Geo-referenced population-specific microsatellite data across American continents, the MacroPopGen Database

Elizabeth R. Lawrence¹, Javiera N. Benavente¹,², Jean-Michel Matte¹, Kia Marin¹,³, Zachery R. R. Wells¹,⁴, Thaïs A. Bernos¹, Nia Krasteva¹, Andrew Habrich¹,⁵, Gabrielle A. Nessel¹, Ramela Arax Koumrouyan¹ & Dylan J. Fraser¹

Population genetic data from nuclear DNA has yet to be synthesized to allow broad scale comparisons of intraspecific diversity versus species diversity. The MacroPopGen database collates and geo-references vertebrate population genetic data across the Americas from 1,308 nuclear microsatellite DNA studies, 897 species, and 9,090 genetically distinct populations where genetic differentiation ($F_{ST}$) was measured. Caribbean populations were particularly distinguished from North, Central, and South American populations, in having higher differentiation ($F_{ST} = 0.12$ vs. $0.07–0.09$) and lower mean numbers of alleles ($MNa = 4.11$ vs. $4.84–5.54$). While mammalian populations had lower $MNa$ ($4.86$) than anadromous fish, reptiles, amphibians, freshwater fish, and birds ($5.34–7.81$), mean heterozygosity was largely similar across groups ($0.57–0.63$). Mean $F_{ST}$ was consistently lowest in anadromous fishes ($0.06$) and birds ($0.05$) relative to all other groups ($0.09–0.11$). Significant differences in Family/Genera variance among continental regions or taxonomic groups were also observed. MacroPopGen can be used in many future applications including latitudinal analyses, spatial analyses (e.g. central-margin), taxonomic comparisons, regional assessments of anthropogenic impacts on biodiversity, and conservation of wild populations.

Background and Summary

Collating large quantities of data is useful not only for assessing large-scale patterns but also for testing theories, informing conservation initiatives, and providing a valuable resource for future data comparisons. In particular, macro-ecological biodiversity assessments are becoming increasingly popular to identify hotspots of species biodiversity that can inform local management strategies¹⁻⁴. However, populations, not species, are generally recognized as the appropriate scale for the management of sustainable harvesting and protection in endangered species legislation⁶⁻⁸. Nevertheless, population diversity – the number of genetically distinct populations within species – is typically excluded from most biodiversity syntheses and large-scale conservation planning (e.g.¹²⁻¹³). This has consequences when assessing biodiversity loss, as population extinction occurs at a much faster rate than species loss, and as such, a species’ vulnerability could be grossly misrepresented¹⁴.

Molecular markers provide an increasingly effective way to differentiate populations and estimate population diversity¹⁵. One example is the global population diversity estimate based on allozymes and restriction fragment length polymorphisms where authors found on average 220 populations per species and estimated annual loss of 16 million populations, a coarse estimate obtained by dividing the number of sampling locations by the sampling area¹⁰. The collated data from this study was not made publicly available for future usage and is outdated following

¹Department of Biology, Concordia University, 7141 Sherbrooke Street W., Montreal, Quebec, H4B 1R6, Canada. ²School of Environment, University of Auckland, PO Box 92019, Auckland, 1142, New Zealand. ³Golder Associates, 7250, rue du Mile End, 3e étage, Montréal, Québec, H2R 3A4, Canada. ⁴BT Engineering Inc., 100 Craig Henry Drive, Suite 201, Nepean, Ontario, K2G 5W3, Canada. ⁵Department of Biology and Centre for Forest-Interdisciplinary Research, University of Winnipeg, Winnipeg, Manitoba, R3B 2E9, Canada. Correspondence and requests for materials should be addressed to E.R.L. (email: elizabeth-lawrence@outlook.com)
the advancement of genetic tools. No study has formally revisited these concepts since this 1997 study (16–18), indicating the need for collating population information.

Population genetic technologies have seen advances in recent years, switching from allozymes to microsatellites to single nucleotide polymorphisms (SNPs), largely due to the better resolution of within-population variation that more recent technologies provide. Population structure studies and vulnerability assessments have used microsatellites as their molecular marker for the past two decades, yet this wealth of data has not been thoroughly collated, although a few authors have collated related information in the form of microsatellite genetic variation (18), population density estimates (19), and pairwise FST estimates (20). Despite the great degree of data collation across these studies, no work has combined the geo-referencing of population-specific genetic variation, FST measurements, and the number of populations within a species to create a single database across a wide variety of taxa and geographic regions.

Here we provide the first description of the release of the Macro-ecological, Population Genetics Database (MacroPopGen Database) – a database that contains geo-referenced population-specific characteristics based on nuclear DNA microsatellites. It contains information on 897 species from 1,308 studies published between 1994–2017, and 9,090 distinct populations of amphibians, birds, fish [anadromous, brackish, catadromous, or freshwater], mammals, and reptiles, totalling 561,605 genotyped individuals. Every population entry is georeferenced to permit large-scale spatial analyses, opening a variety of opportunities for overlaying microsatellite genetic data with environmental, geographic, or anthropogenic variables. It allows for population diversity and FST to be directly compared to species and genetic diversity (e.g., heterozygosity and mean number of alleles) through mapping applications.

MacroPopGen exemplifies the importance and usefulness of collating population genetic data by standardizing data from > 1000 different studies, allowing for large-scale comparisons and many future applications, including latitudinal analyses, spatial or temporal analyses, taxonomic comparisons and regional assessments of genetic diversity across taxa or in relation to anthropogenic effects. Previous works focusing on older markers have already shown incredible usefulness in testing a variety of genetic and ecological theories (14–18). We provide a baseline database for future works to build from and to compare to, particularly for comparing results to different, newer technologies. We urge future population studies using newer technologies to strive for a similar standardized repository for reporting population-specific statistics.

**Methods**

**Data collection.** To collect population-genetic data from vertebrate populations located in the Americas, we first scanned Web of Science and Google Scholar for relevant articles using key search terms including country of occurrence, species common names, author names, and scientific names in combination with “microsatellite”, “distinct population”, and/or “FST”. A full list of the 1304 key terms and combinations used can be found online (22). We also cross-referenced the list of bird microsatellite papers from Willoughby et al. (9).

Search results with over 1000 hits would be filtered where if two consecutive pages did not yield a relevant result, further pages would not be considered (on average the first 15 pages on Google Scholar would be filtered for relevant articles). This preliminary screening limited results down to 6,297 peer-reviewed studies, technical reports, dissertations and government documents, of which only 1,308 fulfilled our criteria, including 142 of which were obtained from Willoughby et al. (9) bird reference list. Once a study was selected, we extracted where possible: population locality name, latitude-longitude coordinates, average population-specific FST (Wright’s FST or Weir & Cockerham’s unbiased FST estimator ̂{\theta}_ST), population-specific observed and expected heterozygosity averaged across loci (H_e/H_o, respectively), sample size (N), population-specific mean number of alleles per locus (MNA), study-specific corrected allelic richness (AR), and the number of microsatellite loci used in the study. For each population, we also documented the taxonomic group (amphibians, birds, fish [anadromous, brackish, catadromous, or freshwater], mammals, or reptiles), family, genus, species, common name, continent, and country. We chose not to include marine species because microsatellites have typically been unable to detect fine-scale population structure in such species, in contrast to the increased power and resolution of more recent genome-scale analyses for such species (23). Instead we focus on terrestrial and aquatic ecosystems.

All populations were georeferenced in decimal degrees; if coordinates were not provided, they were inferred from the text or maps in a study. To calculate a metric of population-specific FST, we consulted pairwise FST tables and averaged across values that included the focal population, or population group if there was no significance between one or more population pairs. When only a global or regional FST was reported then that value would be used for all populations within the study; such FST values are indicated in the database where applicable.

**Inclusion criteria and assumptions.** A study was retained if two criteria were met: 1) microsatellites were used as molecular markers and 2) genetic differentiation was measured by Weir and Cockerham’s pairwise FST as opposed to other differentiation estimators because of its wide usage. Microsatellites were favoured over other molecular markers (e.g., SNPs, mitochondrial DNA, allozymes, RAPD, etc.) because their polymorphic nature allows them to resolve population structure at fine scales, particularly for closely related populations (25,27). Additionally, microsatellites have higher mutation rates than other markers (25,27) and have been one of the most widely used genetic markers in recent decades (26). Therefore, microsatellites presently provide an abundance of collectable data across taxa relative to more recent molecular developments associated with single nucleotide polymorphisms (SNPs) or barcoding. While barcoding can assess phylogenetic signals across populations and species, microsatellites allow for the comparison of genetic characteristics between populations such as heterozygosity and allelic diversity, which has been noted to indicate levels of inbreeding or adaptive potential (28–32).

Studies were assumed to have used selectively neutral nuclear microsatellite loci unless otherwise indicated because microsatellites are located within non-coding regions of the genome (33) and have relatively fast mutation rates (34). Microsatellite loci are often selected based on their polymorphism due to these faster mutation rates.
rates, causing concern that microsatellites may bias measures of genetic diversity compared to whole DNA sequencing-based measures\(^{34,35}\). Polymorphism bias has also been recognized in studies using other genetic markers such as SNPs\(^{21,34,36,37}\), and will continue to present challenges in genetic studies. An inherent assumption of this database is that ascertainment bias is similar across all studies and taxa, and therefore comparable. Additionally, previous work\(^8\) has concluded that the number of loci and primer type (whether cross-species or focal species) were not important in explaining variability in genetic diversity, an indication that ascertainment bias may not be very significant for large quantities of microsatellite data such as this database. Regardless, we tested ascertainment bias with a subset of the database, as described below.

**Demarcating Populations.** Populations were considered genetically distinct above a threshold \(F_{ST}\) value of 0.02. \(F_{ST}\) was used as the statistical measure of differentiation because of its standardized and common use in the literature for measuring genetic differentiation. The chosen threshold was based on a previous analytical review\(^{38}\), which indicated that genetic differentiation is not negligible if \(F_{ST} \geq 0.05\), but an \(F_{ST}\) value as low as 0.01 can also denote statistically significant differentiation\(^8\). While lower values of \(F_{ST}\) (0.02 to 0.01) are sufficient to show significant genetic differentiation, such values are more relevant for distinguishing specific taxonomic groups, such as marine fish populations which exhibit more gene flow\(^{38,39}\). Freshwater and terrestrial species tend to experience lower rates of gene flow than marine species and therefore an \(F_{ST}\) threshold above 0.01 is more appropriate\(^{35,39}\). To avoid accepting biologically insignificant population differentiation (type I error) or rejecting biologically significant differentiation (type II error) when demarcating populations, we considered the significance of \(F_{ST}\) values where available. We ensured that any pairwise comparisons >0.02 were statistically significant; we also checked significance when \(F_{ST}\) was <0.02 and significance implied two separate populations despite a lower \(F_{ST}\). We also accounted for sample sizes with respect to significant \(F_{ST}\). If sample size was from less (occurred in all cases in this study) and populations were found to be significantly different, the populations were instead grouped as one unless an adequate biological explanation was provided (\(n = 5\)). Likewise, if sample size was very large (e.g. >50) but \(F_{ST}\) was <0.02, consideration would be taken to determine if the populations were significantly different given the statistical support large sample sizes provide (usually given by \(p\)-values in the specific study, \(n = 63\) cases where \(n \geq 50\) but \(F_{ST} \leq 0.02\). Additionally, if multiple studies were conducted in the same location for the same species, data from the most recent study or the one with the most microsatellite loci was used (\(n = 268\) populations were duplicates and removed). When \(F_{ST}\) tables were unclear (e.g. many low \(F_{ST}\) values and no significance given), we considered results from population structure analyses (e.g. STRUCTURE, BAPS, etc.) to make informed decisions about population structure.

**Geographic Breadth.** We also report (i) how differentiated each population is in relation to all other populations it was compared to by calculating the average \(F_{ST}\) between a focal population and all other populations within that study, and (ii) the number of populations included in the calculation as well as the geographic distance or breadth that they span. For example, low \(F_{ST}\) values resulting from only a few sampling locations (e.g. 5) in a small geographic region (e.g. 10 km) may have a different interpretation than low \(F_{ST}\) values across many (e.g. >10) sampling locations in a broad geographic range (e.g. 10,000 km). To estimate the geographic breadth that sampled populations cover, we obtained coordinates for each population including locations that had been combined into one population. These data were put into a separate file that contains 10,921 sampling localities. Next, we used custom code\(^8\) utilizing the R package geosphere\(^{22}\) to back transform estimates.

To compare the degree of variation in each taxonomic or continental group, we calculated the coefficient of variation grouped at the species level for \(H_{O}\), \(M_{NA}\), and \(F_{ST}\). Mixed models using the gamma distribution and random effects of reference, genus, and family were constructed. We then used model selection to see which between taxonomic group or continental region best explained differences between groups. We assessed trends of ascertainment bias related to microsatellite loci development using a subset focusing on North American mammalian data (\(n = 1579\) populations, 73 species)\(^22\). In addition to the number of microsatellite loci, we obtained from 230 mammalian studies the number of species used to develop those loci (\(ranged from 1 to 7\), and whether the species were focal (\(n = 384\)), non-focal (\(n = 545\)), or mixed (\(n = 692\)), as well as information on the senior author's country of affiliation. Using IUCN descriptions for each species, we also determined whether the species was harvested and to what extent (\(no n = 317, low n = 957, high n = 347\)), the species' IUCN status (Least Concern \(n = 1335\), Near Threatened \(n = 45\), Vulnerable \(n = 193\), Endangered \(n = 41\), Critically Endangered \(n = 7\)), whether the species was of conservation concern (\(no n = 561, low n = 211, high n = 849\)), charismatic (\(no n = 495, low n = 189, high n = 937\)), or of economic value (\(no n = 602, low n = 887\), or high \(n = 132\)). Extent of harvesting was determined by the degree of harvesting described in IUCN's "Use and Trade" category: none ("no"), subsistence or local harvesting ("low"), or substantial commercial harvesting.
Table 1. Summary statistics for data collected from microsatellite studies published between 1994 and 2017 broken down by taxonomic group. N = sample size; $H_0$ = observed heterozygosity; MNA = mean number of alleles; SD = standard deviation; SE = standard error. Amph = amphibians; Anad = anadromous fishes; FW = freshwater fishes; Mam = mammals; Rep = reptiles; NOR = North America; CEN = Central America; CAR = Caribbean; SOU = South America. Brackish and catadromous fishes are not shown due to their low number of populations (25 and 33, respectively). *Calculated to account for model structure. See text for details.

("high"). Conservation concern was specified to account for species that may have a lower IUCN rank (e.g. Least Concern, LC) but still have populations at risk or aspects of their habitat at risk (e.g. 563 LC species were still of conservation concern and therefore considered as "low"); this was largely described in IUCN's "Threats" and "Conservation Action" categories. Charisma of species was somewhat subjective as it was determined by how generally well-known the species was, and whether the species may be considered a nuisance which would negatively affect their charisma score (e.g. the coyote is well known but can be considered a pest and as such its score was "low"). Economic value of a species was determined by the "Use and Trade" section, where if the species was commercially harvested it would be considered to have economic value ("high"); if the harvest has declined or is relatively low, a species' economic value was considered as "low".

We tested the fixed effects and interactions among these factors for ascertainment bias as well as the random effects of reference, species, genus, and family. We used GLMMs, using a beta distribution for $H_0$ (R package glmmTMB) and a gamma distribution for MNA (R package lme4). Following Zuur et al. guidelines for forwards and backwards model selection, we used the likelihood ratio test to find significant factors for the $H_0$ and MNA models, respectively.

Code and Data Availability. The data and R code used for the analyses are available from FigShare.

Data Records

Data from the MacroPopGen database is hosted at Figshare and can be downloaded as one XLSX file. It consists of 9,098 rows (distinct populations), and 24 columns. The columns include taxonomic identifiers (family, genus, species, common name), population locality information, and study-specific data (sample size, population-specific $F_{ST}$, observed and expected heterozygosity, mean number of alleles, standardized allelic richness, latitude and longitude coordinates, reference ID, and year).

An additional XLSX file containing the corresponding references for each reference ID, and the list of key terms used in searches is also available on Figshare. Most of the references were published in English, although a minority are in Spanish.

Technical Validation

Geographic and taxonomic bias. Between 1994 and 2017, most population microsatellite data came from species studied in North America (85.1%, Table 1). Fish species were the most represented taxonomic group, making up 44.8% of the database (Table 1). Salmonid species made up 55.9% of fish population data and represented 25.0% of data across all taxa.

When accounting for model structure, mean population genetic diversity differed significantly between some continental regions for $H_0$ and MNA (Fig. 1). Populations of South American species had the lowest $H_0$ while...
Caribbean populations showed significantly lower MNA (Table 1, Fig. 1). Despite some significant differences, the range of mean population genetic diversity metrics among continental regions was limited, between 0.57 and 0.61 for HO, and 4.11 and 5.5 for MNA (Fig. 1). Continental population differences in FST were stronger than for

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**Fig. 1** Coefficient of variation and mean values for observed heterozygosity (H₀), mean number of alleles (MNA), and population-specific FST calculated to account for GLMM structure. Error bars represent standard error. Significant differences between groups indicated by letter grouping where groups sharing the same letter(s) are not significantly different from one another. (a,b) Coefficient of variation calculated across (a) taxonomic groups (circles) and (b) between continental regions (squares). (c–e) Mean (c) FST, (d) H₀, and (e) MNA calculated across taxonomic groups. (f–h) Mean (f) FST, (g) H₀, and (h) MNA calculated between continental regions.

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**Fig. 2** Microsatellite observed heterozygosity (H₀), mean number of alleles (MNA), and population-specific FST averaged across each vertebrate group. Colours indicate the taxonomic group each family or genus belongs to: dark green = amphibians, purple = birds, blue = fish, orange = mammals, light green = reptiles. Error bars represent standard error. (a,c,e) Ho, MNA, and FST are averaged across vertebrate families (n = 195). (b,d,f) Ho, MNA, and FST are averaged across vertebrate genera (n = 480).

Caribbean populations showed significantly lower MNA (Table 1, Fig. 1). Despite some significant differences, the range of mean population genetic diversity metrics among continental regions was limited, between 0.57 and 0.61 for H₀, and 4.11 and 5.5 for MNA (Fig. 1). Continental population differences in FST were stronger than for
genetic diversity metrics, wherein Caribbean populations showed significantly higher population-specific FST, suggestive of less gene flow overall for these populations. This result follows general island-mainland expectations where island populations tend to be more isolated than mainland populations41,42.

Among taxonomic groups, populations of anadromous fish had statistically higher mean genetic diversity (MNA = 7.8), and lower average F ST values (0.06) aside from birds (mean F ST = 0.05) (Fig. 1), consistent with previous work12,13. Mammalian populations also had lower mean MNA than all other groups (Fig. 1). However, there were no significant differences in mean HO between taxonomic groups (Fig. 1).

Variation among taxonomic and continental groups. There were significant differences in the coefficient of variation for H0 among taxonomic groups but not continental regions, with bird species showing the least variation (Fig. 1). There were no significant differences in the coefficient of variation for species MNA across taxonomic groups or continental regions (Fig. 1). For F ST, the only statistical difference was for the coefficient

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**Fig. 3** Observed heterozygosity, mean number of alleles, and number of microsatellite loci for populations of each taxonomic group sampled between the years 1994 to 2017. (a–c) All vertebrate groups together; (d–f) only amphibian species; (g–i) bird species; (j–l) all fish species; (m–o) mammalian species; (p–r) reptile species. Linear models are indicated for significant relationships.

**Fig. 4** Funnel plots for all populations; y axis for both plots is the number of microsatellite loci, and (a) x axis is observed heterozygosity (H0) or (b) mean number of alleles (MNA). Vertical line represents the mean value.
of variation to be larger in North American species relative to species in other regions, i.e. no taxonomic group differences in $F_{ST}$ variance were found (Fig. 1). More variance among taxonomic distinctions was observed when considering within-family and within-genera variance in genetic metrics (Fig. 2). For example, the mean family $H_O$ ranged between 0.07–0.88, while MNA ranged from 1.40–24.97, and mean $F_{ST}$ ranged from 0.0008–0.72; genera averages had a similar range for both metrics.

**Bias with microsatellite loci.** We assessed how genetic diversity and the number of microsatellite loci employed in empirical research has changed over time using linear models (Fig. 3). There has been a significant trend for increasing number of loci per year ($R^2 = 0.07$, $p < 0.001$) as well as a weak increase in genetic diversity with year ($H_O$: $R^2 = 0.009$, $p < 0.001$ and MNA: $R^2 = 0.001$, $p = 0.003$). Additionally, we evaluated bias with respect to the number of microsatellite loci and the degree of genetic variation in $H_O$ and MNA using funnel plots (Fig. 4) and linear models. The plots appear to be largely symmetrical and show little bias with respect to number of loci, indicating the data capture a reasonable degree of genetic variation for the number of loci used. Note that we could not use a formal funnel plot test such as the Egger test because we do not have variance for $H_O$ and MNA for each study. However, the number of microsatellite loci was a significant predictor in linear models for both $H_O$ and MNA ($p < 0.001$ for both), although adjusted $R^2$ values were very small (0.002 and 0.03, respectively).

**Ascertainment bias.** After model selection testing for ascertainment bias with respect to loci type and origin, only the interaction between level of harvesting and conservation concern as well as the random effects of reference, family, and genus were significant for the $H_O$ model (Table 2). For the MNA model, the significant factors only included the interaction between conservation concern and charisma, as well as the random effects for reference and genus. None of the factors associated with microsatellite bias were retained in model selection (i.e. number of species used to derive loci, whether those species were focal, non-focal, or mixed). These results are consistent with previous assessments and indicate that microsatellite loci and loci origin do not significantly affect genetic diversity metrics when analyzed across diverse taxa.

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### Table 2. Summary of model selection results for testing ascertainment bias within $H_O$ and MNA. NSpp: number of species used to derive loci; MsatLoci: total number of microsatellite loci; MsatType: microsatellite type (focal, non-native, native); Harvested: level of harvesting; ConservC: degree of conservation concern; Economic: economic value; Charisma: charisma of focal species; AuthorCountry: senior author's country of residence.

| Model | AIC | DF |
|-------|-----|----|
| $H_O \sim 1 + (1|Reference) + (1|Species) + (1|Genus) + (1|Family)$ | −2196.0 | 6 |
| $H_O \sim MsatType + (1|Reference) + (1|Genus) + (1|Family)$ | −2183.6 | 7 |
| $H_O \sim ConservC + (1|Reference) + (1|Genus) + (1|Family)$ | −2202.3 | 7 |
| $H_O \sim Harvested + ConservC + (1|Reference) + (1|Genus) + (1|Family)$ | −2212.2 | 9 |
| $H_O \sim MsatType + ConservC + (1|Reference) + (1|Genus) + (1|Family)$ | −2198.2 | 9 |
| $H_O \sim Harvested + MsatType + ConservC + (1|Reference) + (1|Genus) + (1|Family)$ | −2215.2 | 13 |
| $H_O \sim MsatType + Harvested + ConservC + (1|Reference) + (1|Genus) + (1|Family)$ | −2214.2 | 14 |
| $H_O \sim NSpp + Harvested + MsatType + ConservC + (1|Reference) + (1|Genus) + (1|Family)$ | −2215.3 | 14 |
| $H_O \sim msat + Harvested + MsatType + ConservC + (1|Reference) + (1|Genus) + (1|Family)$ | −2212.1 | 15 |
| MNA $\sim ConservC:Charisma + (1|Reference) + (1|Genus) + (1|Species)$ | 4015.6 | 13 |
| $MNA \sim NSpp + ConservC:Charisma + (1|Reference) + (1|Genus) + (1|Species)$ | 4016.2 | 14 |
| $MNA \sim NSpp + MsatLoci + ConservC + AuthorCountry + ConservC:Charisma + (1|Reference) + (1|Genus) + (1|Species)$ | 4021.6 | 19 |
| $MNA \sim NSpp + MsatLoci + MsatType + Harvested + ConservC + Economic + Charisma + AuthorCountry + ConservC:Charisma + (1|Reference) + (1|Genus) + (1|Species)$ | 4031.5 | 25 |
| $MNA \sim NSpp + MsatLoci + MsatType + Harvested + ConservC + Economic + Charisma + AuthorCountry + MsatLoci + NSpp:MsatType + MsatLoci:MsatType + Harvested:ConservC + Harvested:conservC + ConservC:Charisma + (1|Reference) + (1|Genus) + (1|Species)$ | 4043.7 | 36 |
| $MNA \sim MsatLoci + MsatType + Harvested + ConservC + Economic + Charisma + AuthorCountry + MsatLoci + NSpp:MsatType + MsatLoci:MsatType + Harvested:ConservC + Harvested:Charisma + ConservC:Charisma + (1|Reference) + (1|Genus) + (1|Species)$ | 4050.9 | 37 |
Author Contributions

E.R.L. initiated the database construction, collected data on North American fish, mammals, amphibians, reptiles, and South American mammals; she also combined all files from other contributors, checked data for consistency, ran statistical analyses, built the reference file, and wrote the paper. J.N.B. collected population data on North American fish, and South American mammals; she also combined all files from other contributors, checked data for consistency, ran statistical analyses, built the reference file, and wrote the paper. T.B. and N.K. collected population data on bird species within the American continent, and provided feedback on manuscript revisions. Z.W. collected population data on North American fish and provided feedback on manuscript revisions. K.M. collected population data on North American mammals, fish, reptiles, and amphibians, and assisted with collating reference information; he also provided feedback on manuscript revisions. J.-M.M. completed statistical modeling, ran statistical analyses, built the reference file, and wrote the paper. J.N.B. collected population data on North American fish, and provided feedback on manuscript revisions. D.J.F. conceived the general idea of the study, collected population data on North American fish, and provided feedback on manuscript revisions.
Additional Information

Competing Interests: The authors declare no competing interests.

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