Evolutionary relationships of stonechats and related species inferred from mitochondrial-DNA sequences and genomic fingerprinting

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ABSTRACT Sequence data of the mitochondrial cytochrome b gene and genomic fingerprinting provide good evidence that the geographically separated taxa of the Common Stonechat Saxicola torquata complex represent distinct genetic lineages, which became separated more than one million years ago. The distinct genetic pattern implies that hybridisation and gene flow between these lineages no longer takes place to a significant degree. Since these lineages also differ in morphology, breeding behaviour, vocalisations and physiological control of their annual cycles, we suggest treating European Stonechat S. torquata, African Stonechat S. [t.] axillaris, Reunion Stonechat S. [t.] tectes, Canary Islands Stonechat S. dacotiae and Siberian Stonechat S. [t.] maura as 'good' and distinct species.

Footnote: In this paper, the vernacular and scientific names used for the key taxa of interest are those preferred by the author, although this does not necessarily reflect editorial policy.

Introduction

Systematic classifications are usually based on similarity, with taxa that show the greatest similarity being considered to be closely related. Since taxa which are only distantly related may, however, come to show morphological similarities owing to adaptation to similar ecological constraints – 'convergent evolution', one example being penguins (Spheniscidae) and auks (Alcidae) – evolutionary trees based on the study of adaptive characters can lead to erroneous taxonomic conclusions.

The analysis of nuclear or mitochondrial marker genes has, during the last 15 years, become widely used to reconstruct phylogenetic relationships between taxa in all fields of zoology, including ornithology (see Avise 1994; Mindell 1999). Molecular data have the great advantage that convergent evolution does not impair the analysis to the same degree as it does

for morphological data. If two or more taxa belong to the same species, their marker genes are identical, or almost identical, and intraspecific genetic distances are generally significantly smaller than those between established species. Molecular data may also provide an estimate for the timescale over which a particular evolutionary step has taken place (the 'molecular clock') and, therefore, allow both a phylogenetic and a geographical analysis of the unknown past of a group of organisms. Molecular data have consequently become an important tool for taxonomic and evolutionary studies.

Few molecular studies have addressed phylogenetic relationships in stonechats and other members of the family Turdidae. Based on the analysis of 300 base pairs of the mitochondrial cytochrome b gene, preliminary data have suggested that European Stonechat Saxicola torquata, Siberian Stonechat S. [t.] maura and



220. Male European Stonechat Saxicola torquata, Suffolk, May 2001.

African Stonechat S. [t.] axillaris have already diverged to a high degree (Wittmann 1994; Wittmann et al. 1995).

For the analysis presented here, we sequenced most of the *cytochrome b* gene of 23 turdid taxa in seven genera, and also enlarged the dataset by including more stonechat taxa than in our previous study (Wittmann *et al.* 1995).

The fertilised egg contains only its mother's mitochondria but receives nuclear DNA from both parents. Since mitochondrial DNA (mtDNA) is inherited only through the female line, it does not fully reflect the evolution of all members of the species, and can give a distorted view of phylogenetic relationships. We have, therefore, additionally analysed the structure of the nuclear genome using the polymerase chain reaction (PCR) to produce 'Inter-Simple-Sequence Repeat' ('ISSR')-profiles, as described in an appendix at the end of this paper. ISSR-PCR is a very effective method by which to understand interspecific variation and genetic structure of populations (Damodar Reddy et al. 1999; Ge & Sun 1999), to sex individuals (Wink et al. 1998), to generate species-specific genomic fingerprints (Gupta et al. 1994; Zietkiewicz et al. 1994) and to detect hybridisation between taxa (Wink et al. 2000).

Materials and methods

Blood and tissues were either preserved in an EDTA buffer (0.1 M Tris [pH 7.4], 10% EDTA, 1% NaF, 0.1% thymol) or in ethanol (Wink et al. 1998) and stored at -20°C until processing. Total DNA was extracted from the blood

samples by an overnight incubation at 37°C in lysis buffer (10 mM Tris [pH 7.5], 25 mM EDTA, 75 mM NaCl, 1% SDS) including 1 mg of Proteinase K (Merck, Darmstadt), followed by a standard phenol/chloroform protein extraction. DNA was precipitated from the supernatant with 0.8 volume of cold isopropanol, centrifuged, washed, dried and resuspended in TE buffer.

Full details of the sequencing technology, the ISSR-PCRs, and parameters and methodologies used to draw up the phylogenetic trees can be obtained from the authors.

Results and Discussion

The analysis was restricted mainly to species of the Western Palaearctic, and comprised the following genera: stonechats Saxicola, rock thrushes Monticola, redstarts Phoenicurus, wheatears Oenanthe, robins Erithacus, nightingales Luscinia, and thrushes Turdus. The dippers Cinclus were included as a possibly closely related comparison (ingroup), and the wrens Troglodytes as a distantly related outgroup. Although we sequenced several individuals of each taxon (between six and 20 in the case of stonechat subspecies), within any given subspecies all individuals clustered as a single genealogically related group derived from a single common ancestor (a monophyletic clade). One to three representatives of each taxon were chosen for an analysis which aimed to outline the patterns of phylogeny and speciation within stonechats and related Turdidae.

In fig. 1, the result of maximum parsimony and maximum likelihood analyses of mtDNA

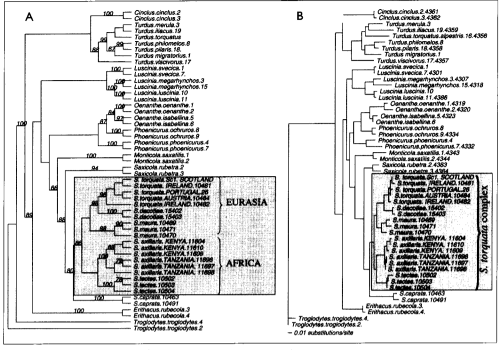


Fig. 1. A molecular phylogeny of stonechats and related turdids based on sequence data of the mitochondrial cytochrome b gene (see text for more details).

A. Maximum parsimony analysis; (bootstrap values above 60% are given at the corresponding bifurcations).

B. Maximum likelihood analysis; branch lengths correspond to genetic distances.

The maximum parsimony method takes the DNA sequences of the extant species and creates a phylogenetic tree linking them all, one which requires the least number of mutational changes during the evolution of all the species. It assumes that no more evolution will have taken place than was absolutely necessary. In contrast, the maximum likelihood method does not make this assumption, it takes specific mathematical models based on current knowledge of how DNA (or proteins) mutates during evolution, accepting that mutation is a random process. Using this knowledge, it is possible to work out the statistical likelihood of any phylogenetic tree based on the observed DNA sequences being the right ones. Of all possible trees, the one calculated as being most likely is accepted.

are given. Both methods arrange the taxa into identical monophyletic groups and are also congruent in branching order. Members of the genus Saxicola form a clade (supported by a bootstrap statistical value of 88%), which derives from a common ancestor that is shared with Rock Thrush Monticola saxatilis. The Monticola/Saxicola clade shares ancestry with the genera Luscinia, Oenanthe, Phoenicurus and Erithacus. These genera are grouped in the apparently monophyletic tribe Saxicolini (supported by a bootstrap value of 89%).

Members of the genus *Turdus*, which form a monophyletic clade (88% bootstrap support), always cluster as a sister group to the tribe Saxicolini. The Dipper *Cinclus cinclus* shows no close affinity to members of the genus *Turdus*, (pairwise genetic distances 12-15%), or to wrens with which they have also been associ-

ated. We need a more complete dataset before considering the real affinities of dippers and wrens.

Relationships within the stonechat complex Mitochondrial-DNA sequences show that, within the genus Saxicola, the Whinchat S. rubetra takes a basal position, followed by Pied Stonechat S. caprata. Both these species are, therefore, assumed to have split from the ancestors of European and Siberian Stonechats early in the evolution of the genus. The S. torquata complex, which includes S. torquata, S. [t.] tectes, S. [t.] maura, S. [t.] axillaris and S. dacotiae, clusters as a monophyletic group (with bootstrap support 86%). Within the complex the following relationships are apparent: the Canary Islands Stonechat S. dacotiae and the European Stonechat S. torquata rubicola/hiber-

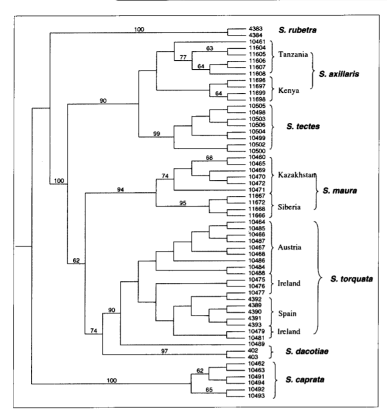


Fig. 2. UPGMA cluster analysis of ISSR-profiles. The pattern of polymorphic-ISSR fragments was recorded in a 0/1 matrix and the results were analysed by UPGMA bootstrap analyses (1,000 replications; bootstrap values above 60% are given at the corresponding bifurcations). Numbers following taxon names refer to internal accession numbers.

nans appear as sister taxa (supported by a bootstrap value of 98%). The Reunion Stonechat S. [t.] tectes and the African Stonechat S. [t.] axillaris form another sister pair. The Siberian Stonechat S. [t.] maura shows affinities to the S. dacotiae/S. torquata

clade. Within the African Stonechat, two populations were studied, from Kenya and Tanzania, which show different and consistent mtDNA sequences (haplotypes), the genetic distances being 0.4-0.6%. Within *S. torquata*, a clear attribution of mtDNA haplotypes to subspecies (*S. t. hibernans* from Ireland, *S. t. rubicola* from central and western Europe) and/or geographic ranges was not apparent. It is not, therefore, possible to define subspecies of the European Stonechat on the basis of their mtDNA alone, although intraspecific variation is obvious (intraspecific distances 0.1-1.3%).

The Canary Islands Stonechat *S. dacotiae* (which occurs on Fuerteventura) is closely related to the European Stonechat *S. torquata*. The genetic distance of 2.7-3.5% implies that these taxa diverged about 1.3-1.7 million years ago, if the '2% per million years' rule (Wilson *et al.* 1987) is taken for calibration of mtDNA. Most likely, the Canary Islands Stonechat derived from a founder population of European Stonechats, which became resident on the Canary Islands, or from a population of Euro-



221. Female European Stonechat Saxicola torquata, Suffolk, May 2001.



222. First-winter European Stonechat Saxicola torquata, Norfolk, September 1996.

pean Stonechats living in northern Africa. Both taxa share common ancestry with the Siberian Stonechat, from which they differ respectively by 2.7% and 5.1% base-pair substitutions.

The African Stonechat S. [t.] axillaris, differs from the European Stonechat by 4.6-5.7% substitutions. It clusters as a sister taxon to the Reunion Stonechat S. [t.] tectes, which has sometimes been recognised as a distinct species (Hall & Moreau 1970). Pairwise distances amount to 4.4-5.1%.

Using genomic fingerprinting with ISSR-PCR, it is apparent that S. t. rubicola/hibernans, S. [t.] axillaris, S. dacotiae, S. [t.] tectes and S. [t.] maura form distinct genetic entities (fig. 2). Once again, S. caprata and S. rubetra cluster at the base of the tree, as in the analyses derived from the mitochondrial cytochrome b gene. The stonechats are again recognised as a monophyletic assemblage (bootstrap support 86%); the relationships within this complex are almost identical to those found for cytochrome b: S. dacotiae clusters as a sister group to S. torquata rubicola/hibernans; S. [t.] axillaris is a sister group to S. [t.] tectes, and S. [t.] maura is intermediate. Also, the phylogeographic divisions within S. [t.] axillaris and S. [t.] maura can be recovered by the ISSR-data. The ISSR-profiles, based on nuclear DNA, clearly indicate that the results obtained from mitochondrial cytochrome b sequences correctly reflect the phylogeny of this group of birds.

ISSR and *cytochrome b* sequences both show a certain degree of geographic variation in the

DNA of stonechats (figs. 1-2). Because we have sampled a limited number of populations so far, our analysis of intraspecific variation must be regarded as preliminary. It would certainly be rewarding for further investigations to sample genetic variation across the complete geographic distribution range of stonechats.

The forms S. [t.] tectes, S. [t.] maura, S. [t.] axillaris and S. dacotiae have often been treated as subspecies of Common Stonechat S. torquata (see Sibley & Monroe 1990). On the basis of diagnostic differences in morphology, breeding biology and distribution, however, these taxa have already been recognised as distinct species in several bird guides and handbooks. If the members of the S. torquata complex were closely related subspecies, we should expect a cluster showing no (or just extremely small) genetic differences between each of the stonechat taxa. Instead, both the cytochrome b sequence data and ISSR-profiles exhibit a welldeveloped phylogenetic pattern with relatively long branch lengths separating them (figs. 1-2). Genetic distances between the taxa of the S. torquata complex range between 6.1% and 2.7%, whereas distances within a given subspecies do not exceed 1.5%. The genetic distances within the S. torquata complex imply divergence times of one to three million years. In our dataset, genetic distances for other sibling species are, for example, 6.6% for Rufous Luscinia megarhynchos and Thrush Nightingales L. luscinia; 5.0% for Northern Oenanthe oenanthe and Isabelline Wheatears O.



223. Siberian Stonechat Saxicola [torquata] maura, Khao Yai, Thailand, November 1991.

isabellina; and 3.4% for Fieldfare Turdus pilaris and Song Thrush T. philomelos. These comparisons indicate that the genetic distance found between members of the S. torquata complex are in a similar range to other closely related, but distinct species. It is, therefore, highly likely that the taxa within the stonechat complex represent distinct genetic lineages. Not only do they differ in genetic terms, but they also differ by their allopatric distribution, ecology, physiology (Gwinner & Scheuerlein 1999; Helm & Gwinner 1999) and morphology (Starck et al. 1995; Helm & Gwinner 2001). As a consequence, European Stonechat, African Stonechat, Canary Islands Stonechat, Reunion Stonechat and Siberian Stonechat can be regarded as distinct species, not only within the Phylogenetic Species Concept but also within the Biological Species Concept.

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Appendix ISSR-PCR

The genomes of all animals and plants contain stretches of small repeated sequences of DNA such as .CACACACACACACACACA, that occur because of faults during the processes of replication and repair of DNA. These are called 'microsatellites'. In general, if there is gene flow between two different populations or subspecies, their patterns of microsatellites will be broadly identical (although mutations may still be found). Once gene flow is restricted, however, such as between closely related species or isolated subspecies that no longer breed with each other, the number, length and arrangement of these repeats starts to diverge randomly. 'ISSR-PCR' attempts to turn this divergence into a form that humans can see and score. It uses a small manufactured stretch of

DNA (a primer) that binds specifically to some of these satellites in DNA isolated from the taxa of interest. An enzyme is used to create many thousands or millions of copies of the DNA that lies between these microsatellites. These copies can be separated and visualised according to their size by running them through a gel that acts like a sieve. Individuals with identical microsatellite patterns in their DNA will produce identical bands on the gel, but if mutations have occurred, different individuals will show slightly different banding patterns. The presence or absence of bands on the gel can be scored for each individual, and these scores fed into the algorithm that draws up phylogenetic trees. Taxa with the most similar banding patterns will tend to be the most closely related.

Looking back

Fifty years ago:

[From: Blue-cheeked Bee-eater in Scilly: a new British Bird] 'Miss Hilda M. Quick, to whom full credit for the observation is due, sent us a short account of the bird immediately after she had seen it. At our request she has supplied the following details of the occurrence:

"On June 22nd, 1951, having a visitor staying with me on St Agnes (Isles of Scilly), I went up the lane before breakfast to get the morning's milk. Something skimmed across the path which registered as strange...only a glimpse but it was definitely odd. I wondered if it could have been a Starling looking greener than usual, or a pale Swallow with a light sheen on it (the flight suggested something of the Swallow tribe). However, I had to go back and get breakfast for my guest, and leave the mystery for a moment.

"While we were breakfasting, a neighbour came in to say that Mr Lewis Hicks had seen a strange and most wonderful bird. So we left breakfast standing and rushed out; collected Mr Hicks and went with him to the fields where he had seen it. (He told me afterwards that at first he could not believe his eyes, and went to fetch his wife to 'come and tell me if you see what I see!') Fairly soon we saw our quarry afar off, and presently it perched at some distance, but in good view. There was no doubt that it was a bee-eater; one knew it from pictures, and the curved bill and elongated tailfeathers could be clearly seen. It seemed to be returning fairly regularly to one spot on the telegraph wires, so I stalked gently up the lane to a position within 20 feet or so of where it came back to perch. It made frequent sallies after insects and brought them back to the wire

to eat. ('Sitting there eating up my bees as fast as it can!' said Mr Hicks, who has hives...but we could not be sure that they were truly bees that it was catching.) Once, its prey escaped from its bill, and it did a lightning dive and turned to recapture it and bring it back to the wire." (Brit. Birds 45: 225, July 1952)

Fifty years ago:

[From: Reports from Observatories, 1951. 'MONKS' HOUSE BIRD OBSERVATORY, 1951, By DR E. A. R. ENNION.] 'MONKS' HOUSE OBSERVATORY commenced its first season's trapping towards the end of April, 1951, after a winter spent in alterations and equipment to provide living and working quarters for the Director and Mrs Ennion, and for up to a full complement of about 25 visitors. It is therefore a rather more permanent establishment than are many of its fellow Observatories and, in its first year, has welcomed some 500 visitors, most, but not all, ornithologists. Four Bird Courses, each lasting a fortnight, have been held, together with a number of "ringing instruction weeks"; also many Courses in marine and other branches of outdoor biology at times when bird activities are to the fore. The Observatory is thus able to cover a wide field and to pursue its recording and ringing throughout the year, especially in winter when about 1,000 birds of 25 spp. (including 450 Greenfinches and 250 Linnets) were trapped between autumn and spring passages, 1951-52. The proportion of Greenfinches retrapped in stackyards and roosts suggests that at least half of the local population of these birds has been ringed: we have barely touched the Linnets!' (Brit. Birds 45: 233, July 1952)