Molecular Ecology Resources (2010) 10, 449-458

DNA BARCODING

DNA barcodes provide new evidence of a recent radiation in the genus *Sporophila* (Aves: Passeriformes)

LEONARDO CAMPAGNA,* DARÍO A. LIJTMAER,* KEVIN C. R. KERR,† ANA S. BARREIRA,* PAUL D. N. HEBERT,† STEPHEN C. LOUGHEED‡ and PABLO L. TUBARO*

*División de Ornitología, Museo Argentino de Ciencias Naturales ''Bernardino Rivadavia'', Av. Ángel Gallardo 470, Ciudad de Buenos Aires, C1405DJR Buenos Aires, Argentina, †Department of Integrative Biology, Biodiversity Institute of Ontario, University of Guelph, 50 Stone Road East, Guelph, ON, Canada N1G 2W1, ‡Department of Biology, Queen's University, 116 Barrie Street, Kingston, ON, Canada K7L 3N6

Abstract

The capuchinos are a group of birds in the genus Sporophila that has apparently radiated recently, as evidenced by their lack of mitochondrial genetic diversity. We obtained cytochrome c oxidase I (COI) sequences (or DNA barcodes) for the 11 species of the group and various outgroups. We compared the patterns of COI variability of the capuchinos with those of the largest barcode data set from neotropical birds currently available (500 species representing 51% of avian richness in Argentina), and subjected COI sequences to neighbourjoining, maximum parsimony and Bayesian phylogenetic analyses as well as statistical parsimony network analysis. A clade within the capuchinos, the southern capuchinos, showed higher intraspecific and lower interspecific divergence than the remaining Argentine species. As most of the southern capuchinos shared COI haplotypes and pairwise distances within species were in many cases higher than distances between them, the phylogenetic affinities within the group remained unresolved. The observed genetic pattern is consistent with both incomplete lineage sorting and gene flow between species. The southern capuchinos constitute the only large group of species among the neotropical birds barcoded so far that are inseparable when using DNA barcodes, and one of few multispecies avian groups known to lack reciprocal monophyly. Extending the analysis to rapidly evolving nuclear and mitochondrial markers will be crucial to understanding this radiation. Apart from giving insights into the evolution of the capuchinos, this study shows how DNA barcoding can rapidly flag species or groups of species worthy of deeper study.

Keywords: capuchinos, cytochrome c oxidase I, DNA barcodes, mitochondrial DNA, Sporophila

Received 22 June 2009; accepted 23 October 2009

Introduction

The avian genus *Sporophila* comprises granivorous species that occur in open and semi-open habitats from the southern USA to southern South America (Meyer de Schauensee 1952). *Sporophila* species are characterized by small size (10–12 cm), stubby bills and strong sexual dimorphism, with colourful and boldly patterned males and drab females (Ridgely & Tudor 1989). The taxonomy of the genus is in flux, with some forms considered dis-

Correspondence: Leonardo Campagna, Fax: +54 11 4982 0306; E-mail: leocampagna@gmail.com tinct species by some researchers, but only subspecies or local variants by others. Moreover, poorly known forms, such as *S. insulata, S. melanops* and *S. zelichi*, may simply be hybrids or aberrant individuals of other better known species (Ridgely & Tudor 1989). Consequently, the number of species included in the genus varies between 28 and 32 (Hellmayr 1938; Meyer de Schauensee 1952; Ridgely & Tudor 1989; Sibley & Monroe 1990; Howard & Moore 1991).

The capuchinos include 11 *Sporophila* species that are smaller than the other members of the genus and characterized by cinnamon-based plumage colour patterns (Ridgely & Tudor 1989). The species of this group show

450 DNA BARCODING

little differentiation in size and shape, which makes females challenging to identify, whereas males differ considerably in adult plumage and vocalizations. Most capuchino species are highly sympatric with one or more other species of the group and many are rare, having limited ranges with populations in decline due to habitat loss and trapping for the pet trade (BirdLife International 2009, see Table 1). The majority of the capuchinos are seasonal migrants, but little is known about the location of the winter grounds. When not breeding, they are commonly seen in mixed flocks showing similar foraging behaviour (Ridgely & Tudor 1989; Silva 1999). A recent phylogenetic analysis performed by our group (Lijtmaer et al. 2004) included 10 of the 11 species (missing S. nigrorufa) and suggested that the capuchinos are monophyletic and further divided into two clades: northern capuchinos (S. castaneiventris and S. minuta) and southern capuchinos (S. bouvreuil, S. cinnamomea, S. hypochroma, S. hypoxantha, S. melanogaster, S. palustris, S. ruficollis and S. zelichi). These are found predominantly north and south of the Amazon River respectively. The phylogenetic relationships among the southern capuchinos

were unresolved, mainly due to the presence of extremely low interspecific sequence divergence and apparent lack of reciprocal monophyly among species.

A short, standardized fragment of the mitochondrial gene cytochrome c oxidase I (COI) has recently been proposed as a tool for species identification; a library of COI sequences from taxonomically verified voucher specimens, constituting 'DNA barcodes', serves as species identifiers (Hebert et al. 2003). This approach to species identification assumes that intraspecific variation in COI is usually lower than interspecific differences (Hebert et al. 2003). COI surveys in several animal groups have already demonstrated high success rates in species identification (e.g. Lepidoptera, Hajibabaei et al. 2006; amphibians, Smith et al. 2008; fish, Ward et al. 2005). Prior studies have shown the effectiveness of COI in identifying bird species as well. In one of the most comprehensive regional analysis performed on vertebrates to date, Kerr et al. (2007) showed that around 94% of North American bird species have COI clusters that do not overlap with those of other species, allowing their unequivocal identification. The remaining 6% included a

 Table 1
 Scientific names, conservation status, estimated number of individuals, breeding habitat and approximate geographic ranges of the species included in the capuchino group (Ridgely & Tudor 1989; BirdLife International 2009)

Scientific name*	Conservation status and estimated number of individuals†	Breeding habitat	Approximate geographic range‡
S. bouvreuil§	LC, unknown¶	Tall grass savannahs	Locally in E and S Brazil; E Paraguay; NE Argentina; S Suriname
S. castaneiventris	LC, unknown¶	Grassy and shrubby clearings, floating vegetation of marshes, lake and river margins	E Colombia; SW Venezuela; E Ecuador; E Peru; N Bolivia; Amazonian Brazil; Guianas
S. cinnamomea	VU, 2500-10 000	Tall grasslands, near marshes	Locally in S Brazil, E Paraguay; NE Argentina; W and extreme SE Uruguay
S. hypochroma	NT, unknown	Tall grasslands near marshes	Very locally in N and E Bolivia; SW Brazil; E and SE Paraguay; NE Argentina; Uruguay
S. hypoxantha	LC, unknown¶	Tall grasslands near marshes	N and E Bolivia; S Brazil; Paraguay; N Argentina
S. melanogaster	NT, unknown	Tall grasslands near marshes and scrub	SE Brazil
S. minuta	LC, 500 000-5 000 000	Tall grass savannahs near water	Colombia; NW Ecuador; Venezuela; Guianas; lower Amazon Brazil; Mexico to Panama
S. nigrorufa	VU, 1000–2500	Tall grasslands near water	E Bolivia; extreme SW Brazil
S. palustris	EN, 1000–2500	Inundated grasslands and marshes	Very locally in S Brazil; SE and central Paraguay; Uruguay; NE Argentina
S. ruficollis	NT, unknown	Grasslands and dry savannah	NE Bolivia; Paraguay; S Brazil; N Uruguay; N Argentina
S. zelichi	CR, 50–250	Tall grass in flooded areas	NE Argentina; S Brazil; E Paraguay; SE Uruguay

*S., Sporophila.

+CR, critically endangered; EN, endangered; LC, least concern; NT, near threatened; VU, vulnerable.

‡N, north; S, south; E, east; W, west; NE, north-east; NW, north-west; SE, south-east; SW, south-west.

SThis species is polytypic, including S. bouvrevil bouvrevil and S. bouvrevil pileata. The former is found in the northern portion of the

species distribution and is rufous below, while the latter is found in the southern portion and is white below.

¶Although the population has not been quantified it is thought to be larger than 10 000 mature individuals.

few pairs or trios of taxa and one relatively large group of eight species (the large white-headed gulls of the *Larus argentatus–fuscus* species complex) that could not be separated based on COI alone because interspecific variation was indistinguishable from variation within single species. Other studies have shown specifically that DNA barcodes can separate and identify sister or closely related species in diverse avian orders (Vilaça *et al.* 2006; Chaves *et al.* 2008; Tavares & Baker 2008).

As part of an ongoing project to barcode all the birds of Argentina (Kerr *et al.* 2009), we obtained the COI sequences for most species of southern capuchinos and subsequently extended our sampling to include all the members of the capuchino group and various individuals within species. Given the high success of DNA barcodes in species identification in animals in general and birds in particular, the objective of this study was to use COI sequences to separate the capuchino species and to gain further insights into their phylogenetic relationships.

Materials and methods

Data set

Most tissue samples used in this study were either collected on field trips organized by the Ornithology Division of the Museo Argentino de Ciencias Naturales 'Bernardino Rivadavia' (MACN) between 2006 and 2007 or loaned by other institutions (see Table 2). A few samples were obtained from other sources, such as donated specimens or birds confiscated from illegal traders and deposited at the MACN. In most cases, a traditional voucher consisting of a study skin, skeleton or specimen in ethanol is deposited either at the MACN or in another institution. In the case of blood samples, digital pictures were taken of the bird before release, providing an electronic voucher.

All 11 species of capuchinos are represented in our data set, and we used four other sympatric species as outgroups: S. collaris, S. caerulescens, S. leucoptera and Volatinia jacarina (all of which lie outside of our ingroup taxa; Lijtmaer et al. 2004). When available, we included multiple individuals per species (range 1-9) and from as many localities of their geographic distribution as possible (range 1-5). This is particularly relevant in the case of recently diverged species like the capuchinos, where incomplete lineage sorting is expected (Maddison & Knowles 2006). In total, our data included 38 southern capuchinos, 10 northern capuchinos and 13 outgroup specimens (Table 2). Both colour patterns and geographic range were used to identify capuchino species. The majority of the samples were taken from males in adult plumage which allowed unequivocal identification. In the case of the southern capuchinos, where most females

are hard to identify using plumage traits, samples belonging to females or individuals of unknown sex (7 of 38) were clearly identified using information from distribution and the species present in the locality of capture.

DNA extraction, COI amplification and sequencing

DNA sources for this study included frozen pectoral muscle, liver, heart, blood and, in the case of S. nigrorufa, toe pads from a museum study skin. Approximately half of the samples were processed at the Canadian Centre for DNA Barcoding (Guelph, ON, Canada) following the extraction procedures described by Kerr et al. (2009). The remaining DNA extracts were obtained using the GenElute mammalian genomic DNA miniprep kit (Sigma-Aldrich) or following the procedures described by Miller et al. (1988). Polymerase chain reactions (PCRs) utilized the primer pair BirdF1 (5'-TTCTCCAACCACAAAGAC ATTGGCAC-3') and COIbirdR2 (5'-ACGTGGGAGATA ATTCCAAATCCTGG-3') to obtain 694 base pairs (bp) of the COI. PCRs were run under the following thermal cycle profile: 1 min at 94 °C followed by six cycles of 1 min at 94 °C, 1.5 min at 45 °C and 1.5 min at 72 °C, followed in turn by 35 cycles of 1 min at 94 °C, 1.5 min at 55 °C and 1.5 min at 72 °C, and finally 5 min at 72 °C (Kerr et al. 2009). As the S. nigrorufa sample was suspected to contain degraded DNA (it was taken from a museum study skin collected in 1885), internal primers were used in conjunction with those above to obtain two shorter, overlapping sequences. These primers were AvMiR1 (5'-ACTGAAGCTCCGGCATGGGC-3') and AvMiF1 (5'-CCCCCGACATAGCATTCC-3') (Kerr et al. 2009), and the same thermal cycle profile was used. Although the sequence obtained was shorter (462 bp) because only the amplification with the AvMiF1/BirdR1 primer pair was successful, most of the variable sites were recovered. PCR products were visualized on a 2% agarose gel and bi-directionally sequenced on an ABI 3730XL DNA Analyzer (Applied Biosystems).

We deposited all sequences in GenBank (for accession numbers, see Table 2). Approximately half of them were deposited as part of a broader study of the Argentine avifauna (Kerr *et al.* 2009), while the remaining sequences were submitted separately.

Genetic variability and phylogenetic analyses

To compare COI variation patterns of the southern capuchinos with other neotropical birds, we used information from the project 'Birds of Argentina – Phase I' at http:// www.barcodinglife.org because this is the largest data set of neotropical birds barcoded thus far (1594 individuals belonging to 500 species; Kerr *et al.* 2009). Kimura 2-parameter (K2P) distances (Kimura 1980) were

452 DNA BARCODING

Table 2 List of the specimens included in this stu
--

Specimen*	Locality	Sex	Type of samplet	GenBank acc. nos.	Museum collection nos.‡	Haplotype
S. caerulescens 1	Estero Poí, Formosa, Argentina	Male	MLHS	FJ028301	MACN-Or-70897	A1
S. caerulescens 2	Estero Poí, Formosa, Argentina	Male	MLHS	FJ028302	MACN-Or-70916	A2
S. caerulescens 3	Estero Poí, Formosa, Argentina	Male	MLHS	FJ028303	MACN-Or-70937	A3
S. caerulescens 4	San Cayetano, Corrientes, Argentina	Male	MLHS	FJ028298	MACN-Or-69625	A4
S. caerulescens 5	San Cayetano, Corrientes, Argentina	Male	MLHS	FJ028299	MACN-Or-69704	A5
S. caerulescens 6	San Cayetano, Corrientes, Argentina	Male	MLHS	FJ028300	MACN-Or-69771	A6
S. collaris 1	Estero Poí, Formosa, Argentina	Male	MLHS	FJ028307	MACN-Or-70907	B1
S. collaris 2	Estero Poí, Formosa, Argentina	Male	MLHS	FJ028310	MACN-Or-70890	B1
S. collaris 3	Estero Valenzuela, Corrientes, Argentina	Male	MLHS	FJ028306	MACN-Or-69888	B2
S. collaris 4	Estero Valenzuela, Corrientes, Argentina	Male	MLHS	FJ028309	MACN-Or-69889	B1
S. collaris 5	Estero Valenzuela, Corrientes, Argentina	Male	MLHS	FJ028308	MACN-Or-69891	B1
S. leucoptera 1	Estero Poí, Formosa, Argentina	Male	MLHS	FJ028316	MACN-Or-70917	C1
S. castaneiventris 1	Isla Sharamentsa, Pastaza, Ecuador	Unknown	MLHS	GU070583	ZMUC 123784	D1
S. castaneiventris 2	Isla Pasto, Loreto, Peru	Male	MLHS	GU070584	LSUMZ 120308	D2
S. castaneiventris 3	Isla Pasto, Loreto, Peru	Female	MLHS	GU070585	LSUMZ 120303	D3
S. minuta 1	Berbice, Guyana	Male	MLHS	GU070586	USNM 621081	E1
S. minuta 2	Wiwitau Mount., Guyana	Male	MLHS	GU070587	USNM 622227	E2
S. minuta 3	Livestock Research Station, Trinidad	Unknown	BS	GU070588	STRI TR-SMI1	E1
S. minuta 4	Guaraunos, Venezuela	Unknown	BS	GU070589	STRI VE-SMI18	E1
S. minuta 5	Guaraunos, Venezuela	Unknown	BS	GU070590	STRI VE-SMI19	E1
S. minuta 6	Guaraunos, Venezuela	Unknown	BS	GU070591	STRI VE-SMI8	E3
S. minuta 7	Guaraunos, Venezuela	Unknown	BS	GU070592	STRI VE-SMI9	E1
S. bouvreuil 1§**	Unknown	Male	MLHS	GU070593	ZMUC 130533	F1
S. bouvreuil 2	San Luis Nat. Park, Concepción, Paraguay	Male	MLHS	GU070594	KUNHM 88403	F2
S. bouvreuil 3	San Rafael Nat. Park, Itapuá, Paraguay	Male	MLHS	GU070595	KUNHM 91411	F3
S. bouvreuil 4	San Rafael Nat. Park, Itapuá, Paraguay	Male	MLHS	GU070596	KUNHM 3664¶	F4
S. bouvreuil 5	San Rafael Nat. Park, Itapuá, Paraguay	Male	MLHS	GU070597	KUNHM 91403	F5
S. bouvreuil 6	San Rafael Nat. Park, Itapuá, Paraguay	Male	MLHS	GU070598	KUNHM 3691¶	F5
S. bouvreuil 7	San Rafael Nat. Park, Itapuá, Paraguay	Male	MLHS	GU070599	KUNHM 91413	F5
S. cinnamomea 1	Iberá, Corrientes, Argentina	Male	BS	FI028305	MACN-Or-ct 3121¶	F6
S. cinnamomea ?	Gualeguaychú, Entre Ríos, Argentina	Male	BS	FI028304	MACN-Or-ct 3122¶	F5
S. hypochroma 1	Gualeguaychú, Entre Ríos, Argentina	Male	BS	FI028311	MACN-Or-ct 3131¶	F5
S. hypoxantha 1	El Bagual, Formosa, Argentina	Unknown	MLHS	FI028312	MACN-Or-70846	F5
S hypoxantha ?	El Bagual Formosa Argentina	Male	MLHS	FI028313	MACN-0r-70957	F7
S. hypoxantha 3	El Bagual Formosa Argentina	Male	MLHS	FI028314	MACN-0r-70958	F5
S. hypoxantha 4	Estero Catalina Formosa Argentina	Male	MLHS	FI028315	MACN-0r-70977	F8
S. hypoxantha 5	Fl Bagual Formosa Argentina	Male	MLHS	GU070600	MACN-0r-70962	F5
S. hypoxantha 6	El Bagual Formosa, Argentina	Male	MLHS	GU070601	MACN-01-70963	F5
S. hypoxantha 7	El Bagual Formosa Argentina	Male	RS	GU070602	MACN-Or-ct 3097¶	F9
S. hypoxantha 8	El Bagual Formosa, Argentina	Male	BS	GU070602	MACN-Or-ct 3098¶	F10
S. hypoxantha 9	Volasco Santa Cruz Bolivia	Malo	MIHS	CU070604	I SUM7 151408	F10
S. nypoxuninu 9 S. melanogaster 1	Bom Jesus Río Grande do Sul Brasil	Female	RS	GU070604	MCP 2072	F11
S. melanogaster 7	Bom Jesus, Río Grande do Sul, Brasil	Female	BS	GU070606	MCP 2072	F12
S. melanogaster 2	Bom Josus, Río Grande do Sul, Brasil	Malo	BS	CU070607	MCP 2074	F10
S. melunoguster S	Bom Josus, Rio Grande do Sul, Brasil	Male	D3 BC	GU070607	MCP 2074	F10 F12
S. melanogaster 5	Bom Josus, Rio Grande do Sul, Brasil	Male	BC	GU070608	MCP 2075	F13 E14
S. melunoguster S	Bom Josus, Nio Grande do Sul, Brasil	Formala	DO	GU070609	MCF 2070	Г14 Г15
S. melunoguster 6	Bom Josus, Nio Grande do Sul, Brasil	Female	DO	GU070610	MCF 2077	F13 F16
S. meiunogusier 7	Mata Crassa Brasil	Francis	05	GU070611	MCF 2070	F10 F17
S. nigroruju 1	Mato Grosso, brasil	Female	55 DC	GU070612	DIVINEI 1885.2.10.119	F1/ E10
5. puiustris 1	Gualeguaycnu, Entre Klos, Argentina	Mala	DO PC	FJU28317	MACN OF 71052	F10 F10
5. puiustris 2	Gualeguaycnu, Entre Klos, Argentina	iviale	DO DC	FJU28318	MACN OF 1052	F19 F20
5. paiustris 3	Ibera, Corrientes, Argentina	Male	B5 DC	EU906931	MACN-Or-ct 3118¶	F2U F17
5. ruficollis 1	Gualeguaychu, Entre Kios, Argentina	Male	D5 DC	EU906932	MACN-Or-ct 3128¶	F1/
5. ruficollis 2	Gualeguaychu, Entre Rios, Argentina	Male	BS	EU906933	MACN-Or-ct 3129¶	F5
5. ruficollis 3§	Argentina	Male	BS	FJ028321	MACN-Or-ct 3130¶	F5
S. ruficollis 4	San Cayetano, Corrientes, Argentina	Male	MLHS	FJ028320	MACN-Or-70176	F21

Specimen*	Locality	Sex	Type of sample†	GenBank acc. nos.	Museum collection nos.‡	Haplotype
S. ruficollis 5	Estero Valenzuela, Corrientes, Argentina	Unknown	MLHS	FJ028319	MACN-Or-70178	F5
S. ruficollis 6	San Luis Nat. Park, Concepción, Paraguay	Male	MLHS	GU070613	KUNHM 129¶	F17
S. ruficollis 7	Trapiche, Beni, Bolivia	Unknown	MLHS	GU070614	ZMUC 123280	F5
S. zelichi 1§	Argentina	Male	BS	FJ028322	MACN-Or-ct 3132¶	F22
V. jacarina 1	San Cayetano, Corrientes, Argentina	Male	MLHS	FJ028565	MACN-Or-69796	G1

Table 2 Continued

For each individual, the species to which it belongs, the locality where it was captured, its sex, the type of sample obtained, the museum collection number, the GenBank accession number and its COI haplotype is detailed.

*S., Sporophila; V., Volatinia.

+MLHS, pectoral muscle, liver or heart sample; BS, blood sample; SS, museum study skin.

‡KUNHM, University of Kansas Museum of Natural History; LSUMZ, Louisiana State University Museum of Zoology; NHM, The Natural History Museum; PUCRS, Coleção de Aves do Museu de Ciências e Tecnologia da Pontifícia Universidad Católica do Rio Grande do Sul; STRI, Smithsonian Tropical Research Institute; USNM, Smithsonian National Museum of Natural History; ZMUC, Zoological Museum University of Copenhagen.

§Captive bird.

**This individual belongs to S. bouvreuil bouvreuil, while the remaining samples are from S. bouvreuil pileata.

¶Tissue numbers provided.

compared using the BOLD Management & Analysis System (Ratnasingham & Hebert 2007) and MEGA4 (Tamura *et al.* 2007). The K2P distance is the best metric when distances have low values (Nei & Kumar 2000), and for this reason this model is used for species-level analysis and identification in DNA barcoding (Hebert *et al.* 2003).

The DNA sequences did not possess any insertions, gaps or stop codons. They were aligned for phylogenetic analyses using BIOEDIT version 7.0.9.0 (Hall 1999). We constructed a neighbour-joining (NJ) tree using K2P distances with MEGA4 (Tamura et al. 2007). To assess robustness of the nodes we performed 1000 standard bootstrap pseudoreplicates (Felsenstein 1985). To analyse the sensitivity of topologies to the method of phylogenetic reconstruction we performed Bayesian analyses using MRBAYES 3.1.2 (Huelsenbeck & Ronquist 2001; Ronquist & Huelsenbeck 2003) and maximum parsimony (MP) analysis using TNT 1.1 (Goloboff et al. 2003). For Bayesian analysis, we selected the model of evolution using the hierarchical likelihood ratio test implemented in MODELTEST version 3.7 (Posada & Crandall 1998). The best fit was produced by the HKY two-parameter model (Hasegawa et al. 1985) with γ -distributed rate heterogeneity. We ran two independent Bayesian analyses with default priors for 5.0×10^6 generations, at which point the standard deviation of split frequencies was <0.01 indicating that both runs had converged. We sampled trees every 100 generations, discarding the first 12 500 as part of the burn-in period. The Potential Scale Reduction Factor (Gelman & Rubin 1992) was very close to 1 for all parameters, indicating that we had a sufficient sample from the posterior probability. We performed MP heuristic searches consisting of 1000 random addition sequences with the TBR

branchswapping algorithm (saving 100 trees per replication). To assess the robustness of the nodes of the resulting phylogenies, we performed 1000 standard bootstrap pseudoreplicates (Felsenstein 1985) consisting of 100 random addition sequences followed by TBR (retaining 10 trees in each pseudoreplicate).

Many COI haplotypes differed by few substitutions and in some instances were shared among species. Thus, a network approach to genealogy might help disentangle relationships. We therefore constructed a statistical parsimony network (Templeton *et al.* 1992) using TCS version 1.21 (Clement *et al.* 2000) to represent the relationship between the COI haplotypes found in the southern capuchinos.

Results

The three methods of phylogenetic reconstruction confirmed with high support that the southern capuchinos are monophyletic in relation to the northern capuchinos and the remaining outgroups. However, none of the phylogenies could distinguish among the southern capuchino species or resolve their phylogenetic affinities. Figure 1a shows the NJ tree based on K2P distances and Fig. 1b shows the virtually identical topology produced using Bayesian analysis. In both trees the northern capuchinos are polyphyletic as S. minuta is the sister species of the southern capuchino clade, while S. castaneiventris is associated with the outgroup species. Constraining the capuchinos to be monophyletic produced a Bayesian tree with the southern capuchinos forming a monophyletic clade nested within the northern capuchinos. In this topology S. castaneiventris was external to the group conformed



Fig. 1 Phylogenetic analysis based on 694 bp of the COI gene. (a) Neighbour-joining tree generated using K2P distances. Numbers indicate nodes supported in more than 50% of 1000 standard bootstrap pseudoreplicates. Bootstrap values of nodes within species clades as well as most nodes within the southern capuchino clade are omitted for simplicity. (b) Bayesian 50% majority rule consensus tree with posterior probabilities indicating node support.

by *S. minuta* and the southern capuchinos, and therefore the northern capuchinos were paraphyletic. The likelihood of both Bayesian topologies did not differ significantly according to a likelihood ratio test $[2(\ln L_1 - \ln L_0) = 0.2, \text{ d.f.} = 1, P > 0.1]$. The MP analysis (not shown) produced a topology similar to that of the constrained Bayesian analysis, although support values for the monophyly of the capuchinos were low (standard bootstrap value of 36).

The results obtained for the southern capuchinos are not surprising given that haplotypes were shared among species and that some individuals showed higher K2P distance when compared with other individuals of their own species than with representatives of different species of the group. This is apparent in the haplotype network (Fig. 2), where the 22 haplotypes found among the 38 southern capuchino individuals studied are shown. Haplotypes differed in up to 11 mutational steps and the most common one (F5) was present in five of the nine species of southern capuchinos (13 of 38 individuals; Table 2).

This case of nine species that are indistinguishable using DNA barcodes appears unique within the birds of Argentina (based on the analysis of 500 species, which represent 51% of the Argentine avifauna and the only large data set of COI sequences from neotropical Fig. 2 Unrooted maximum parsimony network showing 95% probability linkages among 22 COI haplotypes obtained from 38 individuals of the nine species of southern capuchinos. Each line represents a single mutational change. Dashed lines show alternative connections that were not unambiguously resolved by the analysis. Empty squares represent un-sampled or extinct haplotypes. Area of circles is proportional to the number of individuals with that haplotype. Haplotype F5 is present in 13 individuals, F10 and F17 in three individuals and the remaining haplotypes are present only in one individual.



birds currently available for comparison; Kerr et al. 2009). We thus compared the genetic patterns of the southern capuchinos with those of the rest of the Argentine avifauna (Fig. 3). Both the average of intraspecific and interspecific divergences and its range are similar within the southern capuchinos (K2P; 0.65% vs. 0.60% and 0.14-1.2% vs. 0.07-1.2% respectively). The highest intraspecific divergence (1.9%) was found in S. bouvreuil, the only polytypic species of the group, when comparing the rufous morph (S. bouvreuil bouvreuil: sample S. bouvreuil 1) with the white morph (S. bouvreuil pileata: six remaining samples). This finding suggests that further study is needed to clarify the systematics of this species. The average intraspecific distance in the southern capuchinos is higher than that of most Argentine species, an especially striking observation given that many of the species with higher intraspecific distances included in Fig. 3b are now suspected to include more than one lineage deserving species status (Kerr et al. 2009; Sanín et al. 2009). By contrast, the average interspecific divergence among the southern capuchinos is in the lowest extreme of the distribution of the rest of the congeneric comparisons

(Fig. 3a). Therefore, the interspecific divergence within the southern capuchinos is closer to the average intraspecific divergence than to that of the average congeneric divergence for the Argentine avifauna (a similar result is obtained if compared exclusively with the rest of the Argentine passerines). The remaining *Sporophila* species analysed, including the northern capuchinos, showed a marked difference between average intraspecific and interspecific divergence (K2P; 0.21%, range 0.12–0.37% vs. 8.2%, range 6.5–9.3%).

Discussion

In this study we sequenced 694 bp of the COI gene from the 11 species of capuchinos and several outgroups. Consistent with Lijtmaer *et al.* (2004), we found that the southern capuchinos are monophyletic in relation to the northern capuchinos and the remaining outgroups and that they have extremely low levels of interspecific divergence with most species sharing haplotypes. This explains why the COI gene neither separated the species nor resolved the phylogenetic relationships within the group. Different processes could cause shared COI



Fig. 3 Average interspecific congeneric and intraspecific K2P distances of the southern capuchinos compared with that of species present in the Argentine data set (excluding the southern capuchinos). (a) Frequency distribution of interspecific comparisons. (b) Frequency distribution of intraspecific comparisons. The arrowhead shows the position of the southern capuchinos relative to the species in the Argentine data set.

haplotypes among species and lack of reciprocal monophyly between them.

First, some cases could represent single taxa erroneously divided into more than one species (Johnston 1961). For example, it has been recently suggested, on the basis of similarities in song and habitat preference, that S. zelichi should be considered conspecific with S. palustris (Areta 2008). However, this is unlikely to be the case for most species in the group as all males differ considerably in coloration patterns and recent results show that some females can also be distinguished between species when coloration is objectively assessed with a spectrophotometer and an avian visual model is used to evaluate the results (P. Benites, unpublished data). Moreover, most southern capuchino males can be distinguished by song, showing significant differences in parameters related to time, frequency and complexity of vocalizations (L. Campagna, unpublished data) and implying some degree of reproductive isolation.

A second possibility is incomplete lineage sorting, which occurs when recently diverged taxa have not yet accumulated sequence differences in the locus analysed (Funk & Omland 2003). This option appears to be the most common in cases of avian species that lack reciprocal monophyly (Funk & Omland 2003). In the early stages of divergence, when lineage sorting is incomplete, common haplotypes are shared between species (Omland *et al.* 2006). Some COI haplotypes, notably haplotype F5, were widely shared among species of southern capuchinos, suggesting that lineage sorting is still incomplete in this group.

Alternatively, these taxa may share mtDNA because of introgressive hybridization. Around 9% of all bird species are known to have hybridized in nature (Grant & Grant 1992) and there are records of hybridization in Sporophila (Sick 1963; Ouellet 1992; Stiles 1996). Recurrent hybridization could explain the genetic pattern observed in the southern capuchinos as extensive gene flow among multiple species can make it difficult to infer patterns of genetic exchange and strongly affect mitochondrial tree topology (Funk & Omland 2003). The southern capuchino species showed higher average intraspecific and lower interspecific genetic distances than other Argentine species and we found numerous divergent haplotypes that differed by up to 10 mutational steps within single capuchino species. In this sense, incomplete lineage sorting is less likely to involve divergent allelic lineages than is introgression (Funk & Omland 2003), suggesting that in addition to a lack of lineage sorting, introgression of haplotypes via hybridization could also be responsible for the genetic pattern observed in the southern capuchinos.

We found no evidence for monophyly of the northern capuchinos. Instead our data support either paraphyly or

polyphyly. Lijtmaer *et al.* (2004) suggested that the northern capuchinos were monophyletic, although this result was equivocal because a possibly misidentified previously published sequence form *S. castaneiventris* (obtained from GenBank) was far removed from the remaining representatives of the species and therefore excluded from the conclusions. More work is needed to distinguish between these possibilities and to define whether *S. castaneiventris* should be included in the capuchinos.

Our study significantly augments the main findings of Lijtmaer et al. (2004) in relation to the southern capuchinos. We used a more comprehensive sampling, both in relation to the species of southern capuchinos and the geographic distribution of each of them, including all species and more than twice as many individuals. We used fresh tissue samples (except for S. nigrorufa) instead of museum study skins or previously published sequences and therefore minimized the risk of crosscontamination or species misidentification. Finally, the previous study was also done with mitochondrial DNA (498 bp corresponding to 303 bp of the cytochrome bgene and 195 bp of part of the cytochrome oxidase subunit II, the complete lysine transfer RNA and part of the ATP synthase subunit 8); however, an advantage of using COI in the present study is that this gene has been shown to be successful in separating sister species pairs of birds differing by as little as 0.6-0.9% sequence divergence (Baker et al. 2009). Moreover, COI has produced similar results compared with multigene approaches (Baker et al. 2009). A further advantage of DNA barcodes is that there is a quality-assured COI database of many species to which new data can be compared.

This study flags the southern capuchinos as an exceptional radiation of birds. They are the only multispecies group that cannot be identified or separated by DNA barcodes among the neotropical birds barcoded so far (Vilaça et al. 2006; Chaves et al. 2008; Kerr et al. 2009). Moreover, very few cases were identified showing average intraspecific distances <1% and these always involved a pair or trio of species that do not share COI haplotypes and exhibit diagnostic differences in their COI sequences (Kerr et al. 2009). The southern capuchinos are comparable only with the large white-headed gulls (Larus argentatus-fuscus species complex), the only similar case encountered among North American birds (Kerr et al. 2007). As in the southern capuchinos, the large white-headed gulls have very similar COI barcodes and show similarly low divergence at other loci (Hebert et al. 2004). These gulls are thought to have diverged less than 10 000 years ago, and hybridization is common among them (Crochet et al. 2002, 2003). For bird genetic studies generally, where a variety of loci are analysed, the only other large group of species with genetic divergences as

low as in the cases mentioned above are Darwin's finches (Freeland & Boag 1999; Sato *et al.* 1999, 2001), the darkeyed junco (*Junco hyemalis*) species complex (although the number of species in this complex remains a matter for debate; Milá *et al.* 2007) and possibly the crossbills (*Loxia* spp.; Edelaar *et al.* 2003), three groups of recent origin known to hybridize. For capuchinos, extending our analysis to rapidly evolving nuclear (e.g. intronic SNPs and microsatellites) and mitochondrial markers (e.g. control region) will be crucial to understanding the radiation of the southern capuchinos.

Apart from giving insights into the evolution of the capuchinos, the present study clearly shows how a standardized mitochondrial survey, like DNA barcoding, rapidly flags species or groups of species worthy of deeper study. Detecting evidence of gene flow may lead to studies of hybrid zones, mechanisms of reproductive isolation and re-examination of species limits leading to more stable classifications. Cases of incomplete lineage sorting can motivate studies in demography and speciation rates and finally high levels of intraspecific divergence may help discover cryptic species (Funk & Omland 2003). As the project to barcode the birds of the world advances, many other cases of interest to evolutionary biologists will undoubtedly be revealed.

Acknowledgements

We thank two anonymous reviewers for valuable comments on earlier versions of the manuscript. We are indebted to JC Reboreda, AS DiGiacomo, AG DiGiacomo and S Arenas for kindly providing samples and P Benites, M Blanco, PL Calderón and C Kopuchian for help in the collection of samples in the field. We thank the national fauna authorities (ED Ramadori, E Fernández and R Scandalo), the director of fauna for the province of Corrientes (SR Zajarevich), the Administración de Parques Nacionales (P Cichero and N Sucunza) and the people of El Bagual reserve. For providing tissue samples and sharing collection data, we are indebted to AT Peterson and MB Robbins from the University of Kansas Museum of Natural History, RT Brumfield and DL Dittmann from the Louisiana State University Museum of Zoology, K Cook and M Adams from The Natural History Museum, CS Fontana, M Repenning and CE Rovedder from the Coleção de Aves do Museu de Ciências e Tecnologia da Pontifícia Universidad Católica do Rio Grande do Sul and J Fjeldså and JB Kristensen from the Zoological Museum University of Copenhagen. In addition, E Bermingham, O Sanjur and M Miller from the Smithsonian Tropical Research Institute provided samples S. minuta 3-7, collected by E Bermingham and R Ricklefs supported by grants from NSF and the Smithsonian Institution. MJ Braun and J Dean from the Smithsonian National Museum of Natural History provided samples S. minuta 1-2, collected by BK Schmidt and MJ Braun respectively. We also thank the Willi Hennig Society for providing a free edition of TNT 1.1. This project was supported by grant PICT 2004-16-25171 of the Agencia Nacional de Promoción Científica y Tecnológica of Argentina, PIP 112-200801-00741 of the Consejo Nacional de

Investigaciones Científicas y Técnicas of Argentina and by the Richard Lounsbery Foundation to PL Tubaro, also by grants from the Natural Sciences and Engineering Council of Canada to SC Lougheed and from Genome Canada through the Ontario Genomics Institute to PDN Hebert.

References

- Areta JI (2008) Entre Ríos Seedeater (*Sporophila zelichi*): a species that never was. *Journal of Field Ornithology*, **79**, 352–363.
- Baker AJ, Tavares ES, Elbourne RF (2009) Countering criticisms of single mitochondrial DNA gene barcoding in birds. *Molecular Ecology Resources*, 9(Suppl. 1), 257–268.
- BirdLife International (2009) Species factsheets. Downloaded from http://www.birdlife.org.
- Chaves AV, Clozato CL, Lacerda DR, Sari EHR, Santos FR (2008) Molecular taxonomy of Brazilian tyrant flycatchers (Passeriformes: Tyrannidae). *Molecular Ecology Resources*, 8, 1169–1177.
- Clement M, Posada D, Crandall KA (2000) TCS: a computer program to estimate gene genealogies. *Molecular Ecology*, **9**, 1657– 1660.
- Crochet PA, Lebreton JD, Bonhomme F (2002) Systematics of large white-headed gulls: patterns of variation in Western European taxa. *Auk*, **119**, 603–620.
- Crochet PA, Chen JZ, Pons JM *et al.* (2003) Genetic differentiation at nuclear and mitochondrial loci among large whiteheaded gulls: sex-biased interspecific gene flow? *Evolution*, **57**, 2865–2878.
- Edelaar P, Summers R, Iovchenko N (2003) The ecology and evolution of crossbills *Loxia* spp.: the needs for a fresh look and an international research programme. *Avian Science*, **3**, 85–93.
- Felsenstein J (1985) Confidence limits on phylogenies: an approach using the bootstrap. *Evolution*, **39**, 783–791.
- Freeland JR, Boag PT (1999) The mitochondrial and nuclear genetic homogeneity of the phenotypically diverse Darwin's ground finches. *Evolution*, **53**, 1553–1563.
- Funk DJ, Omland KE (2003) Species-level paraphyly and polyphyly: frequency, causes, and consequences, with insights from animal mitochondrial DNA. *Annual Review of Ecology*, *Evolution, and Systematics*, **34**, 397–423.
- Gelman A, Rubin DB (1992) Inference form iterative simulation using multiple sequences. *Statistical Science*, **7**, 457–472.
- Goloboff P, Farris S, Nixon K (2003) TNT: tree analysis using New Technology. Program and documentation, available from the authors, and at http://www.zmuc.dk/public/phylogeny.
- Grant PR, Grant BR (1992) Hybridization of bird species. *Science*, **256**, 193–197.
- Hajibabaei M, Janzen DH, Burns JM, Hallwachs W, Hebert PDN (2006) DNA barcodes distinguish species of tropical Lepidoptera. Proceedings of the National Academy of Sciences, USA, 103, 968–971.
- Hall TA (1999) BioEDIT: a user-friendly biological sequence alignment editor and analysis program for Windows 95/98/NT. *Nucleic Acids Symposium Series*, **41**, 95–98.
- Hasegawa M, Kishino H, Yano T (1985) Dating the human-ape splitting by a molecular clock of mitochondrial DNA. *Journal* of *Molecular Evolution*, **22**, 160–174.
- Hebert PDN, Cywinska A, Ball SL, deWaard JR (2003) Biological identifications through DNA barcodes. *Proceedings of the Royal Society B: Biological Sciences*, **270**, 313–321.

458 DNA BARCODING

- Hebert PDN, Stoeckle MY, Zemlak TS, Francis CM (2004) Identification of birds through DNA barcodes. *Public Library of Science Biology*, 2, 1657–1663.
- Hellmayr CE (1938) Catalogue of Birds of the Americas and the Adjacent Islands in Field Museum of Natural History. Fieldiana Zoology, Volume 13, Part 11. Field Museum of Natural History, Chicago.
- Howard R, Moore A (1991) A Complete Checklist of the Birds of the World, 2nd edn. Academic Press, London.
- Huelsenbeck JP, Ronquist F (2001) MRBAYES: Bayesian inference of phylogeny. *Bioinformatics*, 17, 754–755.
- Johnston DW (1961) *The Biosystematics of American Crows*. University of Washington Press, Seattle, WA.
- Kerr KCR, Stoeckle MY, Dove CJ *et al.* (2007) Comprehensive DNA barcode coverage of North American birds. *Molecular Ecology Notes*, **7**, 535–543.
- Kerr KCR, Lijtmaer DA, Barreira AS, Hebert PDN, Tubaro PL (2009) Probing evolutionary patterns in Neotropical birds through DNA barcodes. *PLoS ONE*, 4, e4379. DOI: 10.1371/ journal.pone.0004379.
- Kimura M (1980) A simple method for estimating evolutionary rate of base substitutions through comparative studies of nucleotide sequences. *Journal of Molecular Evolution*, **15**, 111– 120.
- Lijtmaer DA, Sharpe NM, Tubaro PL, Lougheed SC (2004) Molecular phylogenetics and diversification of the genus *Sporophila* (Aves: Passeriformes). *Molecular Phylogenetics and Evolution*, **33**, 562–579.
- Maddison WP, Knowles LL (2006) Inferring phylogeny despite incomplete lineage sorting. *Systematic Biology*, 55, 21–30.
- Meyer de Schauensee RM (1952) A review of the genus Sporophila. Proceedings of the Academy of Natural Sciences of Philadelphia, **104**, 153–196.
- Milá B, McCormack JE, Castañeda G, Wayne RK, Smith TB (2007) Recent postglacial range expansion drives the rapid diversification of a songbird lineage in the genus Junco. Proceedings of the Royal Society B: Biological Sciences, 274, 2653– 2660.
- Miller SA, Dykes DD, Polesky HF (1988) A simple salting out procedure for extracting DNA from human nucleated cells. *Nucleic Acids Research*, **16**, 1215.
- Nei M, Kumar S (2000) Molecular Evolution and Phylogenetics. Oxford University Press, Oxford.
- Omland KE, Baker JM, Peters JL (2006) Genetic signatures of intermediate divergence: population history of Old and New World Holarctic ravens (*Corvus corax*). *Molecular Ecology*, 15, 795–808.
- Ouellet H (1992) Speciation, zoogeography and taxonomic problems in the neotropical genus *Sporophila* (Aves: Emberizinae). *Bulletin of the British Ornithological Club*, **112**, 125–135.

- Posada D, Crandall KA (1998) MODELTEST: testing the model of DNA substitution. *Bioinformatics*, **14**, 817–818.
- Ratnasingham S, Hebert PDN (2007) BOLD: the barcode of life data system (http://www.barcodinglife.org). *Molecular Ecol*ogy Notes, 7, 355–364.
- Ridgely RS, Tudor G (1989) *The Birds of South America*. University of Texas Press, Austin, TX.
- Ronquist F, Huelsenbeck JP (2003) MRBAYES 3: Bayesian phylogenetic inference under mixed models. *Bioinformatics*, 19, 1572–1574.
- Sanín C, Cadena CD, Maley JM et al. (2009) Paraphyly of Cinclodes fuscus (Aves: Passeriformes: Furnariidae): implications for taxonomy and biogeography. Molecular Phylogenetics and Evolution, 53, 547–555.
- Sato A, O'hUigin C, Figueroa F et al. (1999) Phylogeny of Darwin's finches as revealed by mtDNA sequences. Proceedings of the National Academy of Sciences, USA, 96, 5101–5106.
- Sato A, Tichy H, O'hUigin C et al. (2001) On the origin of Darwin's finches. Molecular Biology and Evolution, 18, 299–311.
- Sibley CG, Monroe BL Jr (1990) *Distribution and Taxonomy of Birds* of the World. Yale University Press, New Haven, CT.
- Sick H (1963) Hybridization in certain Brazilian Fringillidae (Sporophila and Oryzoborus). Proceedings of the 13th International Ornithology Congress, 1962, 161–170.
- Silva JMC (1999) Seasonal movements and conservation of seedeaters of the genus *Sporophila* in South America. *Studies in Avian Biology*, **19**, 272–280.
- Smith MA, Poyarkov NA, Hebert PDN (2008) CO1 DNA barcoding amphibians: take the chance, meet the challenge. *Molecular Ecology Notes*, 8, 235–246.
- Stiles FG (1996) When black plus white equals gray: the nature of variation in the variable seedeater complex (Emberizinae: *Sporophila*). Ornitología Neotropical, 7, 75–107.
- Tamura K, Dudley J, Nei M, Kumar S (2007) MEGA4: molecular evolutionary genetics analysis (MEGA) software version 4.0. *Molecular Biology and Evolution*, 24, 1596–1599.
- Tavares ES, Baker AJ (2008) Single mitochondrial gene barcodes reliably identify sister-species in diverse clades of birds. *BMC Evolutionary Biology*, **8**, 81.
- Templeton AR, Crandall KA, Sing CF (1992) A cladistic analysis of phenotypic associations with haplotypes inferred from restriction endonuclease mapping and DNA sequence data. III. Cladogram estimation. *Genetics*, **132**, 619–633.
- Vilaça ST, Lacerda DR, Sari EHR, Santos FR (2006) DNA-based identification applied to Thamnophilidae (Passeriformes) species: the first barcodes of Neotropical birds. *Revista Brasileira de Ornitologia*, 14, 7–13.
- Ward RD, Zemlak TS, Innes BH, Last PR, Hebert PDN (2005) DNA barcoding Australia's fish species. *Philosophical Transactions of the Royal Society B, Biological Sciences*, 360, 1847–1857.