁴⁷² in Colour Discrimination in the Green-Backed Firecrown Hummingbird (Sephanoides Sephaniodes)". In: Journal of
 ⁴⁷³ Comparative Physiology A 194.9 (2008), p. 785. DOI: 10.1007/s00359-008-0349-8. pmid: 18584181.
- 474 [58] A Ödeen and O Håstad. "Pollinating Birds Differ in Spectral Sensitivity". In: Journal of Comparative Physiology A
 475 196.2 (2010), pp. 91–96. DOI: 10.1007/s00359-009-0474-z.
- ⁴⁷⁶ [59] NS Hart. "Microspectrophotometry of Visual Pigments and Oil Droplets in a Marine Bird, the Wedge-Tailed Shear⁴⁷⁷ water *Puffinus Pacificus*: Topographic Variations in Photoreceptor Spectral Characteristics". In: *Journal of Exper-*⁴⁷⁸ *imental Biology* 207.7 (2004), pp. 1229–1240. DOI: 10.1242/jeb.00857. pmid: 14978063.
- ⁴⁷⁹ [60] DF Stotz, JW Fitzpatrick, TA Parker III, and DK Moskovits. *Neotropical Birds: Ecology and Conservation*. Vol. 3.
 ⁴⁸⁰ OCLC: 32819832. University of Chicago Press, 1996. ISBN: 978-0-226-77629-3.
- [61] MA Stoffel, S Nakagawa, and H Schielzeth. "rptR: Repeatability Estimation and Variance Decomposition by Generalized Linear Mixed-Effects Models". In: *Methods in Ecology and Evolution* 8.11 (2017). Ed. by S Goslee, pp. 1639– 1644. DOI: 10.1111/2041-210X.12797.
- 464 [62] MC Stoddard and RO Prum. "Evolution of Avian Plumage Color in a Tetrahedral Color Space: A Phylogenetic
 Analysis of New World Buntings." In: *The American Naturalist* 171.6 (2008), pp. 755–776. DOI: 10.1086/587526.
- ⁴⁸⁵⁶ [63] M Pagel and F Lutzoni. "Accounting for Phylogenetic Uncertainty in Comparative Studies of Evolution and Adap⁴⁸⁷⁷ tation". In: *Biological Evolution and Statistical Physics*. Ed. by M Lässig and A Valleriani. Vol. 585. Lecture Notes
 ⁴⁸⁸⁸ in Physics. Berlin, Heidelberg: Springer Berlin Heidelberg, 2002. ISBN: 978-3-540-43188-6. DOI: 10.1007/3-540⁴⁸⁹⁹ 45692-9.
- ⁴⁰⁰ [64] OJ Hardy and B Senterre. "Characterizing the Phylogenetic Structure of Communities by an Additive Partitioning of
 ⁴⁰¹ Phylogenetic Diversity". In: *Journal of Ecology* 95.3 (2007), pp. 493–506. DOI: 10.1111/j.1365-2745.2007.01222.x.
- [65] C Baraloto, OJ Hardy, CET Paine, KG Dexter, C Cruaud, et al. "Using Functional Traits and Phylogenetic Trees
 to Examine the Assembly of Tropical Tree Communities". In: *Journal of Ecology* 100.3 (2012), pp. 690–701. DOI:
 10.1111/j.1365-2745.2012.01966.x.
- [66] OJ Hardy. "Testing the Spatial Phylogenetic Structure of Local Communities: Statistical Performances of Different
 Null Models and Test Statistics on a Locally Neutral Community". In: *Journal of Ecology* 96.5 (2008), pp. 914–926.
 DOI: 10.1111/j.1365-2745.2008.01421.x.

- F de Bello, P Šmilauer, JAF Diniz-Filho, CP Carmona, Z Lososová, et al. "Decoupling Phylogenetic and Functional
 Diversity to Reveal Hidden Signals in Community Assembly". In: *Methods in Ecology and Evolution* 8.10 (2017),
 pp. 1200–1211. DOI: 10.1111/2041-210X.12735.
- [68] VN Balasubramanian, S Chakraborty, and S Panchanathan. "Conformal Predictions for Information Fusion". In:
 Annals of Mathematics and Artificial Intelligence 74.1-2 (2015), pp. 45–65. DOI: 10.1007/s10472-013-9392-4.
- [69] RK Suarez. "Hummingbird Flight: Sustaining the Highest Mass-Specific Metabolic Rates among Vertebrates". In:
 Experientia 48.6 (1992), pp. 565–570. DOI: 10.1007/BF01920240.
- ⁵⁰⁵ [70] DL Altshuler, R Dudley, and JA McGuire. "Resolution of a Paradox: Hummingbird Flight at High Elevation Does
 ⁵⁰⁶ Not Come without a Cost". In: *Proceedings of the National Academy of Sciences* 101.51 (2004), pp. 17731–17736.
 ⁵⁰⁷ DOI: 10.1073/pnas.0405260101. pmid: 15598748.
- ⁵⁰⁵ [71] DL Altshuler, FG Stiles, and R Dudley. "Of Hummingbirds and Helicopters: Hovering Costs, Competitive Ability,
 and Foraging Strategies." In: *The American Naturalist* 163.1 (2004), pp. 16–25. DOI: 10.1086/380511. pmid:
 14767833.
- AB Lindberg and JM Olesen. "The Fragility of Extreme Specialization: *Passiflora Mixta* and Its Pollinating Hum mingbird *Ensifera Ensifera*". In: *Journal of Tropical Ecology* 17.2 (2001), pp. 323–329. DOI: 10.1017/S0266467401001213.
- [73] RP Moore, WD Robinson, IJ Lovette, and TR Robinson. "Experimental Evidence for Extreme Dispersal Limitation
 in Tropical Forest Birds". In: *Ecology Letters* 11.9 (2008), pp. 960–968. DOI: 10.1111/j.1461-0248.2008.01196.x.
- [74] LN Céspedes, LI Pavan, JA Hazlehurst, and JE Jankowski. "The Behavior and Diet of the Shining Sunbeam (Aglaeactis Cupripennis): A Territorial High-Elevation Hummingbird". In: The Wilson Journal of Ornithology 131.1 (2019),
 pp. 24–34. DOI: 10.1676/18-79.
- [75] RK Simpson and KJ McGraw. "It's Not Just What You Have, but How You Use It: Solar-Positional and Behavioural
 Effects on Hummingbird Colour Appearance during Courtship". In: *Ecology Letters* 0.0 (2018). DOI: 10.1111/ele.
 13125.
- [76] JA Tobias, R Planqué, DL Cram, and N Seddon. "Species Interactions and the Structure of Complex Communication
 Networks". In: *Proceedings of the National Academy of Sciences* 111.3 (2014), pp. 1020–1025. DOI: 10.1073/pnas.
 1314337111. pmid: 24395769.
- [77] MK McNaught and IPF Owens. "Interspecific Variation in Plumage Colour among Birds: Species Recognition
 or Light Environment?" In: Journal of Evolutionary Biology 15.4 (2002), pp. 505–514. DOI: 10.1046/j.1420 9101.2002.00431.x.
- ⁵²⁷ [78] H Gruson, M Elias, C Andraud, C Djediat, S Berthier, et al. "Hummingbird Iridescence: An Unsuspected Structural
 Diversity Influences Colouration at Multiple Scales". In: *bioRxiv* (2019), p. 699744. DOI: 10.1101/699744.
- [79] CJ Clark, TJ Feo, and I Escalante. "Courtship Displays and Natural History of Scintillant (Selasphorus Scintilla)
 and Volcano (S. Flammula) Hummingbirds". In: The Wilson Journal of Ornithology 123.2 (2011), pp. 218–228. DOI: 10.1676/10-076.1.

- [80] CM Eliason, MD Shawkey, and JA Clarke. "Evolutionary Shifts in the Melanin-Based Color System of Birds". In:
 Evolution 70.2 (2016), pp. 445–454. DOI: 10.1111/evo.12855. pmid: 26044706.
- [81] P Matyjasiak. "Birds Associate Species-Specific Acoustic and Visual Cues: Recognition of Heterospecific Rivals by
 Male Blackcaps". In: *Behavioral Ecology* 16.2 (2005), pp. 467–471. DOI: 10.1093/beheco/ari012.
- [82] D Luther. "The Influence of the Acoustic Community on Songs of Birds in a Neotropical Rain Forest". In: *Behavioral Ecology* 20.4 (2009), pp. 864–871. DOI: 10.1093/beheco/arp074.
- [83] CJ Clark and TJ Feo. "The Anna's Hummingbird Chirps with Its Tail: A New Mechanism of Sonation in Birds".
 In: Proceedings of the Royal Society of London B: Biological Sciences 275.1637 (2008), pp. 955–962. DOI: 10.1098/
 rspb.2007.1619. pmid: 18230592.
- [84] CJ Clark, DO Elias, and RO Prum. "Aeroelastic Flutter Produces Hummingbird Feather Songs". In: Science 333.6048
 (2011), pp. 1430–1433. DOI: 10.1126/science.1205222. pmid: 21903810.
- ⁵⁴³ [85] CJ Clark. "Wing, Tail, and Vocal Contributions to the Complex Acoustic Signals of Courting Calliope Humming birds". In: *Current Zoology* 57.2 (2011), pp. 187–196. DOI: 10.1093/czoolo/57.2.187.
- FG Stiles and LL Wolf. "Ecology and Evolution of Lek Mating Behavior in the Long-Tailed Hermit Hummingbird".
 In: Ornithological Monographs 27 (1979), pp. iii-78. DOI: 10.2307/40166760. JSTOR: 40166760.
- [87] MA Pizo. "Lek Behavior of the Plovercrest (Stephanoxis Lalandi, Trochilidae)". In: The Wilson Journal of Ornithol ogy 124.1 (2012), pp. 106–112. DOI: 10.1676/11-055.1.
- [88] TA Hurly, RD Scott, and SD Healy. "The Function of Displays of Male Rufous Hummingbirds". In: *The Condor* 103.3 (2001), pp. 647–651. DOI: 10.1650/0010-5422(2001)103[0647:TF0D0M]2.0.C0;2.
- [89] S Nakagawa and H Schielzeth. "Repeatability for Gaussian and Non-Gaussian Data: A Practical Guide for Biologists".
- In: Biological Reviews 85.4 (2010), pp. 935–956. DOI: 10.1111/j.1469-185X.2010.00141.x. pmid: 20569253.

Table 3: List of species with their provenance (Confluences = Musée des Confluences, Lyon, France, MNHN = Muséum National d'Histoire Naturelle, Paris, France) and strata. Strata data were extracted from Stotz et al. [60] and used in vision models.

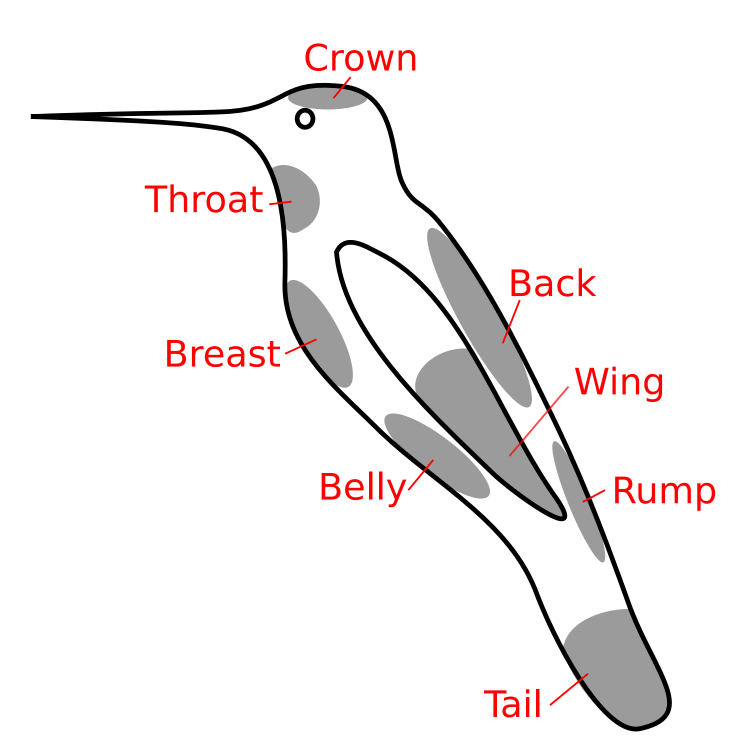
Species	Clade	Provenance	Strata
Adelomyia melanogenys	Coquette	Confluences	Understory
Aglaeactis cupripennis	Brilliant	MNHN	Canopy
Aglaiocercus coelestis	Coquette	MNHN	Canopy
Aglaiocercus kingi mocoa	Coquette	MNHN	Canopy
Amazilia amabilis	Emerald	MNHN	Understory
Amazilia amazilia	Emerald	MNHN	Understory
Amazilia fimbriata fluviatilis	Emerald	MNHN	Canopy
Amazilia franciae	Emerald	MNHN	Canopy

Species	Clade	Provenance	Strata
Amazilia grayi meridionalis	Emerald	MNHN	Canopy
Amazilia rosenbergi	Emerald	MNHN	Understory
Amazilia sapphirina	Emerald	MNHN	Canopy
Amazilia tzacatl jucunda	Emerald	MNHN	Canopy
Androdon aequatorialis	Mangoe	MNHN	Understory
Anthracothorax nigricollis	Mangoe	MNHN	Canopy
Avocettula recurvirostris	Mangoe	Confluences	Understory
Boissonneaua flavescens	Brilliant	MNHN	Canopy
Boissonneaua matthewsii	Brilliant	MNHN	Canopy
Calliphlox amethystina	Bee	MNHN	Canopy
Calliphlox mitchellii	Bee	Confluences	Canopy
Campylopterus falcatus	Emerald	MNHN	Understory
Campylopterus largipennis	Emerald	MNHN	Understory
Campylopterus villaviscensio	Emerald	MNHN	Understory
Chaetocercus bombus	Bee	MNHN	Canopy
Chaetocercus mulsant	Bee	MNHN	Understory
Chalcostigma herrani	Coquette	MNHN	Canopy
Chalcostigma ruficeps	Coquette	Confluences	Understory
Chalcostigma stanleyi stanleyi	Coquette	MNHN	Canopy
Chalybura buffonii intermedia	Emerald	Confluences	Understory
Chalybura urochrysia urochrysia	Emerald	Confluences	Understory
Chlorestes notata obsoletus-puruensis	Emerald	Confluences	Canopy
Chlorostilbon melanorhynchus	Emerald	MNHN	Understory
Chlorostilbon mellisugus phoeopygus	Emerald	Confluences	Understory
Chrysuronia oenone	Emerald	MNHN	Canopy
Coeligena coeligena	Brilliant	MNHN	Understory
Coeligena iris hesperus	Brilliant	MNHN	Understory
Coeligena iris iris	Brilliant	MNHN	Understory
Coeligena lutetiae	Brilliant	MNHN	Understory
Coeligena torquata fulgidigula	Brilliant	MNHN	Understory
Coeligena torquata torquata	Brilliant	MNHN	Understory
Coeligena wilsoni	Brilliant	MNHN	Understory
Colibri coruscans	Mangoe	MNHN	Canopy
Colibri delphinae	Mangoe	MNHN	Canopy

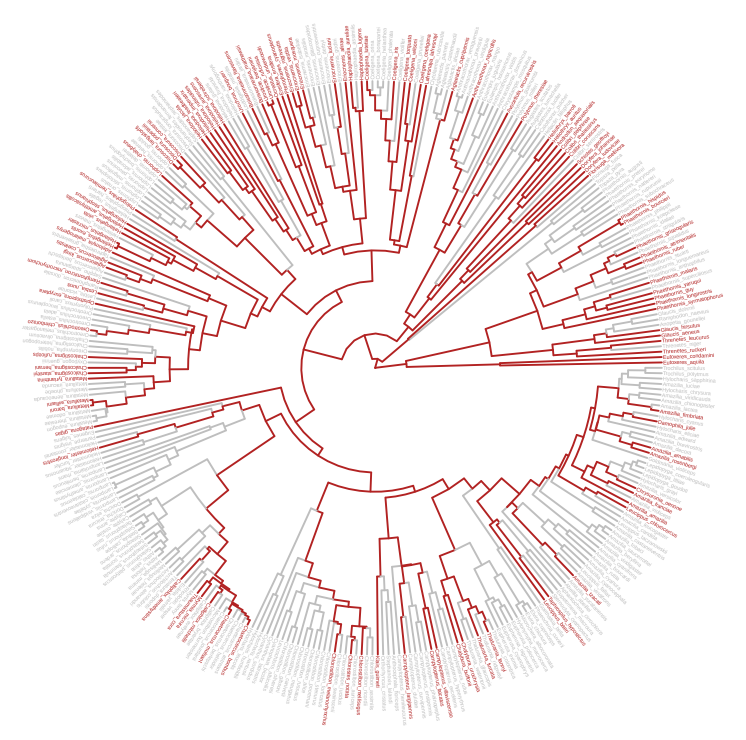
Species	Clade	Provenance	Strata
Colibri thalassinus	Mangoe	MNHN	Canopy
Damophila julie	Emerald	MNHN	Understory
Discosura conversii	Coquette	MNHN	Canopy
Discosura langsdorffi	Coquette	Confluences	Canopy
Discosura popelairii	Coquette	MNHN	Canopy
Doryfera johannae	Mangoe	MNHN	Understory
Doryfera ludovicae	Mangoe	MNHN	Understory
Ensifera ensifera	Brilliant	MNHN	Understory
Eriocnemis alinae	Brilliant	MNHN	Understory
Eriocnemis luciani	Brilliant	MNHN	Understory
Eriocnemis mosquera	Brilliant	Confluences	Understory
Eriocnemis nigrivestis	Brilliant	MNHN	Understory
Eriocnemis vestita smaragdinicollis	Brilliant	MNHN	Understory
Eutoxeres aquila	Hermit	MNHN	Understory
Eutoxeres condamini	Hermit	Confluences	Understory
Florisuga mellivora	Topazes	MNHN	Canopy
Glaucis aeneus	Hermit	MNHN	Understory
Glaucis hirsutus affinis	Hermit	MNHN	Understory
Haplophaedia aureliae russata	Brilliant	Confluences	Understory
Haplophaedia lugens	Brilliant	Confluences	Understory
Heliangelus amethysticollis laticlavius	Coquette	Confluences	Understory
Heliangelus exortis	Coquette	MNHN	Understory
Heliangelus exortis	Coquette	MNHN	Understory
Heliangelus micraster	Coquette	MNHN	Understory
Heliangelus strophianus	Coquette	MNHN	Understory
Heliangelus viola	Coquette	MNHN	Understory
Heliodoxa aurescens	Brilliant	MNHN	Understory
Heliodoxa imperatrix	Brilliant	MNHN	Understory
Heliodoxa jacula jamesoni	Brilliant	MNHN	Understory
Heliodoxa leadbeateri	Brilliant	MNHN	Understory
Heliodoxa rubinoides aequatorialis	Brilliant	MNHN	Understory
Heliodoxa schreibersii	Brilliant	MNHN	Understory
Heliomaster longirostris	MtGem	MNHN	Canopy
Heliothryx auritus	Mangoe	MNHN	Canopy

Species	Clade	Provenance	Strata
Heliothryx barroti	Mangoe	MNHN	Canopy
Klais guimeti	Emerald	MNHN	Understory
Lafresnaya lafresnayi gayi	Brilliant	Confluences	Understory
Lesbia nuna gracilis	Coquette	MNHN	Canopy
Leucippus baeri	Emerald	Confluences	Understory
Leucippus chlorocercus	Emerald	Confluences	Canopy
Lophornis chalybeus verreauxi	Coquette	MNHN	Canopy
Metallura baroni	Coquette	MNHN	Canopy
Metallura tyrianthina tyrianthina	Coquette	MNHN	Understory
Metallura williami primolina	Coquette	MNHN	Canopy
Myrmia micrura	Bee	MNHN	Canopy
Ocreatus underwoodii melanantherus	Brilliant	MNHN	Understory
Opisthoprora euryptera	Coquette	Confluences	Understory
Oreotrochilus chimborazo chimborazo	Coquette	MNHN	Understory
Oreotrochilus chimborazo jamesonii	Coquette	MNHN	Understory
Patagona gigas	Patagona	MNHN	Canopy
Phaethornis atrimentalis atrimentalis	Hermit	Confluences	Understory
Phaethornis bourcieri	Hermit	MNHN	Understory
Phaethornis griseogularis	Hermit	MNHN	Understory
Phaethornis griseogularis	Hermit	MNHN	Understory
Phaethornis guy	Hermit	MNHN	Understory
Phaethornis hispidus	Hermit	Confluences	Understory
Phaethornis longirostris	Hermit	Confluences	Understory
Phaethornis malaris	Hermit	Confluences	Understory
Phaethornis ruber	Hermit	Confluences	Understory
Phaethornis symmatophorus columbianus	Hermit	MNHN	Understory
Phaethornis yaruqui yaruqui	Hermit	MNHN	Understory
Phlogophilus hemileucurus	Coquette	MNHN	Understory
Polytmus theresiae leucorrhous	Mangoe	MNHN	Understory
Pterophanes cyanopterus	Brilliant	MNHN	Understory
Ramphomicron microrhynchum	Coquette	MNHN	Canopy
Schistes geoffroyi	Mangoe	MNHN	Understory
Taphrospilus hypostictus	Emerald	MNHN	Understory
Thalurania fannyi verticeps	Emerald	MNHN	Understory

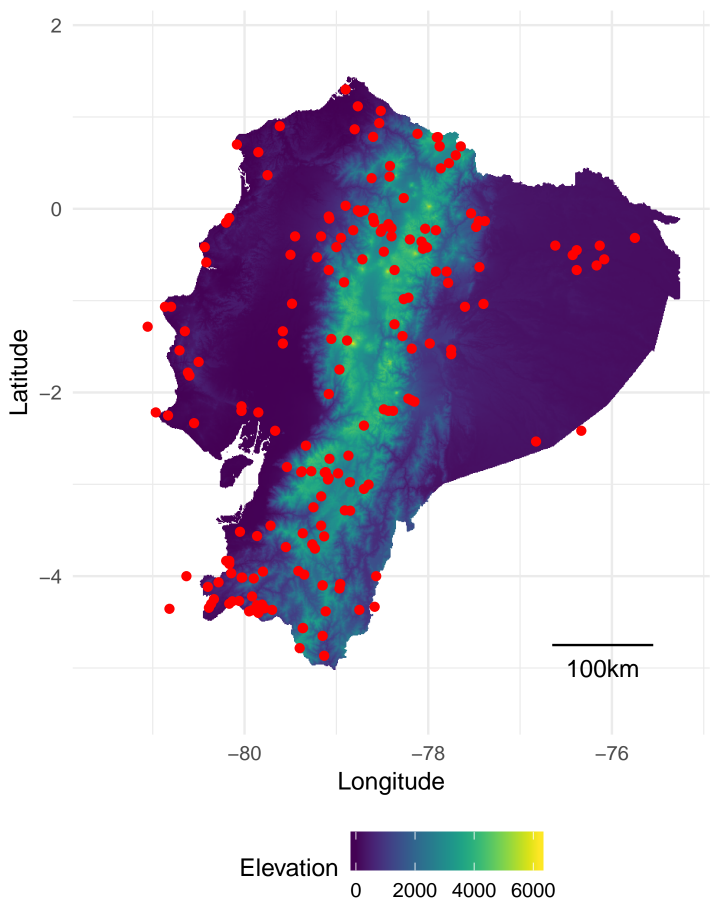
Species	Clade	Provenance	Strata
Thalurania furcata viridipectus	Emerald	MNHN	Understory
Thaumastura cora	Bee	Confluences	Canopy
Threnetes leucurus cervinicauda	Hermit	Confluences	Understory
Threnetes ruckeri	Hermit	MNHN	Understory
Urochroa bougueri	Brilliant	Confluences	Understory
Urochroa bougueri leucura	Brilliant	Confluences	Understory
Urosticte benjamini	Brilliant	MNHN	Understory
Urosticte ruficrissa	Brilliant	Confluences	Understory



Supplementary figure 1: Locations and names of the 8 patches measured on all species. Additional patches were measured for each species as soon as they differed from one of the 8 patches listed here for a human observer, as detailed in the methods section and as in Gomez and Théry [7].



Supplementary figure 2: Phylogenetic coverage of the *Trochilidae* family in our dataset (species and lineages in red).



Supplementary figure 3: Study sites locations (red dots) plotted on an altitudinal map of Ecuador. Communities outside the borders of the map are on islands or close enough to Ecuador borders to be taken into account in our study.

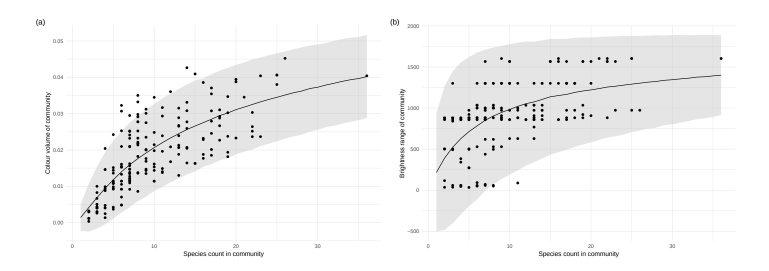
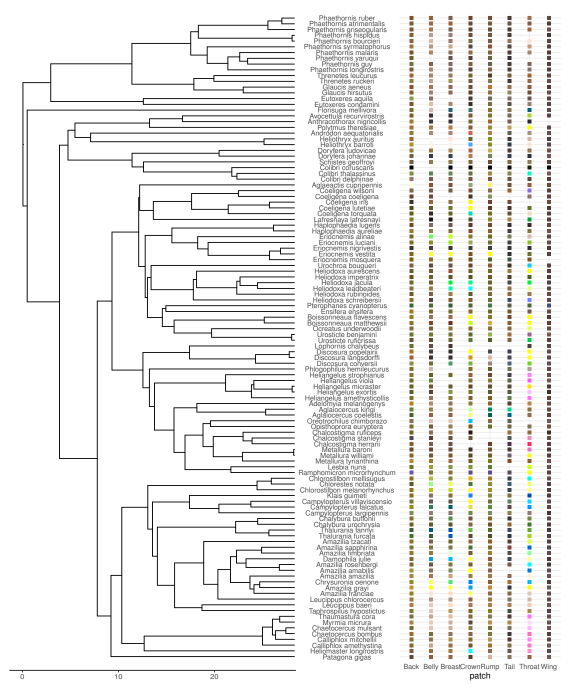


Figure 1: (a) community total colour volume and (b) brightness range increase with the number of species within the community. Each point is a community. The black solid line represents the mean value of (a) colour volume or (b) brightness range from 10 000 random communities with a given species count (null model) and the gray ribbon represents two standard deviations from the mean of the null model.

		D	iffuse	Dire	ectional	Ε	Both
Varia	ble	R	p-value	\mathbf{R}	p-value	\mathbf{R}	p-value
	х	0.734	0.002	0.877	< 0.0001	0.925	< 0.0001
Hue	у	0.923	$<\!0.0001$	0.785	0.0006	0.951	$<\!0.0001$
	\mathbf{Z}	0.780	0.0006	0.880	$<\!0.0001$	0.940	$<\!0.0001$
Bright	ness	0.411	0.090	0.055	0.48	0.373	0.04

Supplementary table 1: We quantified the repeatability R (intra-class coefficient ICC) and the related p-value by boostraping using the rptR R package [89] of indices used in this study by performing the same measurements twice on two patches for 12 species (*Coeligena torquata*, *Colibri coruscans*, *Doryfera ludovicae*, *Heliangelus strophianus*, *Heliodoxa jamesonii*, *Heliothryx barroti*, *Juliamyia julie*, *Lesbia nuna*, *Metallura tyrianthina*, *Ramphomicron microrhynchum*, *Schistes albogularis*, *Urosticte benjamini*). Patches were selected to be of similar hue from a human point of view.



Supplementary figure 4: Colour of the 8 main patches for each species in our dataset. The colour corresponds to the colour in the human visual system (CIE10). The x-axis on the phylogeny is in millions years.

variable	value	Crown	Back	Rump	Tail	Throat	Breast	Belly	Wing
Hue	$ au_{st}$	-0.0073	0.055	0.055	0.044	0.027	0.03	0.05	0.058
	$p_{ au_{st} < 0}$	0.4	1	1	1	0.9	0.9		1
	$p_{ au_{st}>0}$	0.6	0.01	0.01	0.03	0.09	0.06	0.005	0.006
	$dc\tau_{st}$	0.0099	0.026	-0.0021	0.0034	-0.0021	-0.0032	-0.01	0.00073
	$p_{ au_{st} < 0}$	1	1	0.8	1	0.9	0.3	< 0.0001	1
	$p_{ au_{st}>0}$	< 0.001	< 0.0001	1	0.2	1	1	1	1
Brightness	$ au_{st}$	-0.021	0.0078	0.0032	-0.0064	0.00015	0.0041	-0.0031	0.0091
	$p_{ au_{st} < 0}$	0.1	0.7	0.6	0.5	0.5	0.6	0.5	0.6
	$p_{ au_{st}>0}$	0.9	0.3	0.4	0.5	0.5	0.4	0.5	0.4
	$dc\tau_{st}$	-0.0014	0.0028	0.00037	0.00068	0.013	0.023	0.007	-0.0058
	$p_{\tau_{st}<0}$	0.3	1	0.9	1	1	1	1	0.2
	$p_{\tau_{st}>0}$	0.8	0.7	0.7	0.8	< 0.001	<0.001	0.002	1
Hue shift	$ au_{st}$	-0.007	0.051	0.052	0.043	0.027	0.029	0.049	0.058
	$p_{ au_{st} < 0}$	0.4	1	1	1	0.9	0.9	1	1
	$p_{ au_{st}>0}$	0.6	0.01	0.01	0.03	0.08	0.06	0.006	0.006
	$dc\tau_{st}$	0.0087	0.0059	-0.0068	-0.006	-0.0033	0.0023	-0.0098	-0.0018
	$p_{ au_{st} < 0}$	1	1	0.005	0.01	0.6	1	< 0.0001	1
	$p_{ au_{st}>0}$	< 0.001	0.03	1	1	1	0.9	1	1

Supplementary table 2: Numerical values for τ_{st} and decoupled τ_{st} (denoted $dc\tau_{st}$). P-values were computed by comparison of the actual value with the nuu distribution (obtained by randomisation of the communities using method 1s of Hardy [66]). Significant p-values are in bold and green. Positive values of $dc\tau_{st}$ indicate phenotypic clustering whereas negative values indicate overdispersion.