# A Worldwide Perspective on the Population Structure and Genetic Diversity of Bottlenose Dolphins (Tursiops truncatus) in New Zealand

Gabriela Tezanos-Pinto, Charles Scott Baker, Kirsty Russell, Karen Martien, Robin W. Baird, Alistair Hutt, Gregory Stone, Antonio A. Mignucci-Giannoni, Susana Caballero, Tetusya Endo, Shane Lavery, Marc Oremus, Carlos Olavarría, and Claire Garrigue

From the University of Auckland, School of Biological Sciences, Private Bag 92019, Auckland, New Zealand (Tezanos-Pinto, Baker, Russell, Caballero, Lavery, Oremus, and Olavarría); the NOAA, Southwest Fisheries Science Center, 8604 La Jolla Shores Drive, La Jolla, CA 92037 (Martien); the Cascadia Research Collective, 218 1/2 W, 4th Avenue, Olympia, WA 98501 (Baird); the Department of Conservation, Private Bag 4715, Christchurch 8140, New Zealand (Hutt); the New England Aquarium, Boston, MA 02110 (Stone); the Red Caribeña de Varamientos, PO Box 361715, San Juan, PR 00936-1715 (Mignucci-Giannoni); the Health Sciences University of Hokkaido, Ishikari-Tobetsu, Hokkaido 061-0293, Japan (Endo); the Centre de Recherches Insulaires et Observatoire d'Environnement BP 1013, Papetoai Moorea, French Polynesia (Oremus); the Centro de Estudios del Cuaternario, Plaza Munoz Gamero 105, Punta Arenas, Chile (Olavarría); and the Opération Cétacés, BP12827, 98802, Noumea, New Caledonia (Garrigue).

Address correpondence to G. Tezanos-Pinto at the address above, or e-mail: g.tezanospinto@auckland.ac.nz.

## **Abstract**

Bottlenose dolphins (*Tursiops truncatus*) occupy a wide range of coastal and pelagic habitats throughout tropical and temperate waters worldwide. In some regions, "inshore" and "offshore" forms or ecotypes differ genetically and morphologically, despite no obvious boundaries to interchange. Around New Zealand, bottlenose dolphins inhabit 3 coastal regions: Northland, Marlborough Sounds, and Fiordland. Previous demographic studies showed no interchange of individuals among these populations. Here, we describe the genetic structure and diversity of these populations using skin samples collected with a remote biopsy dart. Analysis of the molecular variance from mitochondrial DNA (mtDNA) control region sequences (n = 193) showed considerable differentiation among populations ( $F_{ST} = 0.17$ ,  $\Phi_{ST} = 0.21$ , P < 0.001) suggesting little or no female gene flow or interchange. All 3 populations showed higher mtDNA diversity than expected given their small population sizes and isolation. To explain the source of this variation, 22 control region haplotypes from New Zealand were compared with 108 haplotypes worldwide representing 586 individuals from 19 populations and including both inshore and offshore ecotypes as described in the Western North Atlantic. All haplotypes found in the Pacific, regardless of population habitat use (i.e., coastal or pelagic), are more divergent from populations described as inshore ecotype in the Western North Atlantic than from populations described as offshore ecotype. Analysis of gene flow indicated long-distance dispersal among coastal and pelagic populations worldwide (except for those haplotypes described as inshore ecotype in the Western North Atlantic), suggesting that these populations are interconnected on an evolutionary timescale. This finding suggests that habitat specialization has occurred independently in different ocean basins, perhaps with Tursiops aduncus filling the ecological niche of the inshore ecotype in some coastal regions of the Indian and Western Pacific Oceans.

**Key words:** tursiops truncatus, bottlenose dolphin, ecotype, population structure, genetic diversity, New Zealand

All cetaceans including baleen whales, beaked whales, dolphins, and porpoises are highly mobile and many species undertake long-distance seasonal migrations (Baker et al. 1993; Rosel et al. 1999; Wells et al. 1999). This mobility has

the potential to reduce the isolation and therefore the genetic differentiation in haplotype frequencies among regional populations. However, several studies have revealed demographic isolation (Würsig and Jefferson 1990;

Rossbach and Herzing 1999) or genetic differentiation at both the haplotype and nucleotide level among neighboring dolphin populations, despite no obvious physical barriers to interchange (e.g., Dowling and Brown 1993; Hoelzel 1998; Hoelzel et al. 1998; Pichler et al. 1998; Krützen et al. 2004; Oremus et al. 2007).

The bottlenose dolphin (Tursiops truncatus) occupies a wide range of coastal and pelagic habitats throughout tropical and temperate waters around the world (Leatherwood et al. 1983). At least one related species (currently Tursiops aduncus, although perhaps not a truly congener; refer to LeDuc et al. 1999; Wang et al. 1999; Natoli et al. 2004) is sympatric with T. truncatus along the coast of mainland China, in the Taiwan Strait (Wang et al. 1999), around Australia (Moller and Beheregaray 2001; Krützen et al. 2004) and off South Africa (Ross 1977; Ross and Cockcroft 1990; Natoli et al. 2004).

It appears that *T. truncatus* may have once or repeatedly, adapted to different environmental conditions resulting in several different forms or "ecotypes." In the North Atlantic, for example, Duffield et al. (1983) described 2 T. truncatus ecotypes based on hematology profiles and distribution: "inshore" and "offshore." Later studies confirmed this finding with independent lines of evidence from morphology, genetics, parasite load, and diet (Hersh and Duffield 1990; Mead and Potter 1990; Hoelzel et al. 1998; Natoli et al. 2004). In many regions of the world, however, there is insufficient evidence to distinguish between differential habitat use by individuals and true ecotype specialization of particular bottlenose dolphin genetic lineages. Distinct parapatric (adjacent) populations have been documented in the Western North Atlantic (Duffield et al. 1983; Hersh and Duffield 1990; Hoelzel et al. 1998; Torres et al. 2003; Kingston and Rosel 2004; Natoli et al. 2004) and to a lesser extent in the Eastern North Pacific (ENP), the Gulf of California (Lowther 2006; Segura et al. 2006), as well as along the western coast of South America (Van Waerebeek et al. 1990; Sanino et al. 2005).

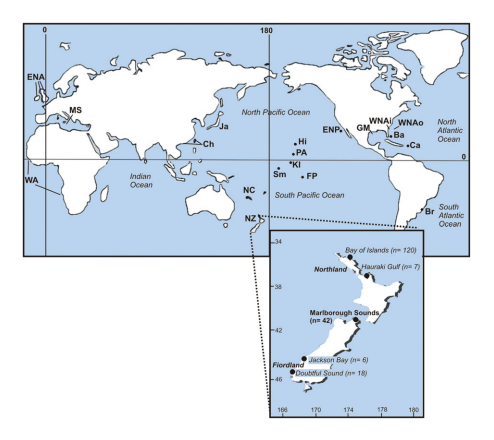
Although it is generally assumed that the inshore ecotype inhabits coastal areas whereas the offshore ecotype inhabits pelagic waters, this assumption can be misleading: individuals described as the offshore ecotype have been reported close to shore in some areas (Wells et al. 1999), and individuals described as the inshore ecotype have been observed far from shore in regions where the continental shelf is broad (Kenney 1990). Moreover, around many islands in the Pacific Ocean, deep ocean habitats are found in close proximity to shallow coastal areas. Information on population structure and ecotype assignment of bottlenose dolphins from these islands has been limited to 1 or 2 populations with small sample sizes (Natoli et al. 2004). Further, there has been some confusion between the inshore ecotype of T. truncatus and the more coastal species of Indo-pacific bottlenose dolphin, T. aduncus (Reeves et al. 2004). For example, mitochondrial DNA (mtDNA) control region sequences of individuals from a coastal South African population previously reported to represent the inshore ecotype of T. truncatus (Goodwin et al. 1996; Smith-Goodwin 1997) were recently shown to match a sequence

of the *T. aduncus* holotype (collected along the Ethiopian coast of the Red Sea; Perrin et al. 2007).

On a worldwide scale, pelagic T. truncatus seem to be characterized by high levels of genetic diversity, whereas coastal populations are characterized by low levels of genetic diversity (Natoli et al. 2004). Moreover, pelagic populations are likely to be the source of independent founder events that have generated somewhat discrete population segments in coastal areas perhaps as a result of resource specialization or philopatry (Hoelzel 1998; Natoli et al. 2004). Intensively studied populations in the Western North Atlantic (WNA) are commonly used as a model for comparison with other regions (Curry 1997; Curry and Smith 1997; Hoelzel et al. 1998; Natoli et al. 2004). However, considering the limited nature of studies conducted in the Central and Western Pacific (CWP) and the taxonomic uncertainty in some studies, it is unknown whether the pattern found in the WNA is representative of the worldwide population structure of the species or if it represents only an ocean basin, or even a region within an ocean. Although, testing of this hypothesis was initiated by Natoli et al. (2004), their sample size for the Pacific Ocean was limited to 18 samples from only 2 regions (1 from the ENP and 17 from China).

In New Zealand waters, bottlenose dolphins are found both in coastal and pelagic habitats (Constantine 2002); but as yet, there has been no independent evidence to classify individuals or populations as genetically more similar to inshore or offshore ecotypes found in other regions. In coastal waters, there are 3 discontinuous populations found in Northland, Marlborough Sounds, and Fiordland (Bräger and Schneider 1998; Schneider 1999; Constantine 2002, Figure 1). The Fiordland population appears to be further subdivided into 3 small communities inhabiting Milford, Doubtful, and Dusky Sounds (Boisseau 2003). Long-term studies using mark-recapture models have estimated abundance of around 446 adults for Northland (confidence interval [CI] = 418–487; Constantine 2002) and around 49 individuals for Doubtful Sound (coefficient of variation = 0.024; Gormley 2002). To date, there is no estimate for Marlborough Sounds; but a photo-identification catalog (Merriman et al. 2005) suggests a population of at least several hundreds. Comparison of individual identification photographs between Northland, Marlborough Sounds, and Fiordland suggests no exchange of individuals among populations (Bräger and Schneider 1998; Schneider 1999; Constantine 2002).

Here, we describe the population structure and genetic diversity of coastal bottlenose dolphins in New Zealand waters based on mtDNA control region sequences. We also describe the genetic relationship of New Zealand bottlenose dolphins to other *T. truncatus* from 18 regions worldwide, including the inshore and offshore ecotypes as described in the Western North Atlantic. With this more comprehensive dataset, we further test the hypothesis that the genetically distinct ecotypes reported in the Western North Atlantic are found worldwide, predicting that New Zealand coastal dolphins would group genetically with individuals representative of the inshore ecotype given their coastal habitat



**Figure 1.** Locations represented by genetic samples of bottlenose dolphins (*Tursiops truncatus*) including New Zealand populations (insert). ENA = Eastern North Atlantic, MS = Mediterranean Sea, WA = West Africa, Ja = Japan, Ch = China—Taiwan, Hi = Hawaiʻi, PA = Palmyra Atoll, KI = Republic of Kiribati, Sm = Samoa, FP = French Polynesia, NC = New Caledonia, NZ = New Zealand, ENP = Eastern North Pacific, GM = Gulf of Mexico, WNAi = Western North Atlantic inshore, WNAo = Western North Atlantic offshore, Ba = Bahamas, Ca = Caribbean, and Br = Brazil.

preference or use. Alternatively, we considered that New Zealand dolphins have adopted a coastal habitat use independent of other coastal or insular populations, perhaps originating from a widespread, pelagic population, or complex of populations. The results provide new insights into the pattern of mtDNA diversity associated with habitat specialization and ecotype formation among *T. truncatus* worldwide.

## **Materials and Methods**

## New Zealand Dataset

A total of 193 samples were collected from bottlenose dolphins in coastal habitats around New Zealand (Figure 1) using a Paxarm biopsy sampling system (Krützen et al. 2002). Of these, 127 samples were from 2 locations in Northland (Bay of Islands 35°14′S, 174°06′E, n = 120 and Hauraki Gulf 36°40′S, 174°50′E, n = 7). Forty-two samples were collected from Marlborough Sounds (41°05′S, 174°15′E), 18 from Doubtful Sound in Fiordland (45°17′S, 167°168′E), and 6 from the neighboring Jackson Bay (44°S, 168°36′E). Analysis of individual identification photographs confirmed that some individuals photographed in Jackson Bay belonged to the Milford Sound community;

therefore, samples collected in this area were assigned to the Fiordland population. Sixteen samples were obtained from strandings around New Zealand; these sequences were included in the worldwide analyses but not in the analyses of population structure for New Zealand as the assignment of individuals to populations was not possible.

#### Pacific Ocean Dataset

Excluding samples collected in New Zealand, a total of 218 samples representing 62 unique mtDNA control region sequences (i.e., haplotypes) were available from 8 populations from the CWP and 1 haplotype (represented by one sample) was available from the ENP. Haplotype sequences were obtained from published sequences, biopsy samples, "whale meat" products, and GenBank sources (Figure 1, Supplementary Appendix 1). From the CWP, 155 skin samples were collected using a biopsy sampling system; of those, 23 were collected from the Republic of Kiribati (Phoenix Archipelago, 2°49′S, 171°40′W), 117 from the main Hawaiian Islands (Oʻahu, Hawaiʻi, Kauaʻi, and Niʻihau, 19°N–22°N, 156°W–160°W), 11 from the Palmyra Atoll (5° 52′N, 162° 06′W), 1 from Samoa (13°25′S, 172°36′W), 2 from French Polynesia (Tuamotu Archipelago 15°S,

148°W), and 1 from New Caledonia (22°51′S, 167°42′E; Figure 1, Supplementary Appendix 1). Previously unpublished sequences from 34 whale meat products identified as *T. truncatus* were obtained from commercial markets of Japan as part of the ongoing molecular monitoring of whale and dolphin products (Baker and Palumbi 1994; Baker et al. 2000; Endo et al. 2005). Most market products from dolphins were supplied by small-type coastal whaling (Endo et al. 2003) and therefore were assumed to originate from coastal areas around Japan.

Six mtDNA haplotype sequences of T. truncatus were obtained from GenBank (accession numbers: AF056231 and AF049101 from Wang et al. 1999; AF459508, AF459509, AF459523, and AF459522 from Ji GQ, Yang G, Liu S, Zhou KY, unpublished data). Additionally, 24 samples representing 19 unique haplotype sequences were reconstructed from 3 publications (Wang et al. 1999; Kakuda et al. 2002; Natoli et al. 2004) representing 3 regions (Japan, China-Taiwan, and ENP; Supplementary Appendix 1). Each publication included one reference sequence (published in GenBank or included in the publication) with a table of variable sites and haplotype frequencies. Haplotype sequences were reconstructed from these by inserting and aligning the reference sequence with the existing T. truncatus dataset using MacClade software Vs. 4.06 (Maddison WP and Maddison DR 2003).

## Atlantic Ocean Dataset

A total of 158 samples representing 50 unique mtDNA haplotype sequences were available from 9 populations in the Atlantic Ocean from published sequences, strandings, and GenBank sources (Figure 1, Supplementary Appendix 1). For this study, 12 samples from Puerto Rico (17°N-18°N, 65°W-67°W) and 1 from the United States Virgin Islands (17°41.23'N, 64°49.32'W) were newly available from stranded individuals. Three haplotype sequences from the Bahamas were obtained from GenBank (accession numbers: AF155160, AF155161, and AF155162 from Parsons et al. [1999]). Additionally, 142 samples representing 37 haplotype sequences were reconstructed from 3 publications (Smith-Goodwin 1997; Parsons et al. 2002; Natoli et al. 2004) representing 8 regions and 2 ecotypes (Figure 1, Supplementary Appendix 1). Haplotype sequences were reconstructed following the procedure described above.

# DNA Extraction, Polymerase Chain Reaction Amplification, and Sequencing

For tissue obtained from biopsy samples and stranded specimens, total genomic DNA was isolated from tissue samples using proteinase K digestion followed by standard phenol/chloroform methods (Sambrook et al. 1989), as modified for small tissue samples by Baker et al. (1994). Amplification of 800 bp of the mtDNA control region was performed using the primers light-strand tPro-whale Dlp-1.5 with the addition of an M13 tag to the 5' end (Dalebout et al. 1998) and heavy-strand Dlp-8G (Pichler et al. 2001). Polymerase chain reaction (PCR) volume was 15 µl per

reaction per sample. PCR conditions were as follows: 0.2 mM deoxynucleoside triphosphate, 2.5 mM MgCl, 1X PCR buffer, 0.4 µM of each primer, and 0.05 µl Platinum *Taq* (Invitrogen, Auckland, New Zealand). PCR cycling profile was 2 min at 94 °C, 35 cycles of 30 s at 94 °C, 40 s at 55 °C, and 40 s at 72 °C. PCR products were purified using ExoI and SAP (Werle et al. 1994) and sequenced with BigDye terminator chemistry using ABI 377 and ABI 3100 DNA sequencers (Applied Biosystem, Foster City, CA). Cycle sequencing used the primer tPro-whale Dlp-1.5. Variable sites of unique haplotypes were confirmed by sequencing the heavy strand using primer Dlp-8G.

For tissue obtained from Japanese whale meat markets, DNA extractions and initial PCR amplifications were conducted using "portable" PCR protocols (e.g., Baker and Palumbi 1994; Baker et al. 2006). In brief, tissue from each product was prepared for PCR amplification using Chelex resin (BioRad Laboratories, Hercules, CA) following Walsh et al. (1991). To comply with Convention on International Trade in Endangered Species (CITES) restrictions (Bowen and Avise 1994; Jones 1994), amplified products were isolated from "native" DNA by biotin labeling of one primer and binding to streptavidin-coated plates (Baker et al. 2006).

#### Taxonomy, Ecotype, and Habitat Classification

In order to avoid potential confusion with T. aduncus, sequences from biopsy samples, strandings specimens, and whale meat products were first compared with sequences from voucher specimens of T. truncatus available from the Witness for the Whales database (Vs. 4.3) within the Web-based program DNA-surveillance (Ross et al. 2003). Sequences used in the worldwide comparison were categorized into previously described ecotypes (i.e., inshore or offshore) by reviewing each published article for independent evidence from at least 2 sources (e.g., molecular or biochemical markers, diet, morphology). However, in some publications, the terms inshore or offshore were used with no evidence other than distribution. We considered that this evidence of classification by habitat (i.e., coastal or pelagic) was insufficient for classification of ecotype. All haplotype sequences from the Western North Atlantic inshore (WNAi), Bahamas, and Gulf of Mexico presented consistent diagnosis as the inshore ecotype, whereas haplotype sequences from the Western North Atlantic offshore (WNAo) presented evidence for diagnosis as the offshore ecotype. Haplotype sequences from all remaining populations were diagnosed as "unknown" in regards to ecotype. Regional populations were also grouped into 3 ocean basins: North Pacific (NP), South Pacific (SP), and Atlantic Ocean (AO; Table 1).

## Sequences Analysis and Phylogenetic Reconstruction

Sequence alignments were performed using Sequencher (Vs. 4.1.2, Genes Codes Corp., Ann Arbor, MI) and edited manually. Unique haplotypes were identified using the software MacClade Vs. 4.06 (Maddison WP and Maddison

**Table 1.** Summary of mtDNA control region sequences available for *Tursiops truncatus* populations worldwide, showing the total number of samples (n), number of haplotypes, sequences length (No. of bp), published ecotype origin (when available), and genetic diversity values

Population	n	No. of haplotypes	No. of bp	Ecotype	Nucleotide diversity $(\pi)$ %	Haplotypic diversity (h)	Source
South Pacific							
New Zealand (NZ)	209	22	391	U	$2.2 \pm 1.1$	$0.91 \pm 0.007$	BS, St
Republic of Kiribati (KI)	23	8	388	U	$0.6 \pm 0.3$	$0.83 \pm 0.05$	BS
New Caledonia (NC)	1	1	386	U	n/a	n/a	BS
Samoa (Sm)	1	1	391	U	n/a	n/a	BS
French Polynesia (FP)	2	2	391	U	n/a	n/a	BS
North Pacific							
Hawai'i (Hi)	117	19	385	U	$2.2 \pm 1.1$	$0.87 \pm 0.016$	BS
Palmyra Atoll (PA)	11	7	385	U	$1.6 \pm 0.9$	$0.93 \pm 0.06$	BS
China (Ch)	22	17	391	U	$1.8 \pm 1$	$0.95 \pm 0.04$	RS (1), GB
Japan (Ja)	41	19	387	U	$1.3 \pm 0.7$	$0.77 \pm 0.07$	RS (2), MP
Eastern North Pacific (ENP)	1	1	297	U	n/a	n/a	RS (3)
Atlantic							
Gulf of Mexico (GM)	10	6	297	I	$0.7 \pm 0.5$	$0.84 \pm 0.1$	RS (3)
Caribbean (Ca)	13	6	387	U	$2.2 \pm 1.2$	$0.82 \pm 0.08$	St
Bahamas (Ba)	7	5	297	I	$0.5 \pm 0.3$	$0.86 \pm 0.14$	RS (3), GB
Western North Atlantic inshore (WNAi)	29	6	297	I	$0.7 \pm 0.5$	$0.43 \pm 0.11$	RS (3)
Western North Atlantic offshore (WNAo)	25	11	297	O	$2.2 \pm 1.2$	$0.88 \pm 0.05$	RS (3, 5)
Eastern North Atlantic (ENA)	38	8	297	U	$0.9 \pm 0.5$	$0.73 \pm 0.047$	RS (3, 4)
Mediterranean Sea (MS)	18	11	294	U	$2.1 \pm 1.2$	$0.94 \pm 0.03$	RS (3)
West Africa (WA)	16	5	297	U	$1.5 \pm 0.9$	$0.72 \pm 0.097$	RS (3)
Brazil (Br)	2	1	297	U	n/a	n/a	RS (1)

bp, base pairs; I, inshore; O, offshore; U, unknown; and n/a, not available. Source: BS, biopsy samples; St, strandings; GB, GenBank sequences; MP, market products; and RS, reconstructed sequences. References: 1) Wang et al. 1999; 2) Kakuda et al. 2002; 3) Natoli et al. 2004; 4) Parsons et al. 2002; and 5) Smith-Goodwin 1997.

DR 2003). The neighbor-joining (NJ) algorithm, as implemented in the software PAUP\* Vs. 4.0b10 (Swofford 2000), was used to reconstruct the phylogenetic relationships among New Zealand haplotypes. Bootstrap confidence estimates were based on 1000 replicates (Felsenstein 1985); the best fitting model of sequence evolution was found using Modeltest Vs. 3.7 (Posada and Crandall 1998). A maximum parsimony (MP) tree was also constructed using the branch and bound algorithm to search through numerous equally parsimonious trees. Because of the poorly resolved phylogeny within the subfamily *Delphininae* (LeDuc et al. 1999; Caballero, Jackson, et al. 2008), we chose a more distantly related species from the subfamily *Stenoninae*, the roughtoothed dolphin (*Steno bredanensis*; Oremus 2008), as an outgroup for all reconstructions (Caballero, Jackson, et al. 2008).

#### Population Structure and Genetic Diversity

Arlequin Vs. 2.001 (Schneider et al. 2000) was used to calculate  $F_{\rm ST}$ ,  $\Phi_{\rm ST}$ , b (haplotype diversity, Nei 1987), and  $\pi$  (nucleotide diversity, Tajima 1983) using Tamura-Nei distance correction (Tamura and Nei 1993). The significance of departure from a random distribution was evaluated using 10 000 permutations among individuals between populations (analysis of the molecular variance [AMOVA], Excoffier et al. 1992). An exact test of population differentiation based on haplotype frequencies (Raymond

and Rousset 1995) was performed to test the null hypothesis of random distribution of individuals between pairs of populations. Populations with less than 5 samples were excluded from the test of differentiation. Sequential Bonferroni corrections were applied to pairwise comparisons where indicated (Rice 1989).

## New Zealand Compared with Worldwide Populations

In order to compare New Zealand populations with the worldwide dataset, average gross  $(d_{xy})$ , and net  $(d_a)$  sequence divergence between populations and sequence diversity within populations  $(d_x, d_y)$  were estimated with Tamura–Nei distance correction, including calculation of standard errors using Mega 2.1. In order to better visualize the similarity of the New Zealand populations to the worldwide dataset, a mid-rooting dendrogram was built with Mega 2.1 (Kumar et al 2001) by NJ using net sequence divergence data  $(d_a)$  among populations.

## Migration Rates among New Zealand Populations

Asymmetric female migration rates among populations were estimated using a Markov Chain Monte Carlo (MCMC) coalescent genealogy as implemented in the software Lamarc Vs. 2.0.1 (Kuhner 2006). Bayesian and maximum likelihood (ML) analyses were employed using 5 replicates per run over 5 different runs, implementing one initial and

final chain, a different random number seed, and 5 heating temperatures (1, 1.1, 1.2, 1.3, and 1.4) for each run. The burn-in option was used to allow the first 5% of each chain to be discarded and avoid unreasonable results as recommended in Kuhner et al. (2005). In order to estimate migration rates with accuracy in reasonable time, the sample size for Northland was reduced to n = 70 by random selection (Kuhner et al. 2005).

## Worldwide Phylogeography

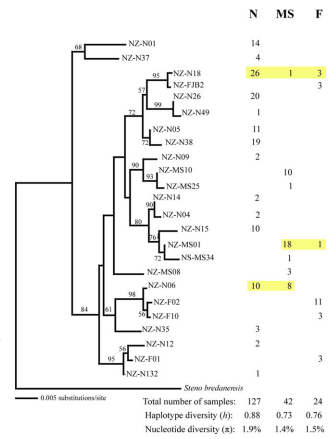
A network of the worldwide haplotype dataset was constructed using the statistical parsimony methodology described in Templeton et al. (1992), as implemented in the software TCS Vs. 1.13 (Clement et al. 2000). This method estimates an unrooted tree and provides a 95% plausible set for all sequence type linkages within the tree, with gaps considered as a fifth character state. To resolve any ambiguities (loops), we used the 3 criteria derived from the coalescent theory (Crandall and Templeton 1993; Templeton and Sing 1993; Crandall et al. 1994): 1) "frequency": high-frequency sequences are more likely to have been present in the population for a longer period of time; therefore, low-frequency sequences are more likely to be connected to sequences with high frequency; 2) "topology": sequences are more likely to be connected to interior sequences than to tip sequences; and 3) "geography": sequences are more likely to be connected to sequences from the same population or region, rather than to sequences occurring in distant populations.

## **Results**

# Phylogeography, Genetic Diversity, and Female Migration Rates among New Zealand Populations

Analysis of the 647-bp consensus fragment from the mtDNA control region sequences (n = 193; 16 samples from strandings were excluded from this analysis; refer to Materials and Methods) representing the 3 New Zealand populations revealed 24 unique maternal lineages (haplotypes; GenBank accession numbers: EU276389-EU276412), defined by 52 variable sites. Overall, there were 46 transition substitutions, 5 transversion substitutions (including one site with both a transition and transversion), and 2 single base insertion-deletions. The model of sequence evolution best fitting the dataset was HKY + I (Hasegawa et al. 1985). The estimated  $T_v/T_i$  ratio was 49.3, and estimated proportion of invariable sites (I) was 0.91.

Phylogenetic reconstructions (both NJ and MP) did not show a pattern of reciprocal monophyly or fixed nucleotide differences among populations, although strong frequency differences were observed. Most haplotypes were found in only one region: 15 unique to Northland, 7 to Marlborough Sounds, and 6 to Fiordland. Only one haplotype was shared among the 3 populations. Another haplotype was shared between Marlborough Sounds and Fiordland, and a third was shared between Northland and Marlborough Sounds (Figure 2).



**Figure 2.** Phylogenetic reconstruction (NJ with HKY + I distance correction) of bottlenose dolphin mtDNA control region sequences, with bootstrap support (>50%) and rooted to the rough-toothed dolphin (*Steno bredanensis*). Shared haplotypes are shaded. N = Northland, MS = Marlborough Sounds, and F = Fiordland.

As expected from the strong frequency differences in haplotypes, the AMOVA results showed a high level of differentiation among the 3 regional populations ( $F_{\rm ST}=0.171,\ P<0.001;\ \Phi_{\rm ST}=0.206,\ P<0.001)$ . Pairwise  $F_{\rm ST}$  and  $\Phi_{\rm ST}$  comparisons showed that all 3 populations differed significantly from one other (Table 2). This was confirmed by an exact test of population differentiation based on haplotype frequencies. For such diverse populations,  $F_{\rm ST}$  values are likely to be less informative regarding population divergence than  $\Phi_{\rm ST}$ , which incorporates both haplotype frequency and sequence divergence among haplotypes (Excoffier et al. 1992).

Northland had the highest estimates of haplotipic ( $b=0.88\pm0.01$ ) and nucleotide diversity ( $\pi=1.9\%\pm1$ ). Fiordland was the next most diverse population ( $b=0.76\pm0.07$ ,  $\pi=1.5\%\pm0.8$ ), and Marlborough Sounds was the least diverse population ( $b=0.73\pm0.04$ ;  $\pi=1.4\%\pm0.7$ ; Figure 2).

The high level of differentiation indicated by the AMOVA was reflected in low levels of female migration estimated in Lamarc (Table 3). The ML coalescent results

**Table 2.** Pairwise  $F_{\rm ST}$  (lower diagonal) and  $\Phi_{\rm ST}$  (upper diagonal) with their respective P values for the 3 New Zealand *Tursiops truncatus* populations

	Northland	Marlborough Sounds	Fiordland		
	n = 127	$\overline{n=42}$	n = 24		
N MS	$0.168 \ (P < 0.001)$	$0.194 \ (P < 0.05)$	0.197 (P < 0.05) 0.298 (P < 0.05)		
F	$0.150 \ (P < 0.001)$	$0.239 \ (P < 0.001)$	0.27 (- 0.00)		

N, Northland; MS, Marlborough Sounds; and F, Fiordland.

were discarded as migration rates and theta  $(\theta)$  values did not stabilize over 5 runs, whereas both parameters stabilized when Bayesian searches were performed. Although CIs overlapped in all pairwise comparisons, some asymmetries in exchange rate was indicated: we found relatively low rates of female migration from both Marlborough Sounds and Fiordland to Northland; low rates of female migration between Marlborough Sounds and Fiordland; and rates of female migration from Northland to Marlborough Sounds and to Fiordland were estimated to be several fold higher than the reverse migration (Table 3).

## Worldwide T. truncatus Genetic Diversity and Population Structure

To explore the phylogeographic relationship of New Zealand bottlenose dolphins to other populations worldwide, we analyzed a total of 586 samples representing 19 regional populations (Supplementary Appendix 1). The total length of sequences varied from 294 to 720 bp, allowing a consensus length of 391 bp for all analyses. For sequences shorter than 391 bp, haplotype identity was inferred from the available length. Although some potentially variable sites were not available for sequences less than 391 bp, visual inspection of the dataset showed that there was no ambiguity in defining unique haplotypes. However, 3 New Zealand haplotypes collapsed (NZ-FJB2 with NZ-N18, NZ-F10 with NZ-F02, and NZ-N38 with NZ-N05) when sequences were shortened to 391 bp. Examination of this 391-bp fragment revealed 89 variable sites defining 135 unique haplotypes. There were 82 transition substitutions, 9 transversion substitutions (including 2 sites showing both a transition and a transversion), and 4 single base insertiondeletions. For analyses of population structure and diversity, 5 regions represented by low sample numbers (a total of 7 samples of 5 haplotypes) were excluded, bringing the total number of samples analyzed to 579 and representing 130 unique haplotypes from 14 populations (including New Zealand, Table 1).

Haplotypic diversity of the 14 populations ranged from  $b = 0.43 \pm 0.11$  for the WNAi to  $b = 0.95 \pm 0.04$  for China (unknown), whereas nucleotide genetic diversity ranged from  $\pi = 0.5\% \pm 0.30$  for Bahamas (inshore) to  $\pi = 2.2\%$  for the Caribbean, Hawai'i, New Zealand (unknown), and WNAo (Table 1).

Overall, regional populations were highly differentiated  $(F_{\rm ST} = 0.16 \text{ and } \Phi_{\rm ST} = 0.34; P < 0.0001)$ . After applying sequential Bonferroni corrections, most pairwise comparisons remained significant for both  $F_{ST}$  (71 out of 91) and  $\Phi_{\rm ST}$  (83 out of 91; Table 4). The small sample size of some populations (i.e., Palmyra Atoll n = 11, Gulf of Mexico n = 10, and Bahamas n = 7; Table 1) may explain these nonsignificant results. There were few shared haplotypes among regional populations worldwide suggesting low levels of female migration (Supplementary Appendix 1). Among oceans, there was one shared haplotype between Japan (North Pacific) and New Zealand/Samoa (South Pacific), 4 between Hawai'i/Palmyra Atoll (North Pacific) and the Republic of Kiribati (South Pacific), and 1 between Palmyra Atoll (North Pacific) and French Polynesia (South Pacific; Supplementary Appendix 1). No haplotypes were shared between the Atlantic and Pacific Oceans.

#### Population Structure by Ecotype and Ocean Basin

We considered population differentiation for 2 higher order groupings: ecotype and ocean basin (Table 1). Unfortunately, a hierarchical analysis of these 2 groupings was not possible because of the imbalance of ecotype classification within oceans. Instead, we conducted 2 nonhierarchical AMOVA analyses including the entire dataset. Pairwise  $F_{\rm ST}$  and  $\Phi_{\rm ST}$  comparisons by ecotype (inshore, offshore, and unknown) showed that all 3 were significantly different, irrespective of ocean origin (overall  $F_{\rm ST}=0.110$ ;  $\Phi_{\rm ST}=0.344$ , P<0.0001 for both; Table 5);  $F_{\rm ST}$  and  $\Phi_{\rm ST}$  values showed far less difference between the offshore and unknown ecotypes than either of those to the inshore ecotype. This pattern was mirrored in the net and gross average sequence divergences (Table 6). Unknown and

**Table 3.** Most probable estimates of female migration rates per generation ( $N_{\rm mf}$ ) using Bayesian analysis between the 3 *Tursiops truncatus* populations in New Zealand

	То						
Migration from	Northland	Marlborough Sounds	Fiordland				
N		3.99  (CI = 0.44-20.52)	4.89 (CI = 0.02–20.32)				
MS	0.40  (CI  = 0.03-2.36)		0.31  (CI  = 0.00-3.12)				
F	0.19  (CI = 0.00-1.70)	0.29  (CI  = 0.00-2.01)	, ,				

N, Northland; MS, Marlborough Sounds; F, Fiordland; and CI, confidence interval.

**Table 4.** Pairwise  $F_{ST}$  (lower diagonal) and  $\Phi_{ST}$  (upper diagonal) for 14 regional bottlenose dolphin populations worldwide (populations with <5 samples were excluded)

	NZ	KI	Ja	Ch	Hi	PA	GM	Ca	Ba	WNAi	WNAo	ENA	MS	WA
New Zealand		0.255	0.174	0.134	0.059	0.196	0.468	0.267	0.473	0.523	0.132	0.364	0.166	0.205
Kiribati	0.125		0.612	0.551	0.349	0.149	0.817	0.576	0.815	0.813	0.466	0.767	0.523	0.619
Japan	0.150	0.204		0.124	0.183	0.550	0.677	0.488	0.717	0.767	0.328	0.643	0.437	0.485
China	0.071	0.109	0.127		0.100	0.461	0.650	0.440	0.667	0.737	0.226	0.620	0.340	0.434
Hawai'i	0.111	0.141	0.161	0.093		0.290	0.517	0.315	0.515	0.585	0.159	0.435	0.205	0.273
Palmyra Atoll	0.091	0.055	0.173	0.068	0.110		0.706	0.379	0.675	0.721	0.323	0.646	0.308	0.428
Gulf of Mexico	0.117	0.164	0.201	0.097	0.142	0.123		0.470	0.669	0.769	0.581	0.771	0.597	0.707
Caribbean	0.127	0.174	0.210	0.110	0.152	0.136	0.168		0.438	0.631	0.201	0.439	0.202	0.189
Bahamas	0.111	0.159	0.199	0.089	0.137	0.115	0.112	0.163		0.711	0.532	0.753	0.569	0.683
WNAi	0.276	0.382	0.389	0.325	0.308	0.385	0.418	0.414	0.432		0.624	0.762	0.640	0.727
WNAo	0.105	0.146	0.181	0.086	0.129	0.109	0.138	0.149	0.131	0.356		0.380	0.100	0.144
ENA	0.169	0.224	0.251	0.166	0.194	0.195	0.224	0.232	0.223	0.413	0.194		0.222	0.369
MS	0.080	0.119	0.157	0.056	0.103	0.077	0.107	0.113	0.099	0.345	0.095	0.102		0.089
WA	0.165	0.220	0.251	0.157	0.191	0.189	0.220	0.229	0.219	0.447	0.194	0.235	0.072	

After Bonferroni correction (P < 0.00055), some pairwise comparisons were not significant (indicated in bold). NZ, New Zealand; KI, Republic of Kiribati; Ja, Japan; Ch, China–Taiwan; Hi, Hawai'i; PA, Palmyra Atoll; GM, Gulf of Mexico; Ca, Caribbean; Ba, Bahamas; WNAi, Western North Atlantic inshore; WNAo, Western North Atlantic offshore; ENA, Eastern North Atlantic; MS, Mediterranean Sea; and WA, West Africa.

offshore ecotypes presented higher values of genetic diversity than the inshore ecotype at both the haplotype and nucleotide level (unknown:  $b = 0.97 \pm 0.002$ ,  $\pi = 2.8\% \pm 1.4$ ; offshore:  $b = 0.88 \pm 0.05$ ,  $\pi = 2.2\% \pm 1.2$ ; and inshore:  $b = 0.76 \pm 0.006$ ,  $\pi = 1.9\% \pm 1.0$ ).

Pairwise  $F_{\rm ST}$  and  $\Phi_{\rm ST}$  comparisons confirmed that the 3 ocean basins were significantly different, irrespective of ecotype classification (overall  $F_{\rm ST}=0.067$ ;  $\Phi_{\rm ST}=0.174$ , P<0.0001 for both; Table 7).

Overall, pairwise  $\Phi_{ST}$  were higher than  $F_{ST}$  values as a result of high haplotypic diversity within populations and some extent of sequence divergence among populations (Supplementary Appendix 1).

Slightly higher values of genetic diversity at both the haplotype and nucleotide level were found in the AO ( $b = 0.95 \pm 0.008$ ,  $\pi = 2.8\% \pm 1.4$ ) compared with the NP ( $b = 0.93 \pm 0.008$ ,  $\pi = 2.2\% \pm 1.1$ ) or the SP ( $b = 0.92 \pm 0.06$ ,  $\pi = 2.6\% \pm 1.0$ ).

## New Zealand Compared with Worldwide Populations

At a regional level, a dendrogram reconstruction based on sequence divergence ( $d_a$ ) among worldwide populations suggested that New Zealand was more divergent from those populations found in the Atlantic Ocean than from those in

**Table 5.** Pairwise  $F_{\rm ST}$  (lower diagonal) and  $\Phi_{\rm ST}$  (upper diagonal) of *Tursiops truncatus* ecotypes: inshore (I), offshore (O), and unknown (U)

	Inshore (I)	Offshore (O)	Unknown (U)		
	$\overline{n=46}$	$\overline{n=47}$	$\overline{n=493}$		
I		0.392	0.423		
O	0.184		0.079		
U	0.121	0.071			

For all comparisons, P < 0.001.

the Pacific Ocean (Figure 3). In terms of ecotype, New Zealand and the CWP were more divergent from populations described as inshore (WNAi, Gulf of Mexico, and Bahamas) than from the offshore form (WNAo; Table 5), regardless of the habitat where samples were collected (coastal or pelagic) or ocean basin.

## Worldwide Phylogeography

A statistical parsimony analysis revealed a very complex network of haplotypes with 31 closed loops, including 6 sequences connected to more than 7 other sequences each. There were 7 loops that could be resolved in more than one way potentially leading to different connections among haplotypes. There was no obvious pattern of monophyly of mtDNA lineages by ocean basin, regional population, or ecotype (Figure 4). However, samples described as inshore in the literature (WNAi, Gulf of Mexico, and Bahamas) clustered together whereas offshore or unknown ecotype origin haplotypes were scattered throughout the reminder of the network. Two haplotypes sampled in the Caribbean (Car-PR610 and Car-PR616) that were of unknown ecotype origin shared one fixed difference with the inshore group, suggesting that these samples belonged to the inshore ecotype.

## **Discussion**

Our study presents one of the most comprehensive analyses of mtDNA structure and diversity of bottlenose dolphins to date in terms of sample size (586 individuals) and geographic sample coverage (19 populations) spanning 3 ocean basins. Our study includes and expands on the analysis of mtDNA by Natoli et al. (2004) by greatly increasing the sample size and geographic coverage for the Pacific Ocean. The scope of the analyses allowed us to place

**Table 6.** Average net  $(d_a;$  lower diagonal), gross  $(d_{xy};$  upper diagonal) sequence divergence between populations and within population diversity  $(d_x \text{ and } d_y;$  diagonal) among New Zealand (NZ), published inshore, offshore, and unknown ecotypes including standard errors (SEs)

	New Zealand	Offshore	Inshore	Unknown	
	n = 209	$\overline{n=25}$	$\overline{n=46}$	$\overline{n=306}$	
NZ Offshore Inshore Unknown	2.5% (SE = 0.7) 0.5% (SE = 0.2) 2.4% (SE = 0.9) 0.1% (SE = 0.1)	2.8% (SE = 0.7) 2.1% (SE = 0.6) 2.1% (SE = 0.8) 0.3% (SE = 0.1)	4.6% (SE = 1.2) 4.2% (SE = 1.1) 1.7% (SE = 0.5) 1.9% (SE = 0.6)	2.8% (SE = 0.7) 2.7% (SE = 0.7) 4.4% (SE = 1.1) 2.6% (SE = 0.7)	

the regional differences found among New Zealand coastal populations in a worldwide context.

# Coastal New Zealand Populations Are Isolated but Retain Surprisingly High Diversity

Results from our study confirmed a high degree of isolation among New Zealand coastal populations. Significant population structure over relatively small geographic distances has been documented in several T. truncatus populations inhabiting coastal areas, including those along the coasts of the Gulf of Mexico (Duffield and Wells 1991; Sellas et al. 2005), the Bahamas (Parsons et al. 2006), and Western Australia, although the latter included individuals of uncertain taxonomy (Krützen et al. 2004). Parsons et al. (2006) suggested that the scale of population subdivision in this species reflects the genetic consequences of their social system and site fidelity. Bottlenose dolphins form stable, long-lasting associations, with individuals often showing strong site fidelity (Wells 1991), even in pelagic groups (Rossbach and Herzing 1999). However, in New Zealand, the only population that shows a high degree of local site fidelity is Doubtful Sound in Fiordland (Schneider 1999; Lusseau 2003). In the Bay of Islands, where the population has been studied intensively, there are no resident individuals, but rather a subset of regular users and infrequent visitors (Constantine 2002). A similar pattern seems to occur in Marlborough Sounds (Merriman et al. 2005).

Despite restricted female migration and significant population structure, all New Zealand populations showed relatively high genetic diversity (overall h = 0.91,  $\pi = 2.2\%$ ) given their relatively small population sizes and degree of isolation. Natoli et al. (2004) reported haplotype diversity values ranging from h = 0.43 to 0.72 for coastal T. truncatus populations and from h = 0.73 to 0.94 for pelagic ones, concluding that coastal populations had comparatively lower genetic diversity. Krützen et al. (2004) analyzed 220 mtDNA control region sequences from coastal Tursiops sp. from Western Australia and identified only 8 unique haplotypes with a diversity of h = 0.66. In contrast, the analysis of 127 mtDNA control region sequences from Northland represented 15 unique haplotypes with a value of b = 0.88. The relatively high genetic diversity encountered in New Zealand, particularly Northland, is not explained by current population sizes or rates of female migration between local populations; the same pattern of high diversity was observed in most populations worldwide, except for those described as belonging to the inshore ecotype.

## Bottlenose Dolphins Experience Long-distance Gene Flow

Results from a test of differentiation and the haplotype network suggested that restricted gene flow with longdistance dispersal events occurs between all populations except for those described as inshore ecotypes in the Western North Atlantic. The extent of movement by pelagic T. truncatus populations is poorly understood but it is thought to include at least occasional long-distance movements (Leatherwood and Reeves 1982; Wells et al. 1999). A recent study in the North Atlantic suggested that pelagic bottlenose dolphins are able to maintain high levels of gene flow over large distances (Quérouil et al. 2007). Additionally, Goodall et al. (2004) reported strandings of 6 bottlenose dolphins along the coast of Tierra del Fuego (55°S) and one live sighting in the Beagle Channel (Tierra del Fuego), suggesting that there is potential for ongoing interchange between the South Atlantic and South Pacific Oceans.

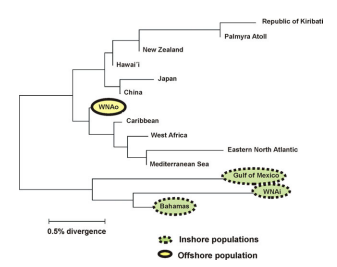
## Habitat Specialization and Ecotypes Occur Independently between Oceans

As suggested by Natoli et al. (2004), the divergence of inshore WNA populations could have occurred for a variety of reasons including founder events from pelagic populations with subsequent philopatry. Without genetic input from other sources, small isolated populations are prone to the effects of genetic drift diverging from the parental population and losing genetic diversity over time (Lacy 1987). On the other hand, differences in resource use could lead to assortative mating or ecological separation resulting in genetic differentiation (Hoelzel 1998).

**Table 7.** Pairwise  $F_{\rm ST}$  (lower diagonal) and  $\Phi_{\rm ST}$  (upper diagonal) of *Tursiops truncatus* by ocean basins

	North Pacific	South Pacific	Atlantic Ocean			
	n = 192	n = 236	n = 158			
NP		0.063	0.251			
SP	0.070		0.216			
AO	0.067	0.063				

AO, Atlantic Ocean; NP, North Pacific; and SP, South Pacific. For all comparisons, P < 0.001.



**Figure 3.** Dendrogram showing mtDNA control region sequence divergence  $(d_a)$  among worldwide regional populations of bottlenose dolphins based on a midpoint rooting NJ reconstruction. Inshore and offshore populations refer to the origins of sequences of known ecotype. WNAo = Western North Atlantic offshore and WNAi = Western North Atlantic inshore.

Analyses conducted here showed that populations described as the inshore ecotype are highly differentiated from all other populations worldwide and restricted to the WNA, supporting previous suggestions that this ecotype could represent a different species or subspecies. Differences in ecology (distribution, foraging, and parasite load), morphology, and genetics led Mead and Potter (1990) to suggest that the WNA inshore and offshore ecotypes could be considered different species. Using nuclear markers amplified fragment length polymorphism (AFLP), Kingston and Rosel (2004) found that inshore and offshore ecotypes of *T. truncatus* in the WNA exhibited greater divergence than the 2 different species of common dolphin (Delphinus delphis and Delphinus capensis), also suggesting that the 2 ecotypes could represent different species. Interestingly, in the Indian Ocean and some (but not all) regions of the Pacific Ocean, populations thought to represent *T. aduncus* fill the ecological niche of this inshore T. truncatus ecotype.

# Tursiops truncatus Offshore and Unknown Ecotypes Are Evolutionary Interconnected

Bottlenose dolphins found in coastal waters of New Zealand and CWP were genetically more divergent from those populations classified as inshore than from those classified as the offshore ecotype as described in the WNA. The WNA offshore ecotype seems to be genetically related to a number of worldwide haplotypes from populations found in coastal and pelagic habitats suggesting that, in contrast to the WNA inshore ecotype, its origins are not habitat specific. This supports the hypothesis that habitat

use and ecotype have evolved independently in different oceans. If so, the pattern and evolutionary processes leading to highly differentiated ecotypes in the WNA are not entirely representative of T. truncatus worldwide. Moreover, T. truncatus populations described as offshore and unknown ecotypes present relatively high levels of genetic diversity and degree of isolation regardless of population habitat use; however, these populations seem to be interconnected through restricted gene flow. A similar pattern was observed in another worldwide distributed dolphin species such as spinner dolphins from French Polynesia (Stenella longirostris longirostris). Significant genetic differentiation and demographic isolation among neighboring communities indicated restricted gene flow; however, the high levels of genetic diversity found contrasted with this isolation suggesting instead a metapopulation structure (Oremus et al. 2007).

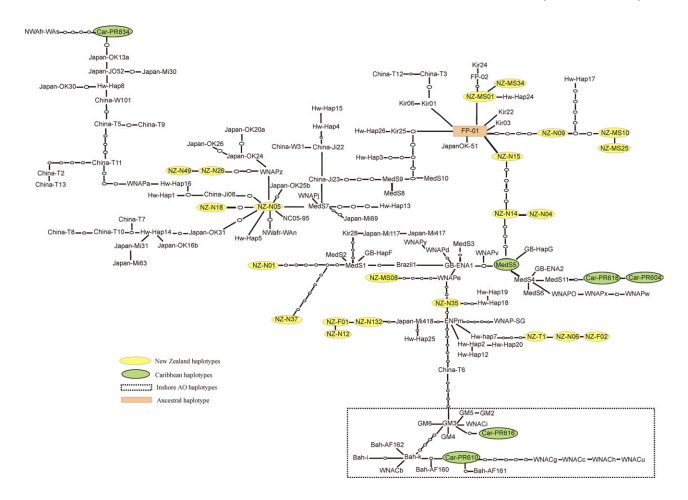
Alternatively, genetic diversity values observed in *T. truncatus* populations worldwide (except for those described as inshore ecotype) could reflect founder events due to recent colonization of coastal habitats. In this case, the observed values of genetic diversity would be a signal of the historical polymorphisms contained in large pelagic populations. However, such diversity values are unlikely to persist in small isolated populations without additional influx from other sources.

#### Conclusion

Our results suggest that the divergence of inshore populations and the formation of ecotypes in the Western North Atlantic do not necessarily reflect the worldwide pattern of T. truncatus; moreover, habitat specialization seems to have occurred independently in different ocean basins. Distinct inshore populations are highly differentiated and restricted to the WNA, potentially representing a different taxonomic unit. All other populations showed significant differentiation of mtDNA lineages among worldwide regions including relatively high mtDNA diversity; however, they were not phylogeographically distinct. These results suggest that offshore and unknown ecotypes are interconnected through long-distance gene flow and/or by interchange with pelagic populations. It is not clear what evolutionary processes have led to this pattern (e.g., foraging or reproductive strategies, environmental factors, social structure). Future research is needed to characterize potential pelagic populations of T. truncatus that might be linking coastal regions in the North and South Pacific Oceans. Independent lines of evidence (e.g., nuclear DNA markers and morphology; Caballero, Trujillo, et al. 2007) would aid in better describing different ecotypes or taxonomic units of this highly versatile species throughout its range.

## Supplementary Material

Supplementary appendix table can be found at http://www.jhered.oxfordjournals.org/.



**Figure 4.** Worldwide parsimony network of *Tursiops truncatus* mtDNA control region haplotype sequences based on 19 regional populations of inshore, offshore, or unknown ecotype origin. Missing or unsampled intermediaries are shown by a small oval (O). Regional origins of haplotype sequences are indicated in the legend and in Supplementary Appendix 1.

## **Funding**

Northland Marine Mammal Trust; Department of Conservation (Northland) Contract 012438-001-AU1; J. Watson Conservation Trust from the Royal Forest and Bird Society; Postgraduate Tuition Fee Bursary (University of Auckland) and the Whale and Dolphin Adoption Project to G.T.P.; Marsden Fund of the Royal Society of New Zealand to C.S.B.; Department of Conservation (West Coast) to K.R; Puerto Rico's Legislature and Dolphin Quest to A.A.M.-G; and Natural Heritage Trust from the Department of the Environment and Heritage (Australian Government) and International Fund for Animal Welfare to C.O.

## **Acknowledgments**

Logistic support in New Zealand was provided by the Department of Conservation (Northland, Marlborough Sounds, Hokitika, and Te Anau); specially T. Beauchamp, J. Beachman, A. Walker (DoC Northland), R. Kemper, B. Masser (DoC Te Anau), and D. Neale (DoC Hokitika). Logistic support in the Republic of Kiribati was provided by the National

Geographic Magazine, the New England Aquarium, and the government of the Republic of Kiribati. We thank the following for assistance in fieldwork: P. Gay, P. Norris, C. Paterson, R. Gosh, and B. Woodward (Doubtful Sound); C. Carraher, A. Engelhaupt and D. Engelhaupt (Marlborough Sounds); R. Constantine, S. Wells, D. Pouwels, F. Mõurao, J. Brueggeman, K. McLeod, E. Newcombe, A. Fleming, C. Clark, D. Heimeier, J. Jackson, and E. Carroll (Northland); N. Wiseman, K. Stockin, B. Doak, B. Murray and A. Cozens (Hauraki Gulf); N. Wiseman and K. Scollay (Jackson Bay); S. Walsh, J. Ward, L. Bell, M. Iakopo, and S. Tufuga (Samoa); N. Funahashi (Japan); D. L. Webster, A. D. Ligon, and D. J. McSweeney (Hawai'i). K. Robertson and B. Hancock conducted laboratory analyses of the samples from Hawai'i and Palmyra. J. Jackson and D. Steel assisted with data analysis. A. Alexander and D. Pouwels reviewed an early version of the manuscript. J. Jackson and 2 anonymous reviewers provided valuable comments on the final manuscript.

Biopsy samples were collected under the following permits and animal ethics protocols: in New Zealand, under permit to C.S.B. from the New Zealand Department of Conservation and animal ethics protocols AEC/02/2002/R9 and AEC/02/2005/R334 from the University of Auckland; in Samoa, under permits to C.O. granted by the Ministry of the Prime Minister and Cabinet, Ministry of Natural Resources and Environment, Police and Prisons Department of the Samoan Government, and the Department of Environment and Heritage (Australia); in New Caledonia to C.G. under letter of authorization 6024-1028-DRN/ENV from the North and South

provinces; in French Polynesia to M.O. under permit from the local government and direction d'Environnement; in the Republic of Kiribati to G.S. and the New England Aquarium under permits granted by the local government; in Hawai'i and Palmyra Atoll to R.W.B. and K.M. under Marine Mammal Protection Act permit 774-1714, National Marine Fisheries Service and CITES permits; in the Caribbean to A.A.M.-G., under Marine Mammal Protection Act permits 779-1339, 779-1633, 774-1714 and CITES permits 04US774223/9 and 05US774223/9, issued to the National Marine Fisheries Service, and under permit 04-EPPE-003 from Puerto Rico's Department of Natural and Environmental Resources and a cooperative agreement with the United States Virgin Islands Department of Planning and Natural Resources.

## References

Baker CS, Lento GL, Cipriano F, Palumbi SR. 2000. Predicted decline of protected whales based on molecular genetic monitoring of Japanese and Korean markets. Proc R Soc Lond B Biol Sci. 267:1191–1199.

Baker CS, Lukoschek V, Lavery S, Dalebout ML, Yong-un M, Endo T, Funahashi N. 2006. Incomplete reporting of whale, dolphin and porpoise 'bycatch' revealed by molecular monitoring of Korean markets. Anim Conserv. 9:474–482.

Baker CS, Palumbi SR. 1994. Which whales are hunted? A molecular genetic approach to monitoring whaling. Science. 265:1538–1539.

Baker CS, Perry A, Bannister JL, Weinrich MT, Abernethy RB, Calambokidis J, Lien J, Lambertsen RH, Urban-Ramirez J, Vasquez O, et al. 1993. Abundant mitochondrial DNA variation and world-wide population structure in humpback whales. Proc Natl Acad Sci USA. 90:8239–8243.

Baker CS, Slade RW, Bannister JL, Abernethy RB, Weinrich MT, Lien J, Urban-Ramirez J, Corkeron P, Calambokidis J, Vasquez O, et al. 1994. Hierarchical structure of mitochondrial DNA gene flow among humpback whales *Megaptera novaeangliae*, world-wide. Mol Ecol. 3:313–327.

Boisseau O. 2003. A summary of research conducted on bottlenose dolphins of Fiordland. Invercargill (New Zealand): Southland Conservancy, Department of Conservation. p. 98.

Bowen BW, Avise JC. 1994. Conservation research and the legal status of PCR products. Science. 266:713.

Bräger S, Schneider K. 1998. Near-shore distribution and abundance of dolphins along the West Coast of the South Island, New Zealand. N Z J Mar Freshwater Res. 32:105–112.

Caballero S, Jackson J, Mignucci-Giannoni AA, Barrios-Garrido H, Beltrán-Pedreros S, Montiel-Villalobos MG, Robertson KM, Baker CS. 2008. Molecular systematics of South American dolphins *Sotalia*: sister taxa determination and phylogenetic relationships, with insights into a multilocus phylogeny of the *Delphinidae*. Mol Phylogenet Evol. 46:252–268.

Caballero S, Trujillo F, Vianna J, Barrios-Garrido H, Montiel M, Beltrán-Pedreros S, Marmontel M, Santos M, Rossi-Santos M, Santos F, et al. 2007. Taxonomic status of the genus *Sotalia*: species level ranking for "Tucuxi" (*Sotalia fluviatilis*) and "Costero" (*Sotalia guianensis*) dolphins. Mar Mamm Sci. 23:358–386.

Clement M, Posada D, Crandall KA. 2000. TCS: a computer program to estimate gene genealogies. Mol Ecol. 9:1657–1659.

Constantine R. 2002. The behavioural ecology of the bottlenose dolphins (Tursiops truncatus) of Northeastern New Zealand: a population exposed to tourism [dissectation]. Auckland (New Zealand)]: The University of Auckland. p. 195.

Crandall KA, Templeton AR. 1993. Empirical tests of some predictions from coalescent theory with applications to intraspecific phylogeny reconstruction. Genetics. 134:959–969.

Crandall KA, Templeton AR, Sing CF. 1994. Intraspecific phylogenetics: problems and solutions. In: Scotland R, Siebert D, Williams D, editors.

Models in phylogeny reconstructions. Oxford (UK): Claredon Press. p. 273–297.

Curry BE. 1997. Phylogenetic relationship among bottlenose dolphins (genus Tursiops) in a worldwide context [dissertation]. [College Station (TX)]: Texas A&M University at Galveston.

Curry BE, Smith J. 1997. Phylogenetic structure of the bottlenose dolphins (*Tursiups truncatus*): stock identification and implications for management. In: Dizon AE, Chivers SJ, Perrin WF, editors. Molecular Genetics of Marine Mammals. Lawrence (KS): Allen Press Inc. p. 227–247.

Dalebout ML, Van Helden A, Van Waerebeek K, Baker CS. 1998. Molecular genetic identification of southern hemisphere beaked whales (*Cetacea: Ziphiidae*). Mol Ecol. 7:687–695.

Dowling TE, Brown WM. 1993. Population structure of bottlenose dolphins (*Tursiops truncatus*) as determined by restriction endonuclease analysis of mitochondrial DNA. Mar Mamm Sci. 9:138–155.

Duffield DA, Ridgway SH, Cornell LH. 1983. Hematology distinguishes coastal and offshore forms of dolphins (*Tursiops*). Can J Zool. 61:930–933.

Duffield DA, Wells RS. 1991. The combined application of chromosome, protein and molecular data for the investigation of social unit structure and dynamics in *Tursiops truncatus*. In: Hoelzel AR, editor. Genetic ecology of whales and dolphins: report of the International Whaling Commission 13. Cambridge (UK): The International Whaling Commission. p. 155–169.

Endo T, Hotta Y, Haraguchi K, Sakata M. 2003. Mercury contamination in the red meat of whales and dolphins marketed for human consumption in Japan. Environ Sci Technol. 37:2681–2685.

Endo T, Hotta Y, Haraguchi K, Sakata M. 2005. Distribution and toxicity of mercury in rats after oral administration of mercury-contaminated whale red meat marketed for human consumption. Chemosphere. 61:1069–1073.

Excoffier L, Smouse PE, Quattro JM. 1992. Analysis of molecular variance inferred from metric distances among DNA haplotypes: applications to human mitochondrial DNA restriction data. Genetics. 131:479–491.

Felsenstein J. 1985. Confidence limits on phylogenies: an approach using the bootstrap. Evolution. 39:783–791.

Goodall RNP, Boy CC, Pimper LE, Macnie SM. 2004. Range extensions and exceptional records of cetaceans for Tierra del Fuego. In: SOLAMAC, editor. The 11th Reunion of Specialists of Aquatic Mammals of South America (11RT), and the 5th Congress of the Latin American Aquatic Mammal Specialists Group (5th SOLAMAC); 2004 Sept 12–17. Quito (Ecuador): Sociedad Latinoamericana de Mamiferos Acuaticos (Latinamerican society of aquatic mammals). p. 223.

Goodwin JA, Durham BD, Peddemors VM, Cockcroft VG. 1996. Genetic variation in the bottlenose dolphin *Tursiops truncatus* along the Kwazulu/Natal Coast, South Africa. S Afr J Marine Sci. 17:225–232.

Gormley AM. 2002. Use of mark-recapture for estimating animal abundance of four marine mammal species in New Zealand [dissertation]. [Dunedin (New Zealand)]: University of Otago.

Hasegawa M, Kishino H, Yano T. 1985. Dating of the human-ape splitting by a molecular clock of mitochondrial DNA. J Mol Evol. 22:1432–1432.

Hersh SL, Duffield DA. 1990. Distinction between northwest Atlantic offshore and coastal bottlenose dolphins based on hemoglobin profile and morphometry. In: Leatherwood S, Reeves RR, editors. The bottlenose dolphin. New York: Academy Press. p. 160–174.

Hoelzel AR. 1998. Genetic structure of cetacean populations in sympatry, parapatry, and mixed assembles: implications for conservation policy. J Hered. 89:451-458.

Hoelzel AR, Potter CW, Best PB. 1998a. Genetic differentiation between parapatric "nearshore" and "offshore" populations of the bottlenose dolphin. Proc R Soc Lond B Biol Sci. 265:1177–1183.

Jones M. 1994. PCR products and CITES. Science. 266:1930.

Kakuda T, Tajima Y, Arai K, Kogi K, Hishii T, Yamada K. 2002. On the resident "bottlenose dolphins" from Mikura waters. Mem Nat Sci Mus. 38:255–272.

Kenney RD. 1990. Bottlenose dolphins off the northeastern United States. In: Leatherwood S, Reeves RR, editors. The bottlenose dolphin. New York: Academic Press. p. 369–386.

Kingston SE, Rosel PE. 2004. Genetic differentiation among recently diverged Delphinid taxa determined using AFLP markers. J Hered. 95:1–10.

Krützen M, Barre LM, Moller LM, Heithaus MR, Simms C, Sherwin WB. 2002. A biopsy system for small cetaceans: darting success and wound healing in *Tursiops spp.* Mar Mamm Sci. 18:863–878.

Krützen M, Sherwin WB, Berggen P, Gales N. 2004. Population structure of an inshore cetacean revealed by microsatellite and mtDNA analysis: bottlenose dolphins (*Tursiops sp.*) in Shark Bay, Western Australia. Mar Mamm Sci. 20:28–47.

Kuhner MK. 2006. LAMARC 2.0: maximum likelihood and Bayesian estimation of population parameters. Bioinformatics. 22:768–770.

Kuhner MK, Yamato J, Beerli P, Smith L, Rynes E, Walkup E, Li C, Sloan J, Colacurcio P, Felsenstein J. 2005. LAMARC version 2.0 [Internet]. [cited 2007 June 4]. Available from: http://evolution.gs.washington.edu/lamarc/

Kumar S, Tamura K, Jakobsen IB, Nei M. 2001. MEGA2: Molecular Evolutionary Genetics Analysis software. Tempe (AZ): Arizona State University.

Lacy RC. 1987. Loss of genetic diversity from managed populations: interacting effects of drift, mutation, immigration, selection, and population subdivision. Conserv Biol. 1:143–157.

Leatherwood S, Reeves R. 1982. Bottlenose dolphin *Tursiops truncatus* and other toothed cetaceans. In: Chapman JA, Feldhamer GA, editors. Wild mammals of North America: biology, management, and economics. Baltimore (MD): The Johns Hopkins University Press. p. 369–414.

Leatherwood S, Reeves R, Foster L. 1983. The Sierra Club handbook of whales and dolphins. San Francisco (CA): Sierra Club Books.

LeDuc RG, Perrin WF, Dizon AE. 1999. Phylogenetic relationships among delphinid cetaceans based on fully cytochrome b sequences. Mar Mamm Sci. 15:619–648.

Lowther JL. 2006. Genetic variation of coastal and offshore bottlenose dolphins, *Tursiops truncatus*, in the Eastern North Pacific Ocean [dissertation]. [San Diego (CA)]: University of San Diego.

Lusseau D. 2003. Male and female bottlenose dolphins *Tursiops spp.* have different strategies to avoid interactions with tour boats in Doubtful Sounds, New Zealand. Mar Ecol Prog Ser. 257:267–274.

Maddison WP, Maddison DR. 2003. MACCLADE (Version 4.06): analysis of phylogeny and character evolution. Sunderland (MA): Sinauer Associates.

Mead JG, Potter CW. 1990. Natural history of bottlenose dolphins along the central Atlantic coast of the United States. In: Leatherwood S, Reeves RR, editors. The bottlenose dolphin. New York: Academy Press. p. 165–195.

Merriman MG, Markowitz T, Harlin AD. 2005. Occurrence and site fidelity of bottlenose dolphins (*Tursiops truncatus*) in the Marlborough Sounds, New Zealand. 16th Biennial Conference on the Biology of Marine Mammals; Dec 12–15, 2005. San Diego (CA): Society of Marine Mammalogy. p. 190.

Moller LM, Beheregaray LB. 2001. Coastal bottlenose dolphins from southeastern Australia are *Tursiops aduncus* according to sequences of the mitochondrial DNA control region. Mar Mamm Sci. 17:249–263.

Natoli A, Peddemors VM, Hoelzel AR. 2004. Population structure and speciation in the genus *Tursiops* based on microsatellite and mitochondrial DNA analyses. J Evol Biol. 17:363–375.

Nei M. 1987. Genetic variation within species. Molecular evolutionary genetics. New York: Columbia University Press. p. 176–207.

Oremus M. 2008. Genetic and demographic investigation of population structure and social system in four delphinid species [dissertation]. [Auckland (New Zealand)]: University of Auckland.

Oremus M, Poole MM, Steel D, Baker CS. 2007. Isolation and interchange among insular spinner dolphin communities in the South Pacific revealed by individual identification and genetic diversity. Mar Ecol Prog Ser. 336: 275–289.

Parsons KM, Dallas JF, Claridge DE, Durban JW, Balcomb KC, Thompson PM, Noble LR. 1999. Amplifying dolphin mitochondrial DNA from faecal plumes. Mol Ecol. 8:1766–1768.

Parsons KM, Durban JW, Claridge DE, Herzing DL, Balcomb KC, Noble LR. 2006. Population genetic structure of coastal bottlenose dolphins (*Tursiops truncatus*) in the northern Bahamas. Mar Mamm Sci. 22:276–298.

Parsons KM, Noble LR, Reid RJ, Thompson PM. 2002. Mitochondrial genetic diversity and population structuring of UK bottlenose dolphins (*Tursiops truncatus*): is the NE Scotland population demographically and geographically isolated? Biol Conserv. 108:175–182.

Perrin WF, Robertson KM, Van Bree PJH, Mead JG. 2007. Cranial description and genetic identity of the holotype specimen of *Tursiops aduncus* (Ehrenberg, 1832). Mar Mamm Sci. 23:343–357.

Pichler FB, Dawson SM, Slooten E, Baker CS. 1998. Geographic isolation of Hector's dolphin populations described by mitochondrial DNA sequences. Conserv Biol. 12:676–682.

Pichler FB, Robineau D, Goodall RNP, Meyer MA, Olavarria C, Baker CS. 2001. Origin and radiation of Southern Hemisphere coastal dolphins (genus *Cephalorbynchus*). Mol Ecol. 10:2215–2223.

Posada D, Crandall KA. 1998. MODELTEST: testing the model of DNA substitution. Bioinformatics. 14:817–818.

Quérouil S, Silva AM, Freitas L, Prieto R, Magalhaes S, Dinis A, Alves F, Matos J, Mendonca D, Hammond PS, et al. 2007. High gene flow in oceanic bottlenose dolphins (*Tursiops truncatus*) of the North Atlantic. Conserv Genet. 8:1405–1419.

Raymond M, Rousset F. 1995. An exact test for population differentiation. Evolution. 49:1280–1283.

Report No.: NOAA-TM-NMFS-SWFSC-363. Reeves RR, Perrin WF, Taylor BL, Baker CS, Mesnick ML. 2004. Report of the workshop on shortcomings of Cetacean taxonomy in relation to needs of conservation and management; April 30–May 2, 2004. La Jolla (CA): U.S. Department of Commerce 94.

Rice WR. 1989. Analyzing tables of statistical tests. Evolution. 43:223-225.

Rosel PE, France SC, Wang JY, Kocher TD. 1999. Genetic structure of harbor porpoise *Phocoena phocoena* populations in the northwest Atlantic based on mitochondrial and nuclear markers. Mol Ecol. 8:S41–S54.

Ross GJB. 1977. The taxonomy of bottlenose dolphins *Tursiops* species in South African waters, with notes on their biology. Ann Cape Prov Mus. 11:135–194.

Ross GJB, Cockcroft VG. 1990. Comments on Australian bottlenose dolphins and the taxonomic status of *Tursiops aduncus* (Ehrenberg 1832). In: Leatherwood S, Reeves RR, editors. The bottlenose dolphin. New York: Academic Press. p. 101–128.

Ross HA, Lento GM, Dalebout ML, Goode M, Ewing G, McLaren P, Rodrigo AG, Lavery S, Baker CS. 2003. DNA surveillance: web-based molecular identification of whales, dolphins and porpoises. J Hered. 94:111–114.

Rossbach KA, Herzing DL. 1999. Inshore and offshore bottlenose dolphin (*Tursiops truncatus*) communities distinguished by association patterns near Grand Bahama Island, Bahamas. Can J Zool. 77:581–592.

Sambrook E, Fritsch F, Maniatis T. 1989. Molecular cloning. New York: Cold Springs Harbor Laboratory Press.

Sanino GP, Van Waerebeek K, Van Bressem MF, Pastene LA. 2005. A preliminary note on population structure in eastern South Pacific common bottlenose dolphins, *Tursiops truncatus*. J Cetacean Res Manag. 7:65–70.

Schneider K. 1999. Behaviour and ecology of bottlenose dolphins in Doubtful Sound, Fiordland, New Zealand [dissertation]. [Dunedin (New Zealand)]: University of Otago. p. 211.

Schneider S, Roessli D, Excoffier L. 2000. Arlequin: a software for population genetics data analysis. ver 2.000 [Internet]. [cited 2006 December 12]. Available from: http://lgb.unige.ch/arlequin/

Segura I, Rocha-Olivares A, Flores-Ramirez S, Rojas-Bracho L. 2006. Conservation implications of the genetic and ecological distinction of *Tursiops truncatus* ecotypes in the Gulf of California. Biol Conserv. 133:336–346.

Sellas AB, Wells RS, Rosel PE. 2005. Mitochondrial and nuclear DNA analyses reveal fine scale geographic structure in bottlenose dolphins (*Tursiops truncatus*) in the Gulf of Mexico. Conserv Genet. 6: 715–728.

Smith-Goodwin JA. 1997. A molecular genetic assessment of the population structure and variation of two inshore genera on the east coast of South Africa [dissertation]. [Grahamstown (South Africa)]: Rhodes University.

Swofford DL. 2000. PAUP\*: phylogenetic analysis using parsimony (\* and other methods). Sunderland (MA): Sinauer Associates.

Tajima F. 1983. Evolutionary relationship of DNA sequences in finite populations. Genetics. 105:437–460.

Tamura K, Nei M. 1993. Estimation of the number of nucleotide substitutions in the control region of mitochondrial DNA in humans and chimpanzees. Mol Biol Evol. 10:512–526.

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.

Templeton AR, Sing CF. 1993. A cladistic analysis of phenotypic associations with haplotypes inferred from restriction endonuclease mapping. IV. Nested analyses with cladogram uncertainty and recombination. Genetics. 134:659–669.

Torres LG, Rosel PE, D'Agrosa C, Read AJ. 2003. Improving management of overlapping bottlenose dolphin ecotypes through spatial analysis and genetics. Mar Mamm Sci. 19:502–514.

Van Waerebeek K, Reyes JC, Read AJ, McKinnon JS. 1990. Preliminary observations of bottlenose dolphins from the Pacific coast of South America. In: Leatherwood S, Reeves RR, editors. The bottlenose dolphin. New York: Academic Press. p. 143–154.

Walsh P, Metzger D, Higuchi R. 1991. Chelex 100 as a medium for simple extraction of DNA for PCR-based typing from forensic material. Biotechniques. 10:506–513.

Wang JY, Chou LS, White BN. 1999. Mitochondrial DNA analysis of sympatric morphotypes of bottlenose dolphins (genus: *Tursiops*) in Chinese waters. Mol Ecol. 8:1603–1612.

Wells RS. 1991. The role of long-term study in understanding the social structure of a bottlenose dolphin community. In: Pryor K, Norris KS, editors. Dolphin societies: discoveries and puzzles. Berkley (CA): University of California Press.

Wells RS, Rhinehart HL, Cunningham P, Whaley J, Baran M, Koberna C, Costa DP. 1999. Long distance offshore movements of bottlenose dolphins. Mar Mamm Sci. 15:1098–1114.

Werle E, Schneider C, Renner M, Völker M, Fiehn W. 1994. Convenient single-step, one tube purification of PCR products for direct sequencing. Nucleic Acids Res. 22:4354–4355.

Würsig B, Jefferson TA. 1990. Methods of photo-identification for small cetaceans. In: Hammond PS, Mizroch SA, Donovan GP, editors. Individual recognition of cetaceans: use of photo-identification and other techniques to estimate population parameters. Special Issue 12. Cambridge: The International Whaling Commission. p. 43–52.

Received November 14, 2007 Accepted April 15, 2008

Corresponding Editor: Steve O'Brien