Hybridization between white-headed ducks and introduced ruddy ducks in Spain

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Abstract

The ruddy duck, Oxyura jamaicensis, was introduced to Great Britain in the mid-20th century and has recently spread to other Western European countries. In Spain, ruddy ducks hybridize with the globally endangered white-headed duck, Oxyura leucocephala. We assessed the effects of hybridization on the Spanish white-headed ducks, which constitute 25% of the global population of this species, using a panel of eight nuclear intron markers, 10 microsatellite loci, and mtDNA control region sequences. These data allowed parental individuals, F1 hybrids, and the progeny of backcrossing to be reliably distinguished. We show that hybrids between the two species are fertile and produce viable offspring in backcrosses with both parental species. To date, however, we found no extensive introgression of ruddy duck genes into the Spanish white-headed duck population, probably due to the early implementation of an effective ruddy duck and hybrid control programme. We also show that genetic diversity in the expanding European ruddy duck population, which was founded by just seven individuals, exceeds that of the native Spanish white-headed duck population, which recently recovered from a severe bottleneck. Unless effective control of ruddy ducks is continued, genetic introgression will compromise the unique behavioural and ecological adaptations of white-headed ducks and consequently their survival as a genetically and evolutionary distinct species.

Keywords: hybridization, introns, microsatellites, mitochondrial DNA, Oxyura, white-headed duck

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Introduction

Human activities have resulted in the extension of certain species’ native geographical ranges or introductions to new areas, which in turn have led to hybridization with native plant and animal species, often with significant negative consequences (e.g. Kanda et al. 2002; Moody & Less 2002). Hybridization is common in birds, and particularly among waterfowl (Grant & Grant 1992). Hybridization between mallards Anas platyrhynchos and several closely related species has resulted from introductions of mallards or shifting geographical ranges. The Hawaiian duck (Anas wyvilliana, Browne et al. 1993) and New Zealand grey ducks (Anas superciliosa superciliosa, Gillespie 1985; Rhymer et al. 1994) have suffered from extensive hybridization with introduced mallards, whereas continental mallard populations hybridize with black ducks (Anas rubripes, Mank et al. 2004) and mottled ducks (Anas fulvigula, McCracken et al. 2001; Williams et al. 2005) in North America, and eastern spot-billed, Anas zonorhyncha, in Russia (Kulikova et al. 2004). The production of hybrid individuals, however, does not necessarily imply genetic introgression between species. Hybrid individuals may be infertile or have low fitness due to reduced survival and/or reproductive success (Liou & Price 1994; Coyne & Orr 2004). In birds, incomplete prezygotic and postzygotic isolation are common. Particularly in waterfowl, hybrids are often fertile (Grant & Grant 1992; Price & Bouvier 2002), but occur infrequently in most natural populations, making introgression unlikely. Therefore, identification of hybrids as well as information on their fertility and reproduction in the wild is crucial to assessing the impact of hybridization on a species.

The white-headed duck is the only stifftail duck (subfamily Oxyurinae) native to the Palaearctic and is classified as Endangered by the IUCN (BirdLife International 2000).
Although white-headed ducks had a wide distribution in the past, extending from Spain and Morocco in the west to the western limit of China in the east, populations became fragmented and suffered major declines in recent decades (Green & Hughes 2001). After extinctions in Egypt, Central and Eastern Europe, and Asia, extant populations have become isolated from each other (Green & Hughes 2001; Li & Mundkur 2003). The largest population in the western half of the species’ range is in Spain, and it is the only population that has been growing in recent decades. Only 22 individuals were counted in 1977 (Torres & Moreno-Arroyo 2000a), but the population has since recovered and appears to have stabilized at about 2500 birds (Hughes et al. 2004), constituting about 25% of the current world population. This successful recovery, however, has been marred by the arrival in Spain of the North American ruddy duck.

Seven ruddy ducks were imported to a zoological collection in England in 1948 and during subsequent years approximately 90 of their descendants escaped to the wild (Hudson 1976). As the feral ruddy duck population increased in Great Britain, it expanded to other Western European countries and Morocco (Hughes et al. 2004), thus invading the white-headed duck’s range [see Muñoz-Fuentes et al. (2005a), Brua (2001), and Hughes (1997) for more information on the geographical ranges of the two species]. In 1984, ruddy ducks numbered 2000 in the UK alone (Collier et al. 2005) and were detected in Spain for the first time (Torres & Moreno-Arroyo 2000b). The first ruddy duck × white-headed duck hybrid was identified in 1991 (Torres & Moreno-Arroyo 2000b), contributing to concerns for the white-headed duck that have prompted efforts to eliminate feral ruddy ducks in Portugal, Spain, France and the UK. Hybridization and up to two generations of backcrossing have been observed in captivity (B. Hughes, unpublished) but the degree of genetic introgression and patterns of hybridization in the wild are unknown.

In this study, we used a panel of molecular genetic markers to (i) confirm that ruddy ducks and white-headed ducks are two well-differentiated species; (ii) compare the genetic diversity of their populations; (iii) determine the parentage of suspected hybrid individuals; and (iv) assess the degree of genetic introgression between the two species in Spain. Our data set includes mitochondrial DNA (mtDNA), 10 nuclear microsatellite loci derived from parallel genomic libraries for Spanish ruddy ducks and white-headed ducks, and fixed differences in nuclear introns. These diagnostic markers in combination with microsatellite data allowed the discrimination of F1 and subsequent generation hybrids, categories that may be difficult to discriminate with microsatellite data alone (Vähä & Primmer 2006), and therefore allowed us to test for successful reproduction of hybrid individuals. Information on mtDNA haplotypes also allowed inferences about potential asymmetries in the composition of hybrid pairs and fertility of different hybrid classes.

Materials and methods

Samples

Our analyses were based on 31 ruddy ducks, 63 white-headed ducks and 29 presumed hybrids collected between 1993 and 2003 across Spain. Samples from white-headed ducks were obtained from individuals found dead in the field, except for seven individuals ringed, sampled and released in 2003. Ruddy duck and hybrid samples were obtained from animals shot as part of the ongoing eradication programme. Initial identification of each individual was based on morphology, mainly plumage characteristics and beak shape (Urdiales & Pereira 1993), and was completed by ornithologists familiar with these species prior to the genetic analysis.

In hybridization studies, it is essential to compare the genetic composition of the species of interest with allopatric individuals of the same species, i.e. collected in areas free of hybridization. Therefore, to identify species-specific alleles, we used ruddy ducks collected between 1987 and 2003 from areas in Europe where they do not co-exist with white-headed ducks (England, France, Iceland and two captive-bred populations) and from North America (for microsatellite markers, ruddy duck populations are not differentiated across Europe, but they are significantly differentiated from North American ruddy ducks; Muñoz-Fuentes et al. 2006), and white-headed ducks from Greece and a captive population in Spain (El Acebuche, Doñana National Park). This latter captive population was established before ruddy ducks were recorded in Spain and is therefore free of introgression. Tissues included blood, brain, muscle and feathers. DNA was extracted as described in Muñoz-Fuentes et al. (2005a).

Sequencing of the mtDNA control region

For all hybrids, and a sample of 27 ruddy ducks and 43 white-headed ducks, we determined mtDNA haplotypes. In previous studies, we sequenced 575 bp of the mtDNA control region for both ruddy ducks and white-headed ducks using primers L81 and H768 (Muñoz-Fuentes et al. 2005a, 2006). Based on these data, we designed species-specific primers to selectively amplify the mtDNA of one species but not the other, allowing the maternal line of additional individuals to be identified with a simple polymerase chain reaction (PCR) test. Each sample was tested with OXJA.CRF1 (5′-CATAYCATGCTCCCACCCATAC-3′) and OXJA.CRR1 (5′-TTTACGTTAGGTGTACGGCT-3′) for ruddy duck mtDNA and OXLE.CRF1 (5′-CATGCTCCACCCATAACCA-3′) and H493 (Sorenson & Fleischer 1996) for white-headed duck mtDNA. DNA was amplified in 25-µL reactions as in Muñoz-Fuentes et al. (2005a). PCRs were performed in a GeneAmp PCR System
Thus, for the Spanish sample, sequence data were available under UV light in 2% agarose with ethidium bromide. Presence or absence of amplification was visualized based on alignments of chicken (Gallus gallus), mallard (Anas platyrhynchos), and selected mammals, as available (see Table S1 for primer sequences). Primers were placed in the exons flanking each intron. For each intron, we obtained sequences for allopatric white-headed ducks and ruddy ducks; ruddy ducks from England or France and white-headed ducks from Greece or the captive population at El Acebuche, Doñana National Park.

Depending on the nature of the difference, we then designed assays to genotype the diagnostic polymorphic base at each locus (Table 1). First, if a fixed difference resulted in a restriction site difference between the two species (four of eight loci), we digested PCR products with the appropriate restriction enzyme and then determined genotypes from the presence or absence of restriction fragments in an agarose gel. For one locus with a transversion difference between the two species (PEPK9), we designed species-specific primers with the 3′ base on the polymorphic site and visualized the presence or absence of amplification in an agarose gel. In the case of another locus (Trop), sequences of the two species differed in length by 13 bp, due to two indels of 5 and 8 bp each, such that the difference in size of PCR products could be visualized in an agarose gel.

### Table 1

<table>
<thead>
<tr>
<th>Locus name</th>
<th>Intron</th>
<th>Primer names</th>
<th>Primer sequence (5′−3′)</th>
<th>T_a (°C)</th>
<th>Detection method</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alpha enolase (Enol, E)</td>
<td>8</td>
<td>OXY.Enol.8F</td>
<td>F: AAATCTTTCAACATTTAGCTCC</td>
<td>55</td>
<td>RE (Hpy 188I)+</td>
</tr>
<tr>
<td></td>
<td></td>
<td>OXY.Enol.8R</td>
<td>R: TCTATATATTTGAATCTTCTCC</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5-aminolevulinate synthase (Alsy, A)</td>
<td>8</td>
<td>OXY.Alsy.8F</td>
<td>F: CTGTATTAGTAGAGAAGAAGA</td>
<td>55</td>
<td>RE (BspHI)+</td>
</tr>
<tr>
<td></td>
<td></td>
<td>ALSY.9R</td>
<td>R: TCCTATATTATTTAAGCATCTTCC</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Rhodopsin (RHDP, R)</td>
<td>1</td>
<td>RHDP.1F</td>
<td>F: TTTCTCTTTGAGTAAAGGTTGA</td>
<td>55</td>
<td>RE (Fnu 4HI)+</td>
</tr>
<tr>
<td></td>
<td></td>
<td>OXY.RHDP.1R</td>
<td>R: CAGCTCGTTAATCGCTTTGCC</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Transforming growth factor-beta2 (TGFb2, G)</td>
<td>5</td>
<td>OXY.TGFb2.5F</td>
<td>F: GACCTTGAAGAGCAAGACTGAGC</td>
<td>55</td>
<td>RE (HaeIII)+</td>
</tr>
<tr>
<td></td>
<td></td>
<td>OXY.TGFb2.5R</td>
<td>R: CTTCTATACAACTTAAATTTCCC</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Tropomyosin (Trop, T)</td>
<td>5</td>
<td>Trop.5F</td>
<td>F: GAATTGAGACGAGCAACCCGG</td>
<td>55</td>
<td>Length difference</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Ox.Trop.1R</td>
<td>R: CTCTGAGCCTCCAAAACAGACTTAGCAC</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Phosphoenolpyruvate carboxykinase (PEPK9, P)</td>
<td>9</td>
<td>OxLe.PEPCK</td>
<td>F: GACCTATATTTATCCCTAAGCTTCTCA</td>
<td>60</td>
<td>PCR success/failure</td>
</tr>
<tr>
<td></td>
<td></td>
<td>PEPCK.9R*</td>
<td>R: GGCCATATGCTGACCACTG</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>OxA.PEPCK</td>
<td>F: CTTATATTTATCCCTAAGCTTCTCA</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>PEPCK.9R</td>
<td>R: CTCTGAGCCTCCAAAACAGACTTAGCAC</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lactate dehydrogenase-B (LDHB, L)</td>
<td>3</td>
<td>LDHB.3P</td>
<td>F: GAAGAYAARCTTAAAGGAAATGATGGA</td>
<td>55</td>
<td>Single-base extension</td>
</tr>
<tr>
<td></td>
<td></td>
<td>LDHB.Oxy.4R</td>
<td>R: GCTTGAATGCAACCATCACC</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>LDHB.Oxy.SNP</td>
<td>S: ACTATATATATGGACACAGA</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Myelin proteolipid protein (MPP, M)</td>
<td>4</td>
<td>OXY.MPP.4F</td>
<td>F: CTTGTCGGAAAGCAGCA</td>
<td>55</td>
<td>Single-base extension</td>
</tr>
<tr>
<td></td>
<td></td>
<td>OXY.MPP.4R</td>
<td>R: TTCCAAGGCCAGACCTTG</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>OXY.SNP</td>
<td>S: CTCTGAGCCTCCCTCAC</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*McCacken and Sorensen (2005); †, cuts white-headed duck allele; ‡, cuts ruddy duck allele.
agarose gel. Finally, when none of these methods could be applied (LDHB, MPP), we used a single-base extension technique, the SNaPshot kit (Applied Biosystems), for which we designed internal SNaPshot primers (Table 1), allowing the species-specific base to be identified by fluorescence in an automatic sequencer.

PCRs were carried out in 20-µL reactions containing 1x Gold Buffer (Applied Biosystems), 2.5 mM MgCl$_2$, 1 mM dNTPs (0.25 mM each), 0.5 µm forward primer, 0.5 µm reverse primer, 25–100 ng of genomic DNA and 0.7 U of AmpliTaq Gold DNA polymerase (Applied Biosystems). PCRs were performed in a GeneAmp PCR System 9700 (Applied Biosystems) or PTC-100 Programmable Thermal Controller (MJ Research) using the following conditions: one segment of 95 °C for 6 min; 35 cycles of 95 °C for 45 s, 55 °C or 60 °C (depending on primers; see Table 1) for 30 s, and 72 °C for 1 min; and a final segment of 72 °C for 7 min. In the case of PEPC9, the number of PCR cycles was set to 30. Restriction digests were in 10-µL reactions with 4 µL of PCR product, 3.5 U of restriction enzyme and the appropriate buffer (New England Biolabs). This was incubated at 37 °C for 1 h and the resulting product visualized in 2% agarose. For PEPC9 and Trop, the PCR product was directly visualized in 2% agarose. For SNaPshot reactions, 5 µL of PCR product was prepared by incubating with 1 µL of ExoSAP-IT (Amersham Biosciences) at 37 °C for 15 min and 80 °C for 15 min. Single-base extensions were in 10-µL extension reactions with 2 µL SNaPshot Ready Reaction Mix (Applied Biosystems), pooled PCR products for two loci (1 µL each) and pooled SNaPshot primers (0.06 µM of LDHB.Oxy.SNP and 0.1 µM of MPP.Oxy.SNP). Cycling conditions were 25 cycles of 96 °C for 10 s, 50 °C for 5 s, and 60 °C for 30 s. We used Sephadex G-50 fine to remove unincorporated dNTPs, and then reaction products were electrophoresed in an ABI PRISM 3100 automated sequencer (Applied Biosystems).

We tested these eight assays on a larger panel of 33 feral European ruddy ducks and 52 wild and captive white-headed ducks from Spain. The results support the conclusion that Spanish white-headed ducks and European ruddy ducks are fixed for alternative alleles at each of these loci. We also tested the performance of our methods with five captive-bred hybrids of known origin (three F$_1$s, one backcross to a white-headed duck and one backcross to a ruddy duck). Genotypes were concordant with expectations given the crosses that produced these birds, indicating that our assays allowed accurate determination of the genotypes of wild hybrids.

Typing of microsatellite alleles

We scored all individuals for 10 microsatellite loci developed specifically for these two Oxyura species (Muñoz-Fuentes et al. 2005b, 2006). Five loci from each species were derived from approximately equal numbers of candidate loci in each species (Muñoz-Fuentes et al. 2005b), thereby controlling for the ascertainment bias typically encountered in interspecific comparisons; polymorphism is often greater in the species from which a microsatellite locus is derived (Ellegren et al. 1995; Wright et al. 2004). The Spanish sample included birds that were morphologically identified as ruddy ducks (n = 31), white-headed ducks (n = 63) and hybrids (n = 29). Microsatellite data for white-headed ducks in allopatry were obtained for birds from Greece (n = 6) and the captive population in Spain (n = 19). Ruddy duck microsatellite data for birds in allopatry were obtained for individuals from France (n = 18), the UK (n = 27), two captive populations in the UK (n = 6 and 7) and North America (n = 51) (Muñoz-Fuentes et al. 2006). Microsatellite allele frequencies in the reference populations are provided as Supplementary material (Table S2).

**Data analysis**

We tested the microsatellite data for Hardy–Weinberg equilibrium and linkage disequilibrium with GENEPOP (Raymond & Rousset 1995) and we applied Bonferroni sequential correction (Rice 1989) to evaluate statistical significance when multiple simultaneous tests are performed. We checked for genotyping errors with MICROCHECKER version 2.1.1 (Van Oosterhout et al. 2004). One additional locus, Oxy4, was also typed but excluded from the analyses because it did not conform to Hardy–Weinberg expectations in ruddy ducks, and there was evidence for linkage disequilibrium in both ruddy ducks (Muñoz-Fuentes et al. 2006) and white-headed ducks.

We used a factorial correspondence analysis (FCA) as implemented in GENETIX version 4.05.2 (Belkhir et al. 1996–2004) to plot each individual in a two-dimensional space according to their microsatellite and intron allele composition independent of any a priori species designations. We used HP-RARE 1.0 (Kalinowski 2005) to correct allelic richness values for differences in sample size. One marker, Oxy14, was included for genetic diversity calculations, but dropped from subsequent analyses due to evidence of preferential amplification of the white-headed duck allele in hybrids (see Results).

We also used a Bayesian assignment method, as implemented in the software NEWHYBRIDS version 1.1 beta (Anderson & Thompson 2002), to identify pure individuals and distinguish among hybrid types. This approach makes no a priori assumptions about population allele frequencies. We ran NEWHYBRIDS for all individuals collected in Spain (presumed white-headed ducks, ruddy ducks and hybrids) and set the program to distinguish the two parental species, F$_1$s, F$_2$s, and first-generation backcrosses to each of the parental species six categories. We repeated the analysis with the program set to also recognize second-generation
backcrosses (eight categories). Some of the individuals identified as the progeny of backcrosses in the first analysis were assigned a low to moderate probability of being the result of second-generation backcrosses (Table S3, Supplementary material), but these probabilities varied depending on the priors used, suggesting that our data were not sufficient to unambiguously distinguish first- and second-generation backcrosses.

Results

Genetic diversity and divergence

The mtDNA control region sequences of white-headed ducks and ruddy ducks are unambiguously distinguished by 18 fixed differences or ∼3% sequence divergence (Table S4, Supplementary material). A single haplotype was detected in the ruddy duck population all over Europe (Muñoz-Fuentes et al. 2006), whereas three closely related white-headed duck haplotypes (maximum sequence divergence of 0.2%) are present in Spain (Muñoz-Fuentes et al. 2005a).

At 7 of 10 microsatellite loci, white-headed ducks and ruddy ducks in Spain had no alleles in common (Table S2). At the remaining loci, alleles shared with ruddy ducks were also present in white-headed ducks from Greece and the captive white-headed duck population. Likewise, at the eight intron markers, no ruddy duck alleles were detected in birds morphologically identified as white-headed ducks. Thus, there was no evidence for introgression of ruddy duck alleles into the Spanish white-headed duck population.

Based on microsatellite data, expected and observed heterozygosities (± SD) were 0.363 ± 0.080 and 0.310 ± 0.027, respectively, in ruddy ducks; and 0.183 ± 0.067 and 0.188 ± 0.016 in white-headed ducks (Table 2). In addition, the total number of alleles in ruddy ducks (n = 32 in 30 individuals) was more than twice as large as in white-headed ducks (n = 15 in 63 individuals). Multilocus genotypes in 36 of 59 (61%) white-headed ducks with complete data were identical to at least one other individual; and up to eight individuals shared the same genotype. In contrast, each ruddy duck in Spain had a unique multilocus genotype. A factorial correspondence analysis of the microsatellite and intron data (Fig. 1) reflects substantial differentiation between the two species at nuclear loci ($F_{ST} = 0.74$; $P < 0.001$) and lower genetic diversity in white-headed ducks than in ruddy ducks. The factorial correspondence analysis also showed that all but one of the individuals identified as hybrids based on morphology occupied an intermediate position between white-headed ducks and ruddy ducks (Fig. 1).

Hybrid identification and type of crosses

One microsatellite marker, Oxy14, was included for genetic diversity calculations of ruddy ducks and white-headed ducks but dropped from subsequent analyses in new hybrids due to evidence of preferential amplification of white-headed duck alleles in hybrids. Individuals 21, 22, 23, 24 and 25 tested as homozygous for allele 131 at locus Oxy14 (these individuals were genotyped twice to confirm this result), but were heterozygous with one allele coming

<table>
<thead>
<tr>
<th>Population</th>
<th>$n$</th>
<th>No. hapl.</th>
<th>$H_d$ ± SD</th>
<th>$\pi$ ± SD</th>
<th>$n$</th>
<th>AR</th>
<th>$AR_C$</th>
<th>$H_E$ ± SD</th>
<th>$H_O$ ± SD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Spanish white-headed ducks</td>
<td>39</td>
<td>2*</td>
<td>0.456 ± 0.053</td>
<td>0.0024 ± 0.0003</td>
<td>63</td>
<td>1.5</td>
<td>1.5</td>
<td>0.183 ± 0.067</td>
<td>0.188 ± 0.016</td>
</tr>
<tr>
<td>Spanish ruddy ducks</td>
<td>13</td>
<td>1</td>
<td>0.000</td>
<td>0.000</td>
<td>30</td>
<td>3.2</td>
<td>3.2</td>
<td>0.363 ± 0.080</td>
<td>0.310 ± 0.027</td>
</tr>
<tr>
<td>European ruddy duck#</td>
<td>34</td>
<td>1</td>
<td>0.000</td>
<td>0.000</td>
<td>78</td>
<td>3.4</td>
<td>3.0</td>
<td>0.369 ± 0.080</td>
<td>0.363 ± 0.017</td>
</tr>
<tr>
<td>North American ruddy ducks</td>
<td>67</td>
<td>23</td>
<td>0.824 ± 0.044</td>
<td>0.0038 ± 0.0003</td>
<td>51</td>
<td>6.3</td>
<td>5.4</td>
<td>0.488 ± 0.088</td>
<td>0.443 ± 0.022</td>
</tr>
</tbody>
</table>

$n$, sample size, number of individuals; No. hapl., number of haplotypes; $H_d$, haplotype diversity; $\pi$, nucleotide diversity; SD, standard deviation; AR, allelic richness (or mean number of alleles per locus); $AR_C$, allelic richness corrected for sample size; $H_E$, expected heterozygosity; $H_O$, observed heterozygosity; *a third white-headed duck haplotype was found among the sampled hybrids (Muñoz-Fuentes et al. 2005a); †includes the Spanish ruddy ducks.
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Fig. 2 Multilocus genotypes of individuals morphologically identified as hybrids. White and black indicate an allele (or haplotype) found in white-headed ducks or in ruddy ducks, respectively. Grey indicates an allele found both in white-headed ducks and ruddy ducks (this applies only to microsatellites as the intron markers represent fixed differences between the species). Capital letters and numbers stand for locus names of introns and microsatellites, respectively (see Material and methods and Table 1). Numbers inside the circles indicate the mtDNA haplotype found in that bird (1, Oleu_01; 2, Oleu_02; 3, Oleu_03 (Muñoz-Fuentes et al. 2005a); all feral European ruddy ducks carried the same haplotype, Ojam_01 (Muñoz-Fuentes et al. 2006)). Large coloured squares separate hybrid types based on results from the program NEWHYBRIDS. Sample ID numbers increase with collection date (1 being the earliest, 1993). Four individuals, marked with (a), were identified by NEWHYBRIDS as having a low to moderate probability of being second-generation backcrosses. Rd, ruddy duck; F1, first generation hybrid; Whd_Bc, progeny of F1 × white-headed duck cross; Rd_Bc, progeny of F1 × ruddy duck cross; m, male; f, female.

from each species (or, in a few cases, homozygous for a microsatellite allele present in both species) at all other loci; or in the case of individual 25, all but one other locus (see Fig. 2). Because allele 131 has not been previously observed in European ruddy ducks, these genotypes are highly unlikely. First, it is unlikely that a first generation backcross would have a genotype consistent with an F1 hybrid at a sample of 16 loci but be homozygous for a species-specific allele at only one locus ($P = 0.00014$). Such a genotype is even less likely in a subsequent generation backcross. Second, it is highly unlikely to sample five different individuals that have this same pattern and which are all homozygous at the same locus ($P = 6 \times 10^{-20}$). The above calculations assume that the white-headed duck allele (length 131) does not occur in Spanish ruddy ducks. If these genotypes were correct, then the sample
Discussion

The native white-headed duck and the introduced ruddy duck

Substantial divergence in mtDNA control region sequences and fixed differences between white-headed ducks and ruddy ducks across multiple nuclear introns support the conclusion that they are distinct species (Livezey 1995; McCracken et al. 2000; McCracken & Sorenson 2005). Indeed, the two species are not each other’s closest relatives and their most recent common ancestor dates to perhaps 2 million years ago (McCracken & Sorenson 2005). These results negate suggestions in the popular media that white-headed ducks and ruddy ducks are not distinct species (see McCracken et al. 2000).

Based on a sample of 10 microsatellite loci, including five obtained from each species, we found that white-headed ducks had lower genetic diversity than the introduced ruddy duck population in Europe. This result is particularly striking considering that the European ruddy duck population apparently derives from just seven founders (Hudson 1976; Muñoz-Fuentes et al. 2006). Rapid population growth following the founding event likely minimized genetic drift and the loss of genetic diversity in ruddy ducks (Muñoz-Fuentes et al. 2006), whereas a severe bottleneck following a long period of declining population size resulted in a greater reduction in genetic diversity in white-headed ducks. Loss of genetic diversity compromises a species evolutionary potential and increases extinction risk (Frankham et al. 2002).

Morphological and genetic identification of parental species and hybrids

All individuals identified either as ruddy ducks or hybrids during the control programme were confirmed to be such genetically, except for one individual (individual 20, see Figs 1 and 2). Hybrid identification during the control programme followed a key based on morphological criteria such as plumage characteristics and beak shape (Urdiales & Pereira 1993), which performed well in identifying hybrids, but failed to identify different hybrid categories (unpublished data).

Individual 20 (Figs 1 and 2) had a phenotype generally similar to a ruddy duck, but with a completely black head and unusual bill shape and was, due to its unusual characteristics, misidentified as a hybrid. Genetic analysis indicated it to be a ruddy duck, with a ‘unique’ microsatellite allele, not found in any other ruddy duck, either in Europe or North America (n = 142) (Muñoz-Fuentes et al. 2006), or in white-headed ducks. It is highly unlikely that all of the remaining microsatellite alleles and nuclear introns would match a ruddy duck genotype if this bird were not a ruddy
duck. Therefore, we conclude that this bird is a ruddy duck with variant plumage.

**Hybridization in the wild**

Microsatellite markers and fixed differences in nuclear introns allowed us to differentiate among first-generation hybrids, the progeny of backcrossing, and pure ruddy ducks and white-headed ducks. Our results demonstrate that first generation hybrids are fertile and mate with individuals of both parental species in the wild. It is possible that our sample of hybrids includes some siblings, but based on sampling dates and localities (see Table S5, Supplementary material), and mtDNA haplotypes (Fig. 2), the individuals identified as backcross progeny were produced in at least two and three independent backcrosses to ruddy ducks and white-headed ducks, respectively. Therefore, backcrossing is not an isolated phenomenon.

Because the majority (89%) of F1 hybrids had ruddy duck mtDNA, most crosses were between female ruddy ducks and male white-headed ducks. This strong asymmetry in the direction of hybridization is likely a consequence of the efficiency of the eradication programme in eliminating male ruddy ducks (males of the two species are easier to differentiate than females), rather than a behavioural asymmetry in the biology of hybridization (Green & Hughes 2001). This effect, however, has likely changed through time: the proportion of females among culled ruddy ducks and hybrids was significantly greater from June 2000 to May 2003, when the control team became nationally coordinated and its effectiveness presumably increased, than from June 1991 to May 2000 ($\chi^2 = 2.9$ for the change of sex ratios between the two periods, $P < 0.05$; Calzada et al. 2003).

A contrasting pattern was observed among individuals identified as the progeny of subsequent generation backcrosses to one of the two parental species. All 10 of these individuals had white-headed duck mtDNA (Fig. 2). Thus, our data provide no evidence of reproduction by F1 females with ruddy duck mtDNA, even though most F1 hybrids had ruddy duck mtDNA. In contrast, the three individuals produced by backcrosses to ruddy ducks had white-headed duck mtDNA, indicating that female hybrids with white-headed duck mtDNA are fertile and successfully reproduced with male ruddy ducks. Likewise, seven individuals that resulted from F1’s backcrossing with white-headed ducks had white-headed duck mtDNA. These individuals may have been produced by either a female white-headed duck mating with a male hybrid or a female hybrid with white-headed duck mtDNA mating with a male white-headed duck.

The production of ruddy duck backcrosses with white-headed duck mtDNA demonstrates conclusively that female hybrids with white-headed duck mtDNA are fertile, whereas direct evidence that female hybrids with ruddy duck mtDNA are fertile is still lacking. Again, this result contrasts with the preponderance of ruddy duck mtDNA among our sample of F1 hybrids. Thus, in accord with Haldane’s rule (in birds, females are heterogametic) and as observed in other birds (Price & Bouvier 2002), hybrid females with ruddy duck mtDNA may have low fertility or are perhaps infertile. Alternatively, given our relatively small sample of hybrids, the absence of subsequent generation hybrids with ruddy duck mtDNA may be due to chance. Future research on ruddy ducks and white-headed ducks should test for hybrid infertility due to genomic incompatibilities that depend on the direction of the cross (Orr 1997; Price & Bouvier 2002).

**Conservation implications**

Although we detected backcrossing between hybrids and white-headed ducks (Fig. 2), we found no evidence of ruddy duck alleles in individuals that were morphologically identified as white-headed ducks ($n = 63$ using microsatellite data, including 43 for which we also gathered intron data). Based only on our intron data, for which there are no alleles shared between the two species, and given that none of the 43 birds identified as white-headed ducks had any ruddy duck alleles at a total of eight loci, the 95% confidence interval on the proportion of ruddy duck genes in the white-headed duck population ($0$ of 688 typed alleles, i.e. 43 individuals $\times$ 8 loci $\times$ 2 alleles per locus) is $0$–$0.7\%$. Likewise, the 95% confidence interval on the proportion of white-headed ducks that have one or more ruddy duck alleles at one or more of the loci we typed ($0$ of 63 birds typed) is $0$–$6\%$. Thus, the continuing ruddy duck eradication programme in Spain has apparently been effective in limiting introgression to date. Our results, however, are alarming: despite an aggressive control programme in place since 1991, backcrosses to both parental species have been found in Spain. These results suggest that there would be a significant risk of widespread introgression if this programme is not continued, and particularly if ruddy ducks’ numbers keep increasing in other parts of Europe. In addition, the situation may be exacerbated by the competitive asymmetry between the expanding population of an invasive species and a recovering native species with greatly reduced genetic diversity. If left to proceed without control, hybridization and expansion of the ruddy duck will likely cause the loss of species-specific characters and adaptations in white-headed ducks and eventually threaten the species with extinction. Alternatively, the result may be a hybrid swarm as recorded in New Zealand as a result of mallard and grey duck, hybridization (Gillespie 1985; Rhymer et al. 1994).

The fact that we did not detect ruddy duck alleles in white-headed ducks indicates that introgression, if it has
occurred, remains at low frequency and that there is still
time to prevent the extinction of the white-headed duck
through hybridization. The maintenance of the white-
headed duck’s genetic integrity will depend on the con-
tinued success of a programme for the elimination of
hybrids and ruddy ducks. In the absence of a ruddy duck eradication programme, as now implemented in the UK,
France and Spain, it is likely that patterns of hybridization
might be different. In the UK, the main source population
of ruddy ducks in Europe, growth rate is estimated at
18% per annum and the number of countries in which the
ruddy duck has been recorded since first breeding in the
wild in the UK in 1960 is increasing (Hughes et al. 2004).
If both the white-headed duck and ruddy duck are to be
preserved, these species cannot co-exist in the same areas
and action against the ruddy duck in Europe should be
intensiﬁed, especially in the UK. The plight of the Spanish
white-headed duck exempliﬁes how human actions in
one country can have dramatic trans-boundary effects, how
different perturbations can act synergistically to decrease
the chances of long-term survival for wild species, and how
coordinated international conservation plans are needed
for effective biodiversity conservation.

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Supplementary material

The supplementary material is available from
http://www.blackwellpublishing.com/products/journals/
suppmat/MEC/MEC3170/MEC3170sm.htm

Table S1 Primer names and sequences used in this study to
amplify and sequence introns at the loci indicated. An asterisk
following the locus name designates loci not selected for genotyping
hybrids because either ampliﬁcation or sequencing was unsuccessful,
or there were no species-diagnostic differences.

Table S2 Allele frequencies at 10 microsatellite loci in white-
headed ducks by population (Spain, Greece, captive) and feral
ruddy ducks from Spain. Sample size (n) is indicated below the
population name. Locus names are in bold and alleles are in italics,
designated by their size in base pairs. For white-headed ducks,
private alleles (present only in one population) appear in bold.
Acebuche stands for the captive population sampled from El
Acebuche in Doñana National Park, Spain, established before
hybridization was detected in Spain. The small number of alleles
shared between ruddy ducks and white-headed ducks in Spain
were also shared between allopatric populations of both
species. Alleles marked with (*) were observed in ruddy ducks
from North America (Muñoz-Fuentes et al. 2006), but in Europe
were found only in white-headed ducks.

Table S3 Probabilities that each individual belongs to a given
hybrid category as estimated by newhybrids.

Table S4 Variable nucleotide positions in mtDNA control region
sequences of white-headed ducks and ruddy ducks in Europe
(1993–2003). Eighteen ﬁxed differences between the two species
are marked with asterisks. Total number of sites is 576; ‘.’ denotes
an identical base and ‘−’ an indel.

Table S5 Hybrid samples used in this study. Sample ID, collection
date, origin and age for each hybrid individual are provided, as
well as the corresponding number code used in the main text. ad,
adult; juv, juvenile.

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