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ORIGINAL ARTICLE

Stronger evidence for genetic ancestry than environmental conditions in shaping the evolution of a complex signalling trait during biological invasion

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Introductions of invasive species to new environments often result in rapid rates of trait evolution. While in some cases these evolutionary transitions are adaptive and driven by natural selection, they can also result from patterns of genetic and phenotypic variation associated with the invasion history. Here, we examined the brown anole (Anolis sagrei), a widespread invasive lizard for which genetic data have helped trace the sources of non-native populations. We focused on the dewlap, a complex signalling trait known to be subject to multiple selective pressures. We measured dewlap reflectance, pattern and size in 30 non-native populations across the southeastern United States. As well, we quantified environmental variables known to influence dewlap signal effectiveness, such as canopy openness. Further, we used genome-wide data to estimate genetic ancestry, perform association mapping and test for signatures of selection. We found that among-population variation in dewlap characteristics was best explained by genetic ancestry. This result was supported by genome-wide association mapping, which identified several ancestry-specific loci associated with dewlap traits. Despite the strong imprint of this aspect of the invasion history on dewlap variation, we also detected significant relationships between dewlap traits and local environmental conditions. However, we found limited evidence that dewlap-associated genetic variants have been subject to selection. Our study emphasizes the importance of genetic ancestry and admixture in shaping phenotypes during biological invasion, while leaving the role of selection unresolved, likely due to the polygenic genetic architecture of dewlaps and selection acting on many genes of small effect.

KEYWORDS

Anolis, dewlap, genetic ancestry, invasion history, invasive species, natural selection

1 | INTRODUCTION

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Biological invasions enable researchers to conduct studies at large spatial and temporal scales that capture substantial environmental variation, facilitating the ability to observe ecological and evolutionary processes over contemporary timescales (Sakai et al., 2001; Sax et al., 2007; Lawson Handley et al., 2011; Hodgins et al., 2018). When non-native populations encounter new environments in terms of predators, competitors or abiotic conditions, this variation can relax selection experienced in their native range and/or ² WILEY-MOLECULAR ECOLOGY

impose novel selective pressures (Bock et al., 2015, 2021; Lee, 2002; Sakai et al., 2001). Invasions have provided many recent examples of rapid adaptation (e.g., Battlay et al., 2023; Colautti & Barrett, 2013; Stern & Lee, 2020). Previous studies have demonstrated phenotypic shifts in non-native species driven by natural selection occurring over the course of tens to hundreds of generations (e.g., Bock et al., 2018; Colautti & Barrett, 2013; Huey, 2000; Shine, 2011). Yet not all invasions result in rapid adaptation, even when non-native populations experience novel environments, due to a variety of factors such as a lack of associated selective pressures, little relevant genetic variation or possible genetic constraints (e.g., Alexander & Edwards, 2010; Baeckens et al., 2023; Bock et al., 2021, 2023).

In addition to examples of rapid adaptation, studies of biological invasions have revealed the importance of the invasion history for patterns of demographic, genetic and phenotypic variation (e.g., Cristescu, 2015; Kolbe et al., 2004). Invasion history includes the contribution to genetic ancestry through the location and number of source populations in the native range (Kolbe et al., 2004; Kolbe, Larson, & Losos, 2007), the spread of populations in the non-native range (e.g., leading to 'expansion load'; Peischl et al., 2013) and the extent of admixture (i.e., intraspecific hybridization of previously isolated sources) within non-native populations (Bock et al., 2021; Rius & Darling, 2014), among other factors. As a result, genetic and phenotypic variation can increase or decrease in invasive populations during an invasion. For example, admixture will often increase variation (Kolbe et al., 2004, 2008; Rius & Darling, 2014), whereas random genetic drift will decrease variation through founder effects and population bottlenecks (Dlugosch & Parker, 2008; Ficetola et al., 2008; Zhu et al., 2017). The invasion history is expected to have a strong effect on patterns of genetic and phenotypic variation within and among non-native populations and can strongly bias the interpretation of drivers of evolution during invasions, when not properly accounted for (Colautti et al., 2009; Hodgins et al., 2018; Keller & Taylor, 2008). Therefore, understanding evolution in introduced populations will often require elucidating the combined effects of invasion history and adaptation to environmental conditions encountered in the novel range.

The biological invasion of the brown anole lizard (Anolis sagrei) provides an excellent opportunity to test hypotheses for phenotypic evolution among non-native populations. Native to the northern Caribbean-Cuba, Bahamas, Cayman Brac and Little Cayman-the brown anole is a very successful invasive lizard that has been repeatedly introduced to Florida (USA) starting in the late 19th century (Garman, 1887) and expanding rapidly beginning in 1950s (Campbell, 1996). At least eight introduction events from genetically distinct native range populations have resulted in non-native populations with greater genetic diversity than those observed in the native range (Bock et al., 2021; Kolbe et al., 2004, 2008; Kolbe, Larson, & Losos, 2007). Also, non-native Florida populations are more morphologically variable in body size and shape, number of toepad lamellae, and number of scales compared to native Cuban populations (Kolbe, Larson, & Losos, 2007; Lee, 1992). This increased phenotypic and genetic variation may be acted upon by natural selection, which could promote evolution and facilitate invasion success (e.g., Shine, 2011, but see Jaspers et al., 2021). Introduced populations differ significantly from each other in genetic variation and certain morphological traits (Bock et al., 2021; Kolbe et al., 2008; Kolbe, Larson, & Losos, 2007). Thus, the invasion history may dominate patterns of phenotypic and genetic variation in these non-native populations. Indeed, genetic ancestry rather than local adaptation better explains variation among non-native brown anole populations in morphology and water loss traits (Baeckens et al., 2023; Bock et al., 2021; Kolbe, Larson, & Losos, 2007). Moreover, novel genetic interactions that result from hybridization (e.g., Bock et al., 2021) and high linkage disequilibrium in chromosomal segments of reduced recombination (i.e., Bock et al., 2023) may play a role in constraining adaptive responses in the invasive range.

Given the lack of previous evidence for local adaptation among non-native brown anole populations, we focused on a phenotypically and genetically complex trait that is subject to multiple selective pressures to increase the chance of detecting evidence of natural selection. We therefore examined evolutionary change across the invasive range in the iconic Anolis signalling ornament: the dewlap. Dewlaps are extendable structures located on the throat of anoles that differ in size, shape, colour and patterning at both the inter- and intraspecific levels (Losos, 2009). This multifaceted signalling structure often plays an important role in territory establishment and defence (e.g., Fleishman, 1992), reproductive interactions (e.g., Crews, 1975), species recognition (e.g., Losos, 1985; Rand & Williams, 1970) and predator deterrence (e.g., Leal & Rodríguez-Robles, 1995, 1997a, 1997b). Considerable evidence supports that the dewlap is subject to a variety of selection pressures (Driessens et al., 2017; Leal & Fleishman, 2002, 2004; Ng, Landeen, et al., 2013; Vanhoovdonck et al., 2009). Previous research has shown strong. albeit contrasting, relationships between environmental conditions and dewlap design in native populations of Anolis lizards (Leal & Fleishman, 2002, 2004; Ng, Landeen, et al., 2013; see Discussion for details). For brown anoles, native range populations inhabiting mesic environments had primarily marginal dewlaps (i.e., red or orange covering most of the dewlap with yellow along the outer margin), showing high reflectance in red, whereas lizards occupying xeric environments had a higher proportion of solid dewlaps with higher ultraviolet (UV) reflectance (Driessens et al., 2017). These results show that Anolis dewlap phenotypes are associated with variation in environmental conditions, likely resulting from differential selection on signal effectiveness in response to climatic conditions and physical habitat characteristics (Endler, 1992, 1993) as well as interactions with competitors, predators and conspecifics (Cole, 2013).

In this study, we investigated multiple aspects of a complex trait in brown anole populations across a large portion of its non-native range in the southeastern United States. Our goal was to determine whether among-population variation in aspects of dewlap design can be best explained by environmental conditions (consistent with local adaptation), genetic ancestry or a combination of both factors. Specifically, we tested whether variation in dewlap characteristics (i.e., colour, pattern and size) was associated with (1) differences in prevailing environmental conditions (i.e., temperature, precipitation

and light conditions) and (2) genetic ancestry as measured by the relative contribution of genome-wide SNP variation inherited from different native range source populations (Bock et al., 2021; Kolbe et al., 2004, 2008; Kolbe, Larson, & Losos, 2007). We complemented the latter ancestry-based tests with genome-wide trait association analyses, taking advantage of recent analytical developments that allow ancestry-specific associations to be detected along the genome (e.g., Skotte et al., 2019). Further, we relied on F_{ST} outlier analyses and genotype-environment association analyses, which can identify loci involved in adaptation. The phenotypically diverse and putatively polygenic nature of complex traits, such as the Anolis dewlap, afford an excellent opportunity to assess whether changing environmental conditions during an invasion shape phenotypic variation in ways consistent with natural selection. The brown anole invasion is a particularly useful case because previous studies of morphological and physiological traits have failed to provide evidence of adaptation during the invasion (i.e., Baeckens et al., 2023; Bock et al., 2021, 2023; Kolbe, Larson, & Losos, 2007). A better understanding of the evolutionary dynamics during biological invasions is useful for predicting future impacts of invasive species, especially as global environments continue to change rapidly.

2 | MATERIALS AND METHODS

2.1 | Study sites and sampling

We sampled 13-28 brown anoles from each of 30 populations in southeastern United States from Florida (N = 28 sites) and Georgia (N=2 sites; Table S1). Sites were organized along three latitudinal transects (Figure S1), completed during three field trips in March-July 2018. Sampling was limited to natural habitats with variable understory light conditions and no evidence of artificial structures. However, we sampled two populations (i.e., LOW and TIF; Table S1) from landscaped vegetation around hotel grounds due to low lizard densities in these areas. In all, we obtained 589 adult males, which were transported to the animal care facility of Harvard University. We kept the lizards for 5-9 weeks in the lab under uniform conditions in individual enclosures (see De Meyer et al., 2019 for a detailed enclosure description), during which time they were measured for a range of related projects. Dewlap characteristics were scored for a subset of 570 lizards 10-30 days after arrival from the field. All procedures described here for housing and measuring lizards were approved by the Harvard University Institutional Animal Care and Use Committee (IACUC protocol #26-11).

2.2 | Spectral data acquisition

Dewlap reflectance was measured with an Ocean Optics USB4000 spectrometer, a pulsed Xenon light source (PX-2, Ocean Optics), and a reflectance probe encased in a black anodized aluminium sheath. The reflectance probe was set at 45° to prevent specular reflection

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(i.e., glare; Endler, 1990). For each lizard, we collected reflectance data at three points: one in the centre of the dewlap and two along the edge (Figure S2a). Dewlaps were extended using stainless-steel, reverse-action forceps. For each of the three points on the dewlap, we took three repeated measurements, resulting in nine measurements per individual. We calibrated the spectrometer after every three measurements using a Spectralon white and black standard and measured reflectance from 300 (ultraviolet, UV) to 700 nm (red) in 10-nm intervals (Cuthill et al., 1999). We included the UV spectral range since *Anolis* lizards have the capacity to perceive UV signals (Fleishman et al., 1993; Fleishman & Persons, 2001).

2.3 | Spectral data processing

Spectrophotometric data were processed to reduce noise, remove technical artefacts and analyse reflectance spectra using the package pavo (Maia et al., 2013) in R v.3.6.1 (R Development Core Team, 2011). Raw spectra were smoothed using the procspec function, which applies the LOESS (Locally Weighted Scatterplot Smoothing) method with a quadratic regression and a Gaussian distribution. To normalize spectra and correct for negative values, we set the minimum value to zero and scaled up other values accordingly using the procspec function. We tested for repeatability of measurements taken at the same position on the dewlap using the R package rptR (Stoffel et al., 2017). Because all repeatability estimates were above 80%, we averaged repeated measurements using the aggspec function in pavo. We extracted four colorimetric variables: total brightness, cut-on wavelength (i.e., hue), UV reflectance and chroma. Total brightness was calculated as the area under the 'uncorrected' spectral curve from 300 to 700nm. To determine the UV reflectance and cut-on wavelength (i.e., the midpoint between baseline and maximum reflectance; Cummings, 2007), we corrected the spectra for brightness by making the area under each curve equal to 1.0 (Endler, 1990). This correction allows for the identification of differences in spectral shape that are unrelated to brightness (Ng, Landeen, et al., 2013).

2.4 | Dewlap colour composition and pattern

Each extended dewlap was photographed using a Nikon D3300 (24.2 MP) digital camera on a white background under standardized room lighting conditions in the lab. We included a colour standard (i.e., X-rite Mini ColorChecker® Classic) and a ruler for scale. Images were calibrated using the *colorChecker* function in the R package *patternize* (Van Belleghem et al., 2018). This function calculates a second-order polynomial regression between the observed and expected RGB (red, green and blue) values and performs the calibration of the image.

Because some dewlaps are a mixture of colours and the spectral data only represent three points on the dewlap, we also wanted to quantify the proportion of each colour present in each dewlap. 4 WILEY-MOLECULAR ECOLOGY

We determined the RGB values of the colours present in each pixel (size≈0.007 mm) of the dewlap image using Colour Inspector 3D (Barthel, 2006), a plugin for ImageJ (Rasband, 2012). This plugin displays the distribution of colours of an image within a 3D colour space. We extracted the RGB values and their frequency and imported them to R. We obtained a list of known colours with their respective RGB values using the base R functions colours and col-2rgb. Then, we classified dewlap RGB values for each pixel to colour categories using Euclidean distance, which determines the nearest known colour in RGB space. This quantitative measure of colour composition was calculated as the percent of red, orange and yellow present in each dewlap.

Brown anole dewlaps vary from a single colour to some combination of red-, orange- and yellow-coloured patches. A previous study of dewlap variation in the native range of the brown anole categorized dewlap patterns into two types (Driessens et al., 2017). 'Solid' dewlaps are uniformly coloured and may contain a distinct marginal colour, such as a reddish colour covering most of the dewlap with a yellowish colour along the outer margin (Figure S2b). 'Spotted' dewlaps have yellowish spots scattered across the reddish centre and may also contain a yellow outer margin (Figure S2c). We scored dewlaps from non-native populations in our study using these same two categories to facilitate comparison to the native range analysis in Driessens et al. (2017).

2.5 Dewlap size

We measured dewlap area and perimeter from digital photos in ImageJ. Dewlap area and perimeter were measured by creating line segments throughout the border of the dewlap, ultimately forming a polygon and obtaining the area and perimeter of it. We measured the dewlap for each lizard twice to verify the repeatability of this method, which was 1.0. We used a digital X-ray system to capture skeletal images from euthanized lizards. Using ObjectJ, a plugin for ImageJ, we estimated body size by measuring snout-vent length (SVL), the distance from the tip of the snout (i.e., the edge of the soft tissue) to the final vertebrae that had forward-facing arms. SVL values were then log-transformed and regressed against logtransformed dewlap area and perimeter. The resulting residual values represent body size-adjusted dewlap size.

2.6 **Environmental data**

To estimate the amount of light filtered through the forest canopy, we took hemispherical photos with a 180° fisheye lens at the location where each lizard was first observed. Canopy photos were analysed in Hemispherical_2.0, a macro for ImageJ, which calculates the percentage of pixels that are open sky (Beckschäfer, 2015).

We characterized environmental variation using Geographic Information System (GIS) climate layers interpolated from average monthly climate data from weather stations with a spatial resolution

of about 1 km² (WorldClim database; Hijmans et al., 2005). For each sampled locality, we extracted two bioclimatic variables that have been considered important for explaining dewlap colour variation in anoles: annual precipitation and annual mean temperature (Driessens et al., 2017; Ng, Landeen, et al., 2013; Williams, 1974).

2.7 **Statistical analyses**

Reflectance spectra can be interpreted as a multivariate phenotype, where reflectance values at each discrete wavelength are a variable. Therefore, dewlap colour can be represented in a multidimensional colour space. To reduce the dimensionality of the spectral data, we binned reflectance values into 10-nm bins using the procspec function in pavo (Maia et al., 2013). Each of the three positions on the dewlap where we took spectral readings was included as a separate variable, and repeated measurements at each position were averaged. We used the base R function prcomp to conduct a principal component analysis (PCA) on 123 variables representing the reflectance spectra.

We conducted linear mixed effect models using Ime4, Imer and ImerTest packages in R (Bates et al., 2015; Kuznetsova et al., 2017). For these models, we assigned principal component (PC) axes 1 or 2, which includes all colorimetric variables across the spectral range, as the response variables. Further, percent Western Cuba ancestry (see 'Population structure and genetic diversity' section) was included as a fixed factor along with the covariates annual mean temperature, annual mean precipitation and canopy openness. We also included site as a random effect to account for variation among populations not related to our main hypotheses. To further investigate dewlap variation, we used models with the same covariates, as well as fixed and random effects as previously described to evaluate individual variables describing aspects of the dewlap: UV reflectance, total brightness and hue across the three dewlap positions; colour composition of red, orange and yellow; and dewlap size (i.e., area and perimeter). We log-transformed data to improve normality when appropriate.

We performed a generalized linear model, specifically a binomial logistic regression, to identify whether any variables predict dewlap pattern (i.e., solid or spotted) using the base R function glm. The predictor variables were Western Cuba ancestry, canopy openness, annual mean temperature and annual mean precipitation. All statistical analyses were performed in R.

Genome-wide SNP genotyping 2.8

We used the same samples and quality-filtered reads included in our previous study on the genetic architecture of limb length in A. sagrei (Bock et al., 2021). This sampling includes the A. sagrei males used here for dewlap measurements, non-native population samples obtained earlier in the invasion (i.e., in 2003), and samples from the native range of A. sagrei. Using quality-filtered reads for all samples

(see detailed methods in Bock et al., 2021), we repeated the SNP calling and variant filtering steps based on version 2.1 of the A.*sa-grei* genome, which recently became available (Geneva et al., 2021). Reads were aligned to the genome using the dDocent v2.2.20 pipe-line (Puritz et al., 2014), and SNPs were called using Freebayes v. 1.3.2 (Garrison & Marth, 2012).

Filtering of the resultant variant calls was implemented using vcflib (https://github.com/vcflib/vcflib) and consisted of sequential steps based on number of alleles (i.e., keeping only biallelic markers), type of variant (i.e., keeping only SNPs), read mapping quality (i.e., using SNPs with a MAPQ score > 20), and depth of sequencing (i.e., keeping only genotypes with DP>7). For the remaining filtered SNPs, we used BCFtools v.1.9 (Narasimhan et al., 2016) to subset genotypes corresponding to the sequenced A. sagrei individuals obtained in 2018 from Florida and Georgia, for which dewlap trait data were also available (N = 561). Of the 561 samples, nine were sequenced in duplicate, resulting in 570 total sequencing libraries. We then kept only SNPs with data at more than 70% of samples and only SNPs with a minor allele freguency > 1%, and calculated identity-by-state (IBS) between samples using the SNPrelate R package (v. 1.19.4; Zheng et al., 2012). Following previous studies (e.g., Bock et al., 2021), we relied on IBS values between DNA replicates to estimate the rate of genotyping errors in our dataset. We then kept one replicate per sample and repeated the 70% call rate and minor allele frequency filters described above across all 561 genetically distinct samples. Finally, we removed one sample that had missing data at more than 30% of the final filtered SNPs, keeping the remaining 560 genotypes for downstream analyses.

Of the resulting filtered SNPs, we removed candidate gametolog SNPs following Bock et al. (2021). These gametolog SNPs occur on the X chromosome of A.*sagrei* (Geneva et al., 2021) and are a result of homology between the X chromosome (i.e., scaffold 7 in the A.*sagrei* v2.1 genome assembly) and the Y chromosome, which is currently not included in the genome assembly. After gametolog removal, we excluded markers that were in strong linkage disequilibrium (i.e., $r^2 > .4$), by scanning the genome in 5000 SNP windows, using '-indep-pairwise' option in PLINK (v1.9; Purcell et al., 2007). We then kept all SNPs located on the largest 14 scaffolds of the genome assembly, which correspond to the known number of chromosomes for A.*sagrei* and cover more than 99% of the total assembly length (Geneva et al., 2021).

2.9 | Population structure and genetic diversity

To summarize population structure, we relied on all LD-filtered autosomal SNPs and used Bayesian clustering as implemented in STRUCTURE v.2.3.4 (Pritchard et al., 2000). We considered *K* values from 1 to 10. For each *K* value, we used 20 independent replicates. Each replicate used the first 150,000 iterations as burn-in, followed 150,000 MCMC repetitions. We then used the *pophelper* (v.2.3.0; Francis, 2017) R package to estimate the most likely

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number of clusters in our data based on the delta K method (Evanno et al., 2005) and to plot the STRUCTURE results. We supplemented these analyses with a discriminant analysis of principal components (DAPC) performed using the *adegenet* R package (v. 2.1.1; Jombart & Ahmed, 2011). We further calculated heterozygosity as the percent heterozygous SNPs out of all SNPs with data for a given individual.

2.10 | Genome-wide association mapping for dewlap traits

The genome-wide association study (GWAS) was based on all LDfiltered markers. Prior to GWAS, missing data were imputed using BEAGLE v5.0 (Browning & Browning, 2016) at default settings. To investigate correlations among the 11 traits that capture dewlap size (i.e., area and perimeter) and colour (i.e., hue, total brightness, mean brightness, UV chroma, red chroma, yellow chroma, and composition of red, orange or yellow), we first used the ggcorrplot R package (v. 0.1.3; Kassambara, 2019). The GWAS was then performed using two complementary approaches, which we refer to as the 'standard GWAS' and the 'ancestry-specific GWAS'. The standard GWAS consisted of a linear mixed model implemented in GEMMA (Zhou & Stephens, 2012). We first used GEMMA to calculate 10 relatedness matrices, by excluding SNPs from one chromosome at a time (i.e., each matrix was based on SNPs from 9 chromosomes). The GWAS for markers on a given chromosome was then performed using the relatedness matrix that excluded markers on that same chromosome. This approach should improve power to detect associations, given that the relatedness matrix does not include markers that are in linkage disequilibrium with the marker being tested (Cheng et al., 2013). We then obtained p values from the Wald test as outputted by GEMMA.

The ancestry-specific GWAS was conducted in asaMap (Skotte et al., 2019). To control for the confounding effect of population structure, we relied on the first 10 PCs from a PCA computed using the *adegenet* R package (v. 2.1.1; Jombart & Ahmed, 2011), which were included as covariates in the asaMap analysis. The PCA was based on all LD-filtered autosomal SNPs. The asaMap software performs eight association tests, with and without different ancestry-specific effects considered for two ancestral populations (Skotte et al., 2019). Following asaMap manual recommendations, we estimated genome-wide ancestry proportions of each sample, as well as SNP allele frequencies for each ancestral population using ADMIX-TURE v. 1.3.0 (Alexander et al., 2009). For each dewlap trait, we obtained genome-wide *p* values for all tests and selected the test with the strongest association (i.e., the lowest observed *P* value among all tested SNPs), following Bock et al. (2021).

For both the standard and the ancestry-specific GWAS results, we used the *GenABEL* R package (Aulchenko et al., 2007) to calculate the lambda genomic inflation factor (λ) and account for any remaining *p* value inflation, which can be caused by unaccounted population stratification. Genome-wide significance thresholds were then set using the Bonferroni method (0.05/total SNPs used for the association test),

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which is the most conservative option used in association studies (e.g., Duggal et al., 2008; Kaler & Purcell, 2019). We also considered a suggestive genome-wide significance threshold (1/total SNPs used for the association test; Duggal et al., 2008), for which one false positive per GWAS is expected. To visualize GWAS results, we used Manhattan plots, made in R v.4.1.0 using publicly available scripts (https://danie Iroelfs.com/blog/how-i-create-manhattan-plots-using-ggplot/). Finally, to estimate the effect size of associations, we build linear models in R. These models had number of alternative alleles (i.e., alleles different from those incorporated in the reference genome) at the lead GWA SNP as the predictor variable, and the corresponding dewlap trait as the response variable. Effect sizes were then estimated as R^2 values, using the 'summary' function in R.

To better understand the observed effect of ancestry on dewlap traits, we next focused on results of the ancestry-specific GWAS. Specifically, we investigated how dewlap traits vary among genotype classes for ancestry-specific associations that passed the Bonferroni or suggestive significance thresholds. For this, we stratified the invasive A. sagrei samples based on ancestry, using a cut-off of 70% Western Cuba ancestry, as inferred using the STRUCTURE analysis. Because of the genetic makeup of invasive A. sagrei populations across the southeastern United States, this approach partitions samples into two groups, one with limited hybridization (i.e., Western Cuba ancestry samples) and another with frequent hybrids for which Central-eastern Cuba ancestry predominates (see also Bock et al., 2021).

Identifying the signature of local adaptation 2.11 for dewlap-associated SNPs

To determine whether SNPs associated with dewlap traits are also involved in local adaptation of invasive A. sagrei populations, we combined F_{ST} outlier analyses and genotype-environment association analyses. The F_{ST} outlier approach aimed to determine whether dewlap-associated SNPs make a large contribution to population genetic differentiation in invasive A. sagrei. We followed the approach described by Bock et al. (2021) for the same set of populations as those used for dewlap measurements here. Briefly, we used VCFtools (v. 0.1.16; Danecek et al., 2011) to calculate SNP-based F_{ST} among 15 independent population pairs found in our dataset. For this, we relied on the full LD-filtered SNP set. We then compared F_{sT} values for all dewlap-associated SNPs with F_{sT} values observed for an equal number of randomly selected non-dewlap SNPs. For this, we used a linear model in R, which had population pair and locus category (i.e., dewlap or nondewlap SNP) as predictor variables, and F_{sT} values as the response variable. As well, we designated as $F_{\rm ST}$ outliers those values that were in the top 5% of the empirical distribution, for each of the 15 population pairs. We then asked whether SNPs that are strongly associated with dewlap traits (i.e., those with the lowest inferred GWAS p value) are also repeatedly classified as F_{st} outliers in multiple independent population pairs.

The genotype-environment association (GEA) analyses aimed to determine whether dewlap-associated SNPs are also associated with environmental variables that are important from the perspective of dewlap signal effectiveness. We followed the approach described by DeVos et al. (2023) and used a latent factor mixed model (LFMM), as implemented in the Ifmm R package (v. 1.1; Frichot et al., 2013). To correct for the confounding effect of population structure, we set K=2, which corresponds to the main genetic subdivision in our dataset (see 'Population structure' results). Similar to the GWAS analyses described above, we adjusted the GEA p values based on the genomic inflation factor. We then relied on the qvalue R package (v. 2.30.0; Storey et al., 2015) to convert p values to q values and to identify genome-wide significant SNPs based on a false discovery rate (FDR) of 5%. We used canopy openness as the environmental variable, given evidence of correlation between this metric and several of the dewlap traits (see 'Associations between the dewlap, genetic ancestry and the environment' results), and because this metric was considerably more variable than temperature or precipitation across our study populations (Figure S3). Finally, we compared the canopy openness GEA results with the standard GWAS results for dewlap total brightness. We focused on this trait for two reasons. First, dewlap brightness was correlated with canopy openness, as might be expected under local adaptation. Second, the GWAS for this trait revealed several ancestry-independent loci. Thus, we could exclude the confounding effects of genomic background, which may occur for loci identified using the ancestry-specific GWAS.

RESULTS 3

3.1 **Dewlap characteristics**

Dewlap reflectance spectra revealed consistent patterns across populations (Figure S4). Despite individual variation in spectral characteristics, most lizards showed peak reflectance values between 600 and 700 nm wavelengths (Figure 1a). The relative intensity of UV wavelengths (300-400 nm) varied across individuals; dewlaps with greater proportions of yellow had higher UV reflectance values.

When comparing PC axes, dewlap reflectance showed strong differentiation between red- and yellow-dewlapped lizards (Figure 1b). PC1 explained 63.7% of the variation, whereas PC2 explained 15.6%. High values of PC1 corresponded to relatively bright dewlaps, whereas low PC1 values corresponded to dull dewlaps. As for PC2, high values corresponded to red dewlaps and low values to yellow ones.

Population structure 3.2

The final LD-filtered SNP dataset contained 25,532 markers, of which 24,574 were located on the autosomes and 958 were located on FIGURE 1 (a) Mean and standard deviation of three representative dewlap phenotypes. The area under the curve of each spectrum was normalized to one prior to averaging. Representative dewlap spectra per population can be found in Figure S4. (b) Results of a PCA for dewlap reflectance for all individuals. Spectral data for each dewlap position (P1, P2, P3) were included as separate variables divided into 10-nm intervals, resulting in 123 variables. Photos of lizards with their dewlaps extended represent the colour variation along each PC axis. High values of PC1 correspond to relatively bright dewlaps, whereas low PC1 values correspond to dull dewlaps. As for PC2, high values correspond to red dewlaps, and low values correspond to yellow ones. The shading of points corresponds to the proportion of Western Cuba ancestry inferred from the K=2 STRUCTURE results (see Figure S5).



the X chromosome. Pairwise IBS values between samples averaged 91.52% (range 88.41%–97.01%). By contrast, IBS values estimated for DNA replicates approached 100% (average 99.91%; range 99.84%-99.96%), as expected if the genotyping error rate is low (i.e., below 0.16%, as estimated using the nine replicates). The STRUCTURE analyses performed using the 24,574 autosomal SNPs confirmed our previous results on genomic variation in these populations (Bock et al., 2021). The delta K criterion indicated that two genetic clusters (K=2) are the best fit for our data (Figure S5a-c). As expected, all individuals were assigned ancestry from both genetically distinct clusters in varying proportions. Based on information from mtDNA sequences (Kolbe et al., 2004; Kolbe, Larson, & Losos, 2007) and genome-wide SNP data from the native range (Bock et al., 2021), these clusters can be interpreted as representative of ancestry from native lineages in Western and Central-eastern Cuba. Southern Florida populations have higher frequencies of Western Cuba ancestry, whereas central to northwestern Florida populations are highly admixed and contain higher frequencies of Central-eastern Cuba ancestry (Figure 2). Note,

however, that DAPC indicated that a clustering of K=3 could be a marginally better fit (Figures S6 and S7).

3.3 | Associations between the dewlap, genetic ancestry and the environment

We found significant correlations between dewlap traits, genetic ancestry and environmental variables. Dewlap variation as represented by PC1 and PC2 showed significant positive correlations with Western Cuba ancestry (p<.001, Table 1). As the frequency of Western Cuba ancestry increased, lizards exhibited brighter and redder dewlaps (Table 1, Figure 1b, Figure S8a,b). Additionally, PC1 was negatively correlated with canopy openness (p=.001; Table 1). As canopy openness increased (i.e., less canopy cover and lighter conditions), lizards exhibited darker dewlaps.

To further investigate the effects of genetic ancestry and environmental variation on distinct aspects of the dewlap and to follow



FIGURE 2 Geographical distribution of genetic ancestry for individuals from the STRUCTURE analysis. Each bar corresponds to one individual. The two genetic clusters are representative of Western Cuba (orange) and Central-eastern Cuba (purple) ancestry, based on information from previous studies using mitochondrial DNA sequences (Kolbe et al., 2004) and genome-wide SNP data (Bock et al., 2021). Pie charts represent the proportion of dewlap colour composition across sampled populations.

up on the PC-based analyses, we built separate models for each dewlap trait (i.e., univariate tests), including UV reflectance, total brightness, hue (cut-on wavelength), colour composition (red, orange, yellow), area and perimeter. Spectral colorimetric variables differed across dewlap positions; therefore, further analyses were also done by position (i.e., P1, P2 and P3). UV reflectance was negatively correlated with Western Cuba ancestry across dewlap positions (P1-edge, p = .003; P2-edge, p = .017; P3-centre, p = .003; Table 1). Centre UV reflectance showed significant positive correlations with canopy openness (P2, p < .001; P3, p = .021) and annual mean precipitation (P2, p=.032). Total brightness was positively correlated with Western Cuba ancestry (P1, P2 and P3, p < .001), negatively correlated with canopy openness (P1, p < .001; P2, p = .028) and negatively correlated with temperature (P3, p = .028). These univariate results were consistent with PC analyses, such that PC1 can be interpreted as brightness. Both PC1 (Figure S8a,I) and total brightness (Figure S8f-h,o,p) resulted in significant positive correlations

with Western Cuba ancestry and negative correlations with canopy openness across dewlap positions (Table 1). PC2 and yellow composition were correlated with Western Cuba ancestry (p < .01; Figure S8); lizards with low PC2 values had yellow dewlaps (i.e., greater yellow composition) and low frequencies of Western Cuba ancestry. As for size, dewlap area and perimeter resulted in significant positive correlations with Western Cuba ancestry (Figure S8j-k) and negative correlations with canopy openness (Figure S8q,r; Table 1). The colorimetric variable hue (cut-on wavelength) was inconclusive due to a lack of model convergence.

Binomial logistic regression indicated that the probability of a lizard having a solid dewlap increased with Western Cuba ancestry (p=.01; Figure S9c). Although we observed variation in dewlap pattern across populations (Figures S9 and S10), this trait was not significantly correlated with any environmental variable (Table 2). Furthermore, we found no significant correlation between red colour composition, genetic ancestry and environmental variables. However, yellow colour

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Dewlap characteristic	Variable	Estimate	Std. error	DF	T-statistic	p Value
P1-P3: Total reflectance	Intercept	22.261	12.034	28.506	1.850	.075
(PC1)	Western Cuba ancestry	13.491	1.761	34.922	7.660	<.001*
	Canopy openness	-0.220	0.063	30.193	-3.498	.001*
	Temperature	-0.548	0.407	27.852	-1.346	.189
	Precipitation	-0.008	0.007	26.827	-1.126	.270
P1-P3: Total reflectance	Intercept	-6.418	8.137	27.294	-0.789	.437
(PC2)	Western Cuba ancestry	5.654	1.130	44.282	5.005	<.001*
	Canopy openness	0.036	0.042	26.962	0.857	.399
	Temperature	0.252	0.275	27.455	0.914	.368
	Precipitation	-0.002	0.005	25.298	-0.394	.697
P1 – UV reflectance	Intercept	-2.822	0.529	25.965	-5.333	<.001*
	Western Cuba ancestry	-0.236	0.074	41.966	-3.211	.003*
	Canopy openness	0.005	0.003	25.670	1.899	.069
	Temperature	0.025	0.018	26.107	1.410	.170
	Precipitation	0.001	<0.001	24.068	1.649	.112
P2 – UV reflectance	Intercept	-2.164	0.385	25.384	-5.624	<.001*
	Western Cuba ancestry	-0.142	0.056	31.035	-2.521	.017*
	Canopy openness	0.008	0.002	26.931	3.883	<.001*
	Temperature	-0.006	0.013	24.787	-0.429	.671
	Precipitation	0.001	<0.001	23.885	2.281	.032*
P3 – UV reflectance	Intercept	-2.367	0.372	25.188	-6.361	<.001*
	Western Cuba ancestry	-0.175	0.054	30.770	-3.217	.003*
	Canopy openness	0.005	0.002	26.734	2.449	.021*
	Temperature	0.023	0.013	24.593	1.824	.080
	Precipitation	0.001	<0.001	23.702	1.108	.279
P1 – Total brightness	Intercept	4.324	0.201	26.390	21.494	<.001*
	Western Cuba ancestry	0.060	0.029	36.690	2.071	.045*
	Canopy openness	-0.004	0.001	26.880	-4.022	<.001*
	Temperature	-0.005	0.007	26.170	-0.709	.485
	Precipitation	<0.001	<0.001	24.640	0.005	.996
P2 – Total brightness	Intercept	4.341	0.144	27.764	30.238	<.001*
	Western Cuba ancestry	0.138	0.021	33.595	6.570	<.001*
	Canopy openness	-0.002	0.001	29.559	-2.308	.028*
	Temperature	-0.006	0.005	27.079	-1.276	.213
	Precipitation	<-0.001	<0.001	26.148	-1.969	.060
P3 – Total brightness	Intercept	4.353	0.156	27.510	27.863	<.001*
	Western Cuba ancestry	0.113	0.023	32.530	4.908	<.001*
	Canopy openness	-0.002	0.001	29.590	-2.034	.051
	Temperature	-0.012	0.005	26.750	-2.325	.028*
	Precipitation	<-0.001	<0.001	25.950	-0.548	.588
Colour composition: Red	Intercept	7.581	39.972	26.148	0.190	.851
	Western Cuba ancestry	8.745	5.458	45.910	1.602	.116
	Canopy openness	0.074	0.208	25.494	0.356	.724
	Temperature	2.062	1.352	26.416	1.525	.139

(Continues)

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Dewlap characteristic	Variable	Estimate	Std. error	DF	T-statistic	p Value
	Precipitation	-0.010	0.025	24.180	-0.386	.703
Colour composition: Orange	Intercept	4.715	0.708	26.005	6.660	<.001*
	Western Cuba ancestry	-0.118	0.099	39.781	-1.183	.244
	Canopy openness	-0.005	0.004	25.885	-1.249	.223
	Temperature	-0.048	0.024	25.984	-2.018	.054
	Precipitation	< 0.001	<0.001	24.189	0.415	.682
Colour composition: Yellow	Intercept	1.428	1.273	26.932	1.122	.272
	Western Cuba ancestry	-0.527	0.172	49.084	-3.056	.004*
	Canopy openness	0.005	0.007	26.139	0.813	.424
	Temperature	-0.010	0.043	27.279	-0.238	.813
	Precipitation	<0.001	<0.001	24.874	0.970	.341
Area	Intercept	0.448	0.159	26.790	2.809	.009*
	Western Cuba ancestry	0.088	0.022	43.020	3.978	<.001*
	Canopy openness	-0.002	0.001	26.460	-2.292	.030*
	Temperature	-0.011	0.005	26.790	-2.024	.053
	Precipitation	<-0.001	<0.001	24.880	-1.547	.135
Perimeter	Intercept	0.214	0.077	26.710	2.784	.010*
	Western Cuba ancestry	0.045	0.011	44.800	4.261	<.001*
	Canopy openness	-0.001	<0.001	26.210	-2.767	.010*
	Temperature	-0.004	0.003	26.820	-1.692	.102
	Precipitation	-0.001	<0.001	24.760	-1.750	.092

Note: Colorimetric variables (i.e., UV reflectance and total brightness) were obtained using edge (P1/P2) and centre (P3) dewlap measurements. We log-transformed UV reflectance, total brightness, colour composition (i.e., red, orange and yellow) and dewlap area to improve normality. Residual area represents body size-adjusted dewlap size. Hue (cut-on wavelength) was inconclusive due to lack of model convergence. Significant *p* values are indicated with an asterisk (*).

TABLE 2Results of the binomial logistic regression modelperformed to detect relationships between dewlap pattern,Western Cuba ancestry and environmental variables (i.e.,temperature, precipitation and canopy openness).

Variable	Estimate	Std. error	Z-value	p Value
Intercept	2.489	2.295	1.085	.278
Western Cuba ancestry	-0.910	0.337	-2.696	.007*
Canopy openness	-0.018	0.012	-1.413	.158
Temperature	0.009	0.076	0.120	.904
Precipitation	-0.002	0.001	-1.165	.244

Note: We categorized dewlap patterns into two types; 'solid' dewlaps are uniformly colored and may contain a margin and 'spotted' dewlaps having yellowish spots scattered across the reddish center and may also contain a yellow outer margin. Significant *P*-values are indicated with an asterisk (*).

composition was negatively correlated with Western Cuba ancestry (p=.004, Figure S8i). These results are consistent with analyses of PC2. Lizards with low PC2 values had yellow dewlaps (Figure 1b) and low frequencies of Western Cuba ancestry (Figure 1b; Figure S8b).

3.4 | Genetic architecture and selection tests for dewlap traits

Analysis of correlations among traits (Figure S11) revealed strong and significant correlations between our measurements of dewlap size (i.e., area and perimeter; r=.91) and between our measurements of dewlap brightness (i.e., mean and total dewlap brightness; r=1). Moderate, albeit still significant, positive correlations were further revealed between dewlap hue, yellow chroma and brightness, as well as between dewlap size and dewlap red chroma. Finally, dewlap UV chroma was negatively correlated with most dewlap traits (Figure S11).

The linear mixed model implemented in GEMMA (i.e., standard GWAS), identified seven loci distributed on chromosomes 2, 4, 6, 7 and 11 as significantly associated with the percent of red colour (Figure 3; Table S2) at the Bonferroni-adjusted significance level. These associations were of moderate effect as estimated using percent variance explained (PVE; 3.5%–9.6%; Figure 3). All other analyses using the GEMMA model did not reveal any associations with dewlap traits at the Bonferroni-adjusted significance level (Table S2). There were, however, SNPs that passed the suggestive

FIGURE 3 Genome-wide association results for percent red colour in dewlaps. (a) Manhattan plots for the LMM implemented in GEMMA (top) and the ancestry-specific association model implemented in asaMap (bottom). The red-dashed lines mark the genome-wide significance threshold calculated using the Bonferroni method. A more permissive suggestive significance level is shown using the black-dashed lines. (b) Dewlap red percent (log-transformed) values shown for the three loci with the largest PVE. For each genotype class, points show the mean, and the black lines indicate standard error. The numbers shown in parentheses under each genotype class indicate sample sizes.



genome-wide significance threshold. These suggestive associations were identified for nine of the dewlap traits and were distributed on all macrochromosomes except chromosome 3, as well as on microchromosomes 9, 11 and 12 (Figure 3; Table S2).

The ancestry-specific GWAS identified one significant locus at the Bonferroni-adjusted level (Figure 4; Table S2), located on macrochromosome 2. This locus was associated with total and mean brightness with a moderate effect in samples of Western Cuba ancestry (PVE=3%; Figure 4), and a much smaller effect for samples in the Central-eastern Cuba ancestry group (PVE=0.7%; Figure 4). Other ancestry-specific associations (N=37 loci) were identified at the suggestive genome-wide threshold for 10 of the dewlap traits, distributed on chromosomes 1, 2, 4, 5, 6 and 11 (Figure 4; Table S2).

The F_{ST} values estimated for the 45 dewlap-associated SNPs (i.e., loci with both significant and suggestive associations with dewlap traits; Table S2) ranged from 0 to 0.62 (average 0.04), whereas F_{ST} for the 45 random SNPs ranged from 0 to 0.43 (average 0.03). The difference in F_{ST} between the dewlap-associated SNPs and the random SNPs was not, however, significant (p=.08; Figure S12a). As well, most dewlap-associated SNPs (i.e., 89%, or 40/45 SNPs) were not classified as F_{ST} outliers (Figure S12b). The remaining five dewlap-associated SNPs were F_{ST} outliers in one or two population pairs at

most (Figure S12b). The GEA analysis identified one SNP on macrochromosome 2 at coordinate 22,431,024 that was significantly associated with canopy openness (q=0.009; Figure S13a). However, this SNP was not associated with any dewlap traits (Figure S13b; Table S2), and it was not classified as an F_{ST} outlier, displaying F_{ST} values that ranged from 0 to 0.33 (average 0.05).

4 | DISCUSSION

Rapid adaptation is a common feature of biological invasions (e.g., Battlay et al., 2023; Colautti & Barrett, 2013; Stern & Lee, 2020). Counter to results for many invasions, previous studies of the brown anole in the southeastern United States have revealed a lack of evidence for adaptation to the novel environmental conditions experienced there (Baeckens et al., 2023; Bock et al., 2021, 2023; Kolbe, Larson, & Losos, 2007). This is surprising given that the colonization of the non-native range in this species likely coincided with substantial changes in ecological interactions, as well as a shift from a tropical to a temperate climate. Consistent with previous studies of phenotypic evolution in the brown anole invasion, our results show that variation in dewlap characteristics among non-native



FIGURE 4 Ancestry-specific associations identified for dewlap traits. Only the three loci showing the strongest association signals are shown (all other associations are listed in Table S2). Of these, the association for dewlap brightness passed the Bonferroni significance threshold, whereas the other two loci passed the suggestive significance threshold. The effect of alleles at each SNP on trait values was calculated separately for the 'Western Cuba ancestry' and the 'Central-eastern Cuba ancestry' sample groups. Sample sizes for each sample group are given below each genotype.

populations is primarily explained by genetic ancestry. Admixture of multiple native range lineages strongly influences the current genetic composition of non-native populations (Bock et al., 2021; Kolbe et al., 2004, 2008; Kolbe, Glor, et al., 2007). GWAS results corroborated these findings and indicated that most dewlap traits appear to have a complex genetic architecture. While we find that multiple aspects of the dewlap are related to local environmental conditions, genomic analyses did not indicate that dewlapassociated SNPs retain a signature consistent with local adaptation. Whether natural selection plays a role in signal divergence during the brown anole invasion thus requires further investigation. This study provides insight into the ongoing evolutionary processes occurring in biological invasions, highlighting the importance of genetic ancestry in brown anole dewlap variation among populations. Below, we put our findings in the context of previous studies of the brown anoles to better understand how populations have evolved during this invasion.

4.1 | Genetic ancestry and its role in shaping dewlap phenotypes

Genetic clustering analyses revealed that genetic ancestry of invasive brown anoles can be summarized at a coarse level as a combination of two genetically distinct native lineages: Western Cuba and Central-eastern Cuba. Genetic ancestry had strong effects on several aspects of the dewlap; lizards with increasing Western Cuba ancestry have larger, brighter red dewlaps with lower UV reflectance and less yellow (Table 1, Figure 1b; Figure S8a-k). As well, the probability of lizards having a solid dewlap increased with the frequency of Western Cuba ancestry (Figure S9c). This result is consistent with findings from the native range which, while based on more limited sampling, also find that Western Cuba populations of *A. sagrei* have a greater proportion of solid dewlaps, whereas Central-eastern Cuba populations have a greater proportion of spotted dewlaps (Driessens et al., 2017). These results point to an important role of genetic ancestry in shaping multiple dimensions of dewlap variation across invasive populations in this system.

Our genome-wide association analyses further explored the effect of genetic ancestry on dewlap variation. We identified several SNPs significantly associated with red coloration and brightness of dewlaps, and dozens of SNPs with suggestive associations with all dewlap traits (Figure 3, Table S2). Our finding that genetic ancestry has strong and widespread effects on dewlap differentiation among non-native populations is consistent with previous studies of invasive species (e.g., Silene vulgaris, Keller & Taylor, 2010; sticklebacks, Lucek et al., 2010). We previously used the same samples to identify a large-effect locus significantly associated with limb length in A. sagrei (Bock et al., 2021). Our results here indicate that the genetic architecture of the dewlap is likely different and more complex than that characterizing limb length, at least for A. sagrei of Cuban ancestry. This interpretation is further supported by among-trait correlations. While all limb length traits show strong and significant pairwise correlations (Bock et al., 2021), dewlap traits are characterized by more moderate among-trait correlations that are occasionally non-significant (Figure S11).

The associations identified here between SNPs and dewlap traits should be interpreted with caution for two reasons. First, aside from associations reported for red composition and total and mean dewlap brightness, the strength of the association signal is reduced for any one SNP. Second, in-depth analyses of standard and ancestry-specific associations (Figures 3 and 4) revealed that genotype classes with the largest difference in trait values are also the ones with the lowest sample size (e.g., genotype GG for Western Cuba ancestry, at Chr2: 166.587 Mb; Figure 4). Therefore, some of these results could be due to uneven sample sizes among genotype classes. With these caveats in mind, we note that the direction of the ancestry-specific effects reported here is consistent with our observations of the effect of Western Cuba ancestry on dewlap trait variation. Specifically, linear mixed effects models of dewlap traits based on genetic ancestry as well as the ancestry-specific GWAS indicated that Western Cuba ancestry is associated with increased brightness of dewlaps.

In many organisms, carotenoid pigments have been shown to be a source of variation in red, orange and yellow ornamental coloration within and between species (reviewed in Toews et al., 2017). In most animal species, carotenoids cannot be synthesized and must be ingested. The extent to which dewlap colour variation is influenced by nutritional condition has so far been investigated in a few anole species, A. distichus (Ng, Kelly, et al., 2013) and A. sagrei (Steffen et al., 2010). Both studies found no difference in colour between carotenoid and control treatments, and dewlap coloration was heritable in A. distichus lizards. Moreover, biochemical investigations of dewlap pigment composition in several Anolis species have found that pterin pigments are important sources of coloration, although carotenoids were detected as well (Macedonia et al., 2000; Ortiz, 1962; Steffen & McGraw, 2007, 2009). In contrast to carotenoids, pterins are produced endogenously (reviewed in Andrade & Carneiro, 2021). Even so, identifying genes involved in pterin synthesis has been challenging because of the complexity of underlying biochemical pathways. For example, studies contrasting skin patches of different colours in other reptiles have identified tens to hundreds of genes that are differentially expressed, and that are likely involved in the production of these pigments (e.g., McLean et al., 2017). Our finding that dewlap phenotypes are correlated with genomewide estimates of genetic ancestry across non-native populations is in line with these previous studies and suggests a complex genetic architecture for dewlap traits.

4.2 | Potential adaptation to light environments

We tested the hypothesis that dewlap phenotypes are correlated with environmental variation, which is expected under local adaptation. Supporting this prediction, we found significant relationships between multiple environmental variables and dewlap characteristics. Our strongest results show that dewlaps tend to be darker with relatively high UV reflectance as canopy openness (i.e., habitat light) increases. Also, we found that brown anoles occurring in habitats with greater precipitation had more UV reflectance. These findings are consistent with those obtained by previous studies in the native range of *A.sagrei*. Driessens et al. (2017) found that lizards occupying open forest habitats had higher UV reflectance than populations inhabiting environments with little light exposure.

Studies in other Anolis species show different correlations between UV reflectance and the environment, suggesting the relationship between aspects of the dewlap and the local environmental conditions might be species specific. For example, Ng, Landeen, et al. (2013) found no difference in UV reflectance among A. distichus populations inhabiting environments that differed in light characteristics. Also, Leal and Fleishman (2002, 2004) reported that A. cristatellus populations in Puerto Rico occupying closed forest habitats with little light penetration exhibit highly reflective and transmissive dewlaps with more UV reflectance. As light exposure increased (i.e., greater canopy openness), dark dewlaps with low UV reflectance were favoured (Leal & Fleishman, 2004).

Several studies on local adaptation of Anolis dewlaps have contrasted populations in xeric and mesic habitats. These habitats differ in characteristics such as habitat light, temperature or precipitation. We sampled brown anole populations across a latitudinal gradient throughout Florida and into southern Georgia. The southeast region of the United States is known to have relatively uniform climatic conditions within seasons. As expected, we found less variation in annual mean temperature and precipitation patterns among our sampled populations (Figure S3b,c) as compared to previous studies of tropical mesic and xeric habitats in Puerto Rico and the Dominican Republic (Leal & Fleishman, 2004; Ng, Landeen, et al., 2013). Despite this reduced variation in temperature and precipitation, we found that lizards inhabiting cooler environments tend to have dewlaps with greater orange composition, which is consistent with previous studies of A. distichus (Ng, Landeen, et al., 2013). While this may indicate convergent evolution of dewlap orange composition in relation to temperature in A. distichus and A. sagrei, the underlying mechanism is unknown.

In contrast to analyses at the trait level discussed above, analyses at the SNP level did not provide evidence of adaptation. Specifically, SNPs associated with dewlap traits had similar $F_{\rm ST}$ values as random genome-wide SNPs (Figure S12a). As well, dewlap-associated SNPs were rarely classified as $F_{\rm ST}$ outliers (Figure S12b). Moreover, the small number of dewlap-associated SNPs that do have extreme $F_{\rm ST}$ values are highly genetically differentiated in one or two population pairs at most (Figure S12b). These results therefore indicate that, if some of the dewlap-associated SNPs reported here are involved in local adaptation, their contribution is likely specific to a small number of populations, with different SNPs being recruited by natural selection in different populations. The genotype-environment association analyses further supported these results, indicating that dewlap-associated SNPs are not also associated with the environmental variables predicted

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to shape dewlap traits under local adaptation (Figure S13). Given evidence presented here that points to a complex genetic architecture for dewlaps and considering that invasive brown anole populations are highly genetically diverse, it is possible that adaptation occurs via small changes in allele frequency at a large number of loci. Identifying the genetic signature of natural selection at loci involved in the control of the dewlap is needed to confirm the occurrence of adaptation, as inferred from trait-environment correlations. Achieving this task will be challenging and will require much denser genome-wide data, in line with observations for other highly polygenic traits (Lowry et al., 2017).

In conclusion, the brown anole invasion allowed us to study the evolution of a complex signalling phenotype during a biological invasion, revealing how genetic ancestry strongly influences amongpopulation variation in dewlaps in the non-native range. Our study supports the importance of invasion history and admixture in determining patterns of phenotypic divergence during biological invasions (Keller & Taylor, 2010; Kolbe et al., 2004). Although we found some evidence that aspects of the dewlap are correlated with environmental variation among non-native populations, which is consistent with previous studies and suggestive of local adaptation, the loci underlying these dewlap characteristics did not show a genetic signature consistent with the action of natural selection. Future studies should consider denser sampling of SNPs along the genome, which may allow the signature of natural selection to be recovered for these polygenic traits. Additionally, to better understand dewlap evolution in non-native brown anole populations, future studies should consider other potential sources of selection, including species recognition (Baeckens, Driessens, & Van Damme, 2018; Losos, 1985; Vanhooydonck et al., 2009), sexual selection (Vanhooydonck et al., 2009), and intrasexual selection (i.e., male-male competition; Baeckens, Driessens, Huyghe, et al., 2018; Lailvaux & Irschick, 2007; Vanhooydonck et al., 2005).

AUTHOR CONTRIBUTIONS

J.N.P.A., D.G.B., and J.J.K. designed research; J.N.P.A., D.G.B., and S.B. performed research; J.N.P.A. and D.G.B. analysed data; and J.N.P.A., D.G.B., S.B., J.B.L., and J.J.K. contributed to writing the paper.

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CONFLICT OF INTEREST STATEMENT

The authors declare no competing interest.

DATA AVAILABILITY STATEMENT

Raw sequence data used for this project are available in the Sequence Read Archive (SRA) https://www.ncbi.nlm.nih.gov/sra (Bio-Project accession PRJNA737437). The scripts used for analyses, the dewlap trait data, the ancestry data and the environmental data are available from FigShare (doi: 10.6084/m9.figshare.24009063.v2).

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