Genetic evidence supports a distinct lineage of American crocodile (*Crocodylus acutus*) in the Greater Antilles

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ABSTRACT

Four species of true crocodile (genus *Crocodylus*) have been described from the Americas. Three of these crocodile species exhibit non-overlapping distributions—*Crocodylus intermedius* in South America, *C. moreletii* along the Caribbean coast of Mesoamerica, and *C. rhombifer* confined to Cuba. The fourth, *C. acutus*, is narrowly sympatric with each of the other three species. In this study, we sampled 113 crocodiles across *Crocodylus* populations in Cuba, as well as exemplar populations in Belize and Florida (USA), and sequenced three regions of the mitochondrial genome (D-loop, cytochrome b, cytochrome oxidase I; 3,626 base pair long dataset) that overlapped with published data previously collected from Colombia, Jamaica, and the Cayman Islands. Phylogenetic analyses of these data revealed two, paraphyletic lineages of *C. acutus*. One lineage, found in the continental Americas, is the sister taxon to *C. intermedius*, while the Greater Antillean lineage is most closely related to *C. rhombifer*. In addition to the paraphyly of the two *C. acutus* lineages, we recovered a 5.4% estimate of Tamura-Nei genetic divergence between the Antillean and continental clades. The reconstructed paraphyly, distinct phylogenetic affinities and high genetic divergence between Antillean and continental *C. acutus* populations are consistent with interspecific differentiation within the genus and suggest that the current taxon recognized as *C. acutus* is more likely a complex of cryptic species warranting a reassessment of current taxonomy. Moreover, the inclusion, for the first time, of samples from the western population of the American crocodile in Cuba revealed evidence for continental mtDNA haplotypes in the Antilles, suggesting this area may constitute a transition zone between distinct lineages of *C. acutus*. Further study using nuclear character data is warranted to more fully characterize this cryptic diversity, resolve taxonomic uncertainty, and inform conservation planning in this system.
INTRODUCTION
The order Crocodylia is currently comprised of 27 recognized or proposed living species distributed among three families (Alligatoridae, Crocodylidae, and Gavialidae) and nine genera (Alligator, Caiman, Melanosuchus, Paleosuchus, Crocodylus, Mecistops, Osteolaemus, Tomistoma, and Gavialis) representing, along with birds, the unique surviving members of the Archosauria (Grigg & Kirshner, 2015). The genus Crocodylus alone comprises almost 50% (12 out of 27) of all extant crocodylian species and is unique for its global distribution in the tropics and recent diversification (Oaks, 2011). Yet, ambiguity remains regarding the phylogenetic relationships within the genus Crocodylus (Meganathan et al., 2010; Man et al., 2011; Meredith et al., 2011; Hekkala et al., 2011; Oaks, 2011; Shirley et al., 2014). For example, the sister relationships of the four New World Crocodylus species have not been consistent and significantly vary depending on the origin of the species sampled (Milián-García et al., 2015, 2018).

Cuba represents an important area in the distribution of New World Crocodylus, given its geographic proximity to mainland North America (≈150 km from Florida, USA) and Central America (≈210 km from the Yucatán peninsula, Mexico) and the fact that it hosts two Crocodylus species. The Cuban crocodile (Crocodylus rhombifer), endemic to the main island of Cuba, is largely restricted to Zapata Swamp (Matanzas Province) where it is sympatric with C. acutus. In most other coastal areas in Cuba, including Birama Swamp in the southeast and the Guanahacabibes’ peninsula (Pinar del Río province) on the western tip, C. acutus is found in allopatry with no evidence of historical overlap with C. rhombifer (Tabet et al., 2014). Previous studies have indicated that the phylogenetic placement of C. acutus is sensitive to sampling location, with most studies relying solely on individuals sampled from continental populations in Central and/or South America (Meredith et al., 2011; Milián-García et al., 2011, 2015; Oaks, 2011). In these cases, a sister taxon relationship between C. acutus and C. intermedius has been consistently recovered with high support (Brochu & McEachran, 2000; Meganathan et al., 2010; Meredith et al., 2011; Hekkala et al., 2011; Oaks, 2011). However, studies that included Antillean C. acutus sampled in Cuba have revealed a well-supported sister relationship with C. rhombifer, to the exclusion of continental C. acutus that continued to cluster with C. intermedius (Milián-García et al., 2011, 2015, 2018). The observed inconsistency in phylogenetic placement of C. acutus is likely a consequence of excluding intraspecific genetic variation when trying to resolve higher order relationships.

Although Cuban and American crocodiles have been included in previous phylogenetic studies, insufficient sampling has precluded a definitive investigation of cryptic diversity in C. acutus and what impact that might have on phylogenetic reconstruction, taxonomic status, and conservation management. To fill this knowledge gap, here we collected mitochondrial DNA character data for C. acutus sampled throughout Cuba
to characterize patterns of variation and lineage diversity across the island. We combined these data with exemplar sampling and previously collected data from Central America (Colombia, Belize), North America (Florida, USA) and elsewhere in the Greater Antilles (Jamaica and Cayman Islands) to reconstruct phylogenetic relationships across the species distribution and relative to all living Crocodylus species. Reconstructed relationships using newly combined data sets allowed us to test a hypothesis of monophyly of C. acutus as currently described and explore implications for current taxonomic and conservation status in Cuba and elsewhere.

MATERIALS AND METHODS

Sampling

We sampled a total of 65 Crocodylus individuals between the years 2007 and 2014 from four main areas across the Cuban archipelago (Fig. 1). These included three areas where C. acutus is found in allopatri to C. rhombifer (Birama Swamp (n = 27); Province of Cienfuegos (n = 1); Province of Pinar del Río along the Guanahacabibes peninsula (n = 2)) and Zapata Swamp, where the two species are sympatric and known to hybridize (C. rhombifer (n = 31); C. acutus (n = 2); suspected hybrids (n = 2); Milián-García et al., 2015). In addition, we sampled the most western ex situ population for C. acutus at the Sabanalamar captive breeding facility (n = 24), also located in the province of Pinar del Río. The facility was founded in 1986 with a small group of adult breeders caught in the surrounding areas of Pinar del Río, up to five individuals from Lanier Swamp, and a group of neonates from Birama Swamp (R. Rodríguez-Soberón, 2018, personal communication).

All individuals were classified as C. rhombifer, C. acutus, or as suspected hybrids based on 26 external morphological characters (Targarona, 2013) and a piece of a caudal scale was clipped from each animal’s tail for genetic analysis. We collected and transported all samples in accordance with an agreement established between the Faculty of Biology at the University of Havana and the National Enterprise for the Protection of Flora and Fauna in Cuba, and CITES export and import permits, C002135 and E-00134/14, respectively.

We also included tissue samples from continental C. acutus collected from the Florida Everglades, USA (n = 12) and Belize (n = 12). All of these individuals have been previously classified as C. acutus based on morphology and genetics (Hekkala et al., 2015; Rossi, 2016).

Mitochondrial DNA sequencing

Total DNA was isolated for the samples from tail scale tissue preserved in 95% ethanol, using the NucleoSpin® kit for DNA extraction (Macherey-Nagel, Düren, Germany) following manufacturer’s instructions. A total of 716 base pair (bp) segment of mtDNA including the tRNAPro-tRNAPhe-D-loop region (hereafter referred to as D-loop) was amplified as a single fragment for all individuals using primers drL15459 (5′-AGGAAAGCGCTGGCCTTGTAA-3′) and CR2HA (5′-GGGGCCACTAAAAACTGGGGGGA-3′) (Weaver et al., 2008) except for individuals from the Florida Everglades and Belize. Two overlapping fragments of cytochrome b (cytb) covering a total of 1,320 bp were amplified by polymerase chain reactions (PCR) using the primers L14212 (5′-TTG GGC...
TTT AGACCA AGA CC-3′ with CB3H (5′-GGC AAA TAG GAA RTATCA-3′) (Palumbi, 1996), and L14849 (5′-TCCTCCACGAACGCGGAR C-3′) with H15453 (5-CCKTCCAYYTCTGTCTTACAAG-3′) (Milián-García et al., 2011). We also amplified a 665 bp fragment from the 3′ end of cytochrome oxidase I (COI) using the primer pair COIa (5′-AGT ATA AGC GTC TGGGTA GTC-3′) with COIf (5′-CCT GCA GGA GGA GAY CC-3′) (Kessing et al., 1989).

We carried out the PCR on either a BioRad T100™ or an Eppendorf thermal cycler. Each PCR was prepared in a 15 μL total volume, containing: six μL of VWR® Taq
DNA-polymerase (5 U/μL) Master Mix 2X, seven μL of distilled water, 0.5 μL of each primer (10 μmol), and 20–50 ng of DNA. Cycling conditions for the D-loop were as follows: 94 °C (2 min), 33 cycles of 94 °C (30 s), 58 °C (30 s), 72 °C (45 s), and a final extension of 72 °C (7 min). For cytb and COI, cycling conditions were: 94 °C (2 min), 35 cycles of 94 °C (45 s), 48 °C (45 s), 72 °C (90 s), and a final extension of 72 °C (10 min). All PCR products were purified by ExoSAP-IT (USB® Products) and sequenced using Big Dye terminators on an Applied Biosystems 3130xl Genetic Analyzer (Applied Biosystems, Foster City, CA, USA). The regions sequenced per sample are shown in Table S1.

Data analysis
We edited and aligned mitochondrial DNA sequences in Sequencher 5.2.4 (Gene Codes, Ann Arbor, MI, USA). We also calculated haplotypic (h) and nucleotide (π) diversity (Nei, 1987) estimates based on mtDNA sequences after excluding gaps and missing data, as executed in DNASP v5 (Librado & Rozas, 2009). We then generated a haplotype network using statistical parsimony as implemented in TCS (Clement, Posada & Crandall 2000). Haplotype networks were visualized and edited using the web-based program tcsBU (Múrias dos Santos et al., 2016). In addition, we calculated Tamura-Nei genetic distances in MEGA v 7 (Kumar, Stecher & Tamura, 2016).

To examine observed haplotype variation recovered in Cuban Crocodylus within the context of all extant Crocodylus, we analyzed our sequences relative to previously published data at overlapping mtDNA fragments for all living Crocodylus spp. (Table 1). We tested for significant geographic structure among populations using analysis of molecular variance (AMOVA) (Excoffier, Smouse & Quattro, 1992) as implemented in ARLEQUIN v3.5 (Excoffier & Lischer, 2010). We unambiguously aligned sequences using

| Table 1 Origin of previously published reference mtDNA sequences used per Crocodylus spp. |
|-----------------------------------------------|
| Species        | GenBank accession number | Mitochondrial sequences                      | Authors                                      |
| C. rhombifer   | JF502247.1               | Partial mitochondrial genome                  | Meredith et al. (2011)                       |
| C. acutus      | HM636894.1, HQ595003, HQ595041, HQ595050, KF273834, HF273842, HF273844, HF273849, HF568314 | Complete mitochondrial genome, Cyt b, COI, D-loop | Ray et al. (2004); Rodríguez et al., 2011; Man et al. (2011); Milián-Garcia et al. (2011); Bloor, Ibáñez & Viloria-Lagares, (2015) |
| C. moreletii   | HQ585889.1, HQ595007, HQ595045 | Complete mitochondrial genome, Cyt b, COI     | Meganathan et al. (2010); Milián-Garcia et al. (2011) |
| C. intermedius | JF502242.1               | Partial mitochondrial genome                  | Meredith et al. (2011)                       |
| C. niloticus   | JF502245.1               | Partial mitochondrial genome                  | Meredith et al. (2011)                       |
| C. suchus      | JF502244.1               | Partial mitochondrial genome                  | Meredith et al. (2011)                       |
| C. johnsoni    | HM488008.2               | Complete mitochondrial genome                 | Meganathan et al. (2010)                     |
| C. porosus     | DQ273698.1               | Complete mitochondrial genome                 | Li et al. (2007)                             |
| C. palustris   | HM488007.1               | Complete mitochondrial genome                 | Meganathan et al. (2010)                     |
| C. mindorensis | GU144287.1               | Complete mitochondrial genome                 | Feng et al. (2010)                           |
| C. novaeguineae| HM636896.1               | Complete mitochondrial genome                 | Man et al. (2011)                            |
| C. siamensis   | EF581859.1               | Complete mitochondrial genome                 | Srikulnath et al. (2012)                     |
MAFFT as implemented in GENEIOUS 10.2.4 (Biomatters Ltd., Auckland, New Zealand; *Kearse* et al., 2012) with default settings and concatenated the regions following their clockwise order in the mitochondrial genome (cytb-D-loop-COI). We conducted a Bayesian phylogenetic analysis for the combined dataset using MrBayes 3.0b4 (*Ronquist* & *Huelsenbeck*, 2003), assuming a mixed-model of nucleotide substitution using the best-fit model inferred for each partition separately according to the Bayesian information criterion as implemented in jModelTest 2 (*Posada* & *Crandall*, 1998). We used the African Slender-snouted crocodile (*Mecistops cataphractus*; EF551000.1) as an outgroup to root the tree. We ran four simultaneous chains for a chain length of $2 \times 10^6$, each using a random tree as a starting point, the default heating scheme, and a subsampling frequency of 100. The first 2,000 trees were discarded as burn-in samples and the remaining trees were used to construct a majority-rule consensus tree and derive posterior probability values.

**RESULTS**

### Haplotypic variation and population differentiation

#### D-loop

The number of identified mtDNA D-loop haplotypes within Cuban *Crocodylus* ranged from one (Birama Swamp and Zapata Swamp) to four (Pinar del Río). Haplotypic and nucleotide diversities ranged from 0.000–0.634 to 0.000–0.01791, respectively, with the highest levels found in Pinar del Río, particularly the captive population (Table 2). Among the wild caught individuals, the most common haplotype was identical to the $\alpha$ haplotype (*Weaver* et al., 2008; *Milián-García* et al., 2015), which is shared by *C. rhombifer* ($n = 30$) and hybrids ($n = 2$) from Zapata Swamp, as well as for *C. acutus* from Cienfuegos ($n = 1$) and Zapata Swamp ($n = 1$) (Fig. 2). The second most common haplotype was identical to the $\beta$ haplotype (*Weaver* et al., 2008; *Milián-García* et al., 2015), differing by nine steps from the $\alpha$ haplotype, and was found in *C. acutus* from Zapata Swamp ($n = 1$) and Birama Swamp ($n = 19$). The third most common haplotype was $\delta$ (identical to the Ca1 haplotype in *Rodriguez* et al. (2011); GenBank accession no. MH753361), which has been considered the most frequent haplotype in North and Central America (Fig. 2). It differs by 23 steps from the $\alpha$ haplotype and was found in all wild *C. acutus* from Pinar del Río ($n = 2$).

#### Cytochrome oxidase I

The COI fragment resolved into four haplotypes (Fig. 2) with an overall haplotype and nucleotide diversity per population ranging from 0.000–0.558 to 0.000–0.02420, respectively (Table 2). Similar to patterns at the D-loop, the most common haplotype was identical to the $\alpha$ haplotype and was shared among *C. rhombifer* ($n = 25$), *C. acutus* ($n = 1$), and hybrids ($n = 2$) from Zapata Swamp as well as for *C. acutus* from Cienfuegos ($n = 1$) (Fig. 2). The second most common haplotype $\delta$ (GenBank accession no. MH758763) was shared for *C. acutus* individuals from Pinar del Río ($n = 2$), Belize ($n = 12$), and Florida ($n = 11$). The third most common haplotype was identical to the $\beta$ haplotype and was shared among *C. acutus* from Birama ($n = 7$), and
Zapata Swamp \((n = 1)\). The fourth mtDNA COI haplotype was unique to \textit{C. acutus} from Birama \((n = 1)\) and differed by just one step from the \(\alpha\) and \(\beta\) haplotypes (GenBank accession no. MH758764).

\textbf{Cytochrome b}

Three mtDNA cytb haplotypes were recovered among the wild individuals sampled from Pinar del Río, Birama Swamp, Zapata Swamp, Belize, and Florida. Following the same pattern as for the D-loop, the most common haplotype was identical to the \(\alpha\) haplotype and was shared among \textit{C. rhombifer} \((n = 31)\) and hybrids \((n = 2)\) from Zapata Swamp, as well as for \textit{C. acutus} from Cienfuegos \((n = 1)\) and Zapata Swamp \((n = 1)\) (Fig. 2). The second most common haplotype, \(\delta\), was shared for \textit{C. acutus} from Pinar del Río \((n = 2)\), Belize \((n = 12)\), and Florida \((n = 12)\) differing by 37 steps from the \(\alpha\) haplotype (GenBank accession no. MH758762). The third haplotype was identical to the \(\beta\) haplotype and was shared among \textit{C. acutus} from Birama \((n = 26)\) and Zapata Swamps \((n = 1)\), differing by six steps from the \(\alpha\) haplotype. Overall haplotypic and nucleotide diversities ranged from 0.000–0.583 to 0.000–0.02948, respectively (Table 2).

The fully concatenated sequences revealed unique haplotypes for all Cuban samples for which all regions were successfully sequenced (Table S1). The AMOVA using the concatenated mtDNA sequences recovered significant levels of genetic structure, with a highest fraction distributed among \((>99\%)\), rather than within \((<1\%)\), the populations of Cuban and American crocodiles studied \((P < 0.0001)\).
Captive population

Thirteen individuals with the D-loop α haplotype were identified in captivity, while two possessed the D-loop β haplotype and seven presented the D-loop δ haplotype. The other two individuals presented a fourth new haplotype differing by only one mutational step away from the δ haplotype (GenBank accession no. MH719199). When analyzing the COI fragment, captive individuals were distributed into three haplotypes. Twelve individuals presented the COI α haplotype, one the COI β haplotype and nine individuals shared the continental haplotype (δ). When analyzing the cytb, three haplotypes were recovered for the 24 captive individuals. A total of 13 individuals presented the cytb α haplotype, two exhibited the cytb β haplotype, while nine shared the same haplotype (δ) showed by the continental individuals.

Phylogenetic analysis

The analysis of the concatenated mtDNA regions for the wild-caught samples revealed that Cuban Crocodylus are split into three independent clades. Two of these correspond to the previous clades recovered for Antillean C. acutus (central and eastern Cuba, Jamaica, Cayman Islands) and C. rhombifer (Fig. 3) revealing a well-supported split (posterior probability = 1), and a sister relationship with C. moreletti. The third clade
grouped wild *C. acutus* from western Cuba with *C. acutus* from continental South, Central, and North America, sister to *C. intermedius*. Tamura-Nei genetic distances revealed shallow divergence between Antillean *C. acutus* and *C. rhombifer* (0.4%), substantially lower than the value resolved between western and eastern *C. acutus* populations within Cuba (5.4%).

**DISCUSSION**

Recent studies have uncovered cryptic diversity across a diverse group of crocodylians (e.g., *C. niloticus*, *C. suchus*, *M. cataphractus*, *Osteolaemus*; *McAliley et al.*, 2006; *Eaton et al.*, 2009; *Hekkala et al.*, 2011; *Oaks*, 2011; *Shirley et al.*, 2014; *Cunningham, Shirley & Hekkala*, 2016). Consistent with previous findings (*Milián-García et al.*, 2011,
2015, 2018), our results revealed a high degree of genetic differentiation within the American crocodile, *C. acutus*. Based on the most expansive geographic and mtDNA character sampling to date, we reconstructed *C. acutus* as paraphyletic, exhibiting two distinct lineages composed of Antillean and continental populations with high genetic divergence (5.4%). Moreover, these two lineages are more closely related to other species of crocodiles than they are to each other. Antillean *C. acutus* exhibits a close affiliation with *C. rhombifer* in a clade sister to *C. moreletii*. Conversely, continental *C. acutus* is most closely related to *C. intermedius*.

At a finer-scale, we found significant differentiation between haplotypes from western Cuba and the rest of the sampled populations, including Zapata Swamp and Bimara Swamp. Interestingly, *C. acutus* haplotypes recovered from Pinar del Río province do not exhibit significant differentiation from continental *C. acutus*, and have individuals placed in the same clade (Fig. 3). This result further suggests that the western tip of Cuba—which is closer to the Yucatan peninsula than it is to Zapata Swamp—may either be a transition zone or an area of recent colonization from continental *C. acutus*. The fact that individuals sampled within the most western on-island *C. acutus* captive population share the continental haplotype is consistent with the hypothesis of a transition zone in the area. Most founders at the Sabanalamar captive population were captured locally back in the 1980s, while others were transported from Bimara and Lanier Swamps (R. Rodríguez-Soberón, 2018, personal communication). The latter helps account for the admixture exhibited within the captive population. Our results also demonstrate that human-mediated translocation should no longer be a supported practice as it produces a mix of highly divergent genetic lineages.

The close genetic relationship between *C. rhombifer* and Antillean *C. acutus* based on mtDNA has been previously hypothesized to be a consequence of mitochondrial capture through hybridization, and *C. rhombifer* and *C. acutus* have been documented to hybridize in Zapata Swamp (Milián-García et al., 2015). Although karyological differences exist between *C. rhombifer* (2n = 30) and *C. acutus* (2n = 32) (Cohen & Gans, 1970), there is evidence of fertile hybrids (at least F1) between the species (E. Pérez-Fleitas, 2018, personal communication). However, most *C. acutus* populations in the Antilles have never been recorded to overlap in distribution with the Cuban crocodile (Tabet, 2010; Tabet et al., 2014; Brochu & Jiménez-Vázquez, 2014). For example, the American crocodile population in Bimara Swamp, recognized as the second most extensive wetland in the insular Caribbean, is considered to be the largest and most protected population for the species in the Antilles (Tabet, 2010; Tabet et al., 2014). As such, mitochondrial capture through hybridization would be a feasible hypothesis to explain genetic similarity between *C. rhombifer* and Antillean *C. acutus* if the event occurred before the dispersion of *C. acutus* around the Cuban archipelago and the rest of the Greater Antilles. It would also imply that Antillean *C. acutus* with *C. rhombifer* captured-mitochondria would have higher fitness than the continental *C. acutus*, given the absence of the latter across the Antilles.

An alternative explanation for this pattern could be one of very recent divergence of the Antillean lineages from continental congeners coupled with the slow mtDNA mutation rate in crocodylians (Hekkala et al., 2011; Oaks, 2011). This hypothesis is consistent
with the observation of low levels of genetic differentiation between *C. rhombifer* and Antillean *C. acutus* (Milián-García et al., 2011, 2015, 2018), albeit at a lower level than the average genetic distance between conspecifics within *Crocodylus* (~1%; Hekkala et al., 2011). These findings, in tandem with observed morphological, ecological, and ethological characteristics (Cuvier, 1807; Targarona, 2013), suggest that *C. rhombifer* constitutes a unique lineage, but further study is warranted (see below).

These patterns reconstructed from mtDNA character data represent clear challenges for the current taxonomy of *C. acutus*. Combined with the results of previous studies (Weaver et al., 2008; Milián-García et al., 2011, 2015, 2018), we found evidence for a cryptic lineage of American crocodile currently found in Cuba and other localities in the Antilles. Moreover, the paraphyletic lineages of *C. acutus* show a genetic distance (5.4%) similar to interspecific comparisons within *Crocodylus* and exhibit closer phylogenetic affinities with other species than each other, suggesting a formal taxonomic reassessment may be in order. Yet it is imperative to highlight that this work was limited to insights from mitochondrial DNA character data; additional analyses using nuclear data are essential for resolving taxonomic uncertainty in this system and to more directly test phylogeographic hypotheses. That said, our results do have immediate relevance for conservation, as the cryptic diversity uncovered here and elsewhere is not currently accounted for in status assessments and management plans for new world *Crocodylus*.

**CONCLUSIONS**

Overall, the reconstructed paraphyly, distinct phylogenetic affinities and high genetic divergence between Antillean and continental *C. acutus* populations reinforce the need for a formal taxonomic reassessment. Such an endeavor would benefit from a nuclear genome-wide study of *Crocodylus*, as has been demonstrated in other systems that exhibit a complex history and high prevalence of cryptic diversity (e.g., Galapagos tortoise, *Chelonoidis* sp.; Miller et al., 2018). This work benefited from the most comprehensive geographic and character data sampling conducted to date for *C. acutus* and other congenerics. Importantly, the inclusion, for the first time, of samples from the western population of the American crocodile in Cuba revealed evidence for continental mtDNA haplotypes in the Antilles, suggesting this area may constitute a transition zone between distinct lineages of *C. acutus*. More broadly, detection of cryptic diversity has important implications for conservation. Although *C. acutus* population sizes have been increasing since CITES protections have been in place (Platt, Rainwater & Nichols, 2004), assessments to date have not evaluated the status of Antillean populations as potentially constituting an independently evolving lineage of New World crocodiles. Likewise, cryptic diversity may be reflective of ongoing evolutionary processes as closely related species and populations adapt to changing environments (Bickford et al., 2007), punctuating the need for formal characterization to resolve taxonomic uncertainty and inform conservation planning (Struck et al., 2018), in this case for the American and Cuban crocodiles.
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Competing Interests
The authors declare that they have no competing interests.

Author Contributions
- Yoamel Milián-García conceived and designed the experiments, performed the experiments, analyzed the data, contributed reagents/materials/analysis tools, prepared figures and/or tables, authored or reviewed drafts of the paper, approved the final draft.
Michael A. Russello conceived and designed the experiments, analyzed the data, contributed reagents/materials/analysis tools, prepared figures and/or tables, authored or reviewed drafts of the paper, approved the final draft.

Jessica Castellanos-Labarcrena analyzed the data, prepared figures and/or tables, authored or reviewed drafts of the paper, approved the final draft.

Martin Cichon performed the experiments, analyzed the data, prepared figures and/or tables, authored or reviewed drafts of the paper, approved the final draft.

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Georgina Espinosa contributed reagents/materials/analysis tools, authored or reviewed drafts of the paper, approved the final draft.

Natalia Rossi contributed reagents/materials/analysis tools, authored or reviewed drafts of the paper, approved the final draft.

Frank Mazzotti contributed reagents/materials/analysis tools, authored or reviewed drafts of the paper, approved the final draft.

Evon Hekkala analyzed the data, contributed reagents/materials/analysis tools, prepared figures and/or tables, authored or reviewed drafts of the paper, approved the final draft.

George Amato analyzed the data, contributed reagents/materials/analysis tools, prepared figures and/or tables, authored or reviewed drafts of the paper, approved the final draft.

Axel Janke conceived and designed the experiments, analyzed the data, contributed reagents/materials/analysis tools, prepared figures and/or tables, authored or reviewed drafts of the paper, approved the final draft.

Field Study Permissions
The following information was supplied relating to field study approvals (i.e., approving body and any reference numbers):

All samples were collected and transported in accordance with an agreement established between the Faculty of Biology at the University of Havana and the National Enterprise for the Protection of Flora and Fauna in Cuba, and CITES export and import permits, C002135 and E-00134/14, respectively.

Data Availability
The following information was supplied regarding data availability:

GenBank accession no. MH719199, MH753361, MH758762–MH758764.

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